

Darwin[™] Python SDK Guide

 $A SparkCognition^{TM} Education Document$

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SparkCognition Darwin Python SDK Guide

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About this guide

This guide describes using the $Darwin^{\mathbb{M}}$ SDK to access and use the Darwin API in automated model building. It is intended for data scientists, software engineers and analysts who want to use the Darwin API to interact with Darwin to create and train models, test the generated models, monitor jobs and perform analysis. The SDK also provides some convenience functions. Note that throughout this document, long key and token values are truncated, indicated by ellipses (\dots) .



Expectation

This document assumes experience of the data scientist or software engineer that is commensurate with data science techniques and associated programming tasks.

Darwin overview

Darwin is a SparkCognition $^{\text{IM}}$ tool that automates model building processes to solve specific problems. This tool enhances data scientist potential because it automates various tasks that are often manually performed. These tasks include data cleaning, latent relationship extraction, and optimal model determination. Darwin promotes rapid and accurate feature generation through both automated windowing and risk generation. Darwin quickly creates highly-accurate, dynamic models using both supervised and unsupervised learning methods.

The general workflow for simple modeling includes:

- Upload training data
- Create model
- Upload test data
- · Test the model
- · Download result artifact

Note: Darwin expects all uploaded ingestion files to be in a *rectangular* format. This means a flat file with features that span columns and data samples that span rows. Plan your data file so it fits this expectation to help prevent errors.

See the SDK example for a modeling example.

For additional information on Darwin, contact your local SparkCognition partner for access to the white paper titled: *Darwin - A Neurogenesis Platform*.

Accessing the API

This document describes the python SDK and explains how to access the Darwin API and its functionality. Additional methods to access the Darwin API include:

- through the https://darwin-api.sparkcognition.com/v1 end point
- optionally, through user created curl commands

For additional information on the Darwin API, contact your local SparkCognition partner for access to see the *SparkCognition Darwin API User Guide*.

Notes:

- An API key is necessary to use the Darwin SDK.
 Contact SparkCognition or your IT manager for an appropriate key.
- All methods return a 2-tuple, for example:

```
(True, <context-dependent-return-object>)
(False, <some-helpful-message>)
```



Darwin SDK interface

Setup Darwin SDK

Perform the following to download and setup the Darwin SDK:

- 1. Install Python 3.5 or greater. Alternatively, install *Miniconda*, from https://conda.io/miniconda.html.
- 2. Create a directory to receive the git repository clone.
- 3. Change (cd) into the new directory.
- 4. Clone the darwin-sdk repository:

```
git clone https://github.com/sparkcognition/darwin-sdk
```

5. Change into the new root directory of the *darwin-sdk* cloned darwin-sdk project:

```
cd <NewCloneRootDirectory>
```

Note: By default this is the *master* trunk.

6. Ensure code is from master trunk:

```
git pull
```

7. Setup the SDK:

```
python setup.py install
```

The SDK defaults to using the production URL: https://darwin-api.sparkcognition.com/v1/

ON-PREM ONLY: For on-prem installations, the domain name will be in the form:

customerdomainname.customerdomain.com

Connect to the Darwin interface

Obtain an api key.

To use the Darwin SDK, an api key is required. A key can be obtained from SparkCognition support or your IT manager. An api_key is a long string, for example:

 $\verb|'RsJ74ZS5AvwznbHh0AfVSgrchhS9KxACDy3jefaQnxb9f6QTSDBFmhnGa0cOIWtNAIFRAG9ToOTpi0mn'| and the control of the$

Register the api key using auth_register().

The purpose of this method is to create a password and an email address for an api_key. Each api_key is synonymous with a service. This method must be invoked once for each api_key to establish a password for that key.

Note: After successful registration, the service uses auth_login() to login as a service.



```
>>> from amb_sdk.sdk import DarwinSdk
>>> s = DarwinSdk()
>>> s.auth_register('asdf', 'iRgwut4kGs0ymULiuKtMd0WFvBYLMWSj16q2uysQeteqH9ssc+\
EETUvcysnPojRpfycLVHa2IlN1IlrfEk1YMA', 'admin@company.com')
(True,'Bearer eyJ0eXAiOiJKV1QiLCJhbGciOiJIUzI1NiJ9.eyJleHAiOjE1MTU1MzM4NjEsImlh\
dCI6MTUxNTUzMDI2MS ... F56xZQiBT-89nrRz1nIXD5LfawHIj_MlUHQqM36vU')
```

Login as a service or create a user under the service and login as a user.

The following explains how to log in as a service.

Notes:

- Although Bearer <auth-token>, returned by auth_login(), is used in subsequent calls to validate authenticity, it is not required for each method.
- The SDK remembers the auth token for the DarwinSdk object. Although an auth token is currently valid for 1 hour, if the DarwinSdk object session life time exceeds 1 hour, the SDK will request another auth token until the session ends.

Example

```
>>> s.auth_login('asdf', 'iRgwut4kGs0ymULiuKtMd0WFvBYLMWSj16q2uysQeteqH9ssc+EET\
UvcysnPojRpfycLVHa2IlN1IlrfEk1YMA')
(True,'Bearer eyJ0eXAiOiJKV1QiLCJhbGciOiJIUzI1NiJ9.eyJleHAiOjE1MTU1MzQxNzIsImlh\
dCI6MTUxNTUzMD ... UQQfoXqYFKJSoRXXDNPE985-a08cE6_o')
```

When login (as a service) successfully completes, the SDK can be used to create and model a workflow.

Note, there are also auth_register_user() and auth_login_user() methods that allow you to create users and login as a specific user. You can choose to use the SDK as a service or create users underneath the service to partition datasets/models to be owned by specific users. It is more convenient to employ user accounts because the api_key is not necessary for logging in as a user.

Example

```
>>> s.auth_register_user('atestuser', 'apassword', 'anemail')
(True,
    'Bearer eyJhbGciOiJIUzI1NiIsInR5cCI6IkpXVCJ9.eyJqdGkiOiJkNjY0MmJjOC1iMmU5LTQxO\
DctODFlNS00YjI2MD ... 5zMp_1FfxU')
>>> s.auth_login_user('atestuser', 'apassword')
(True,
    'Bearer eyJhbGciOiJIUzI1NiIsInR5cCI6IkpXVCJ9.eyJqdGkiOiI3NGYzYmUxZS0yOTlmLTRhN\
zMtODU5ZCO1NGRmM2F ... u1zGCeCONA')
```

1. Verify the connection.

The default url in the SDK is https://darwin-api.sparkcognition.com/v1/. Use get_url() and set_url() to verify connection to the right Darwin service.

ON-PREM ONLY: For on-prem installations, the domain name will be in the form:

```
customerdomainname.customerdomain.com
```



Darwin SDK methods

URL Get/Set methods

DarwinSdk.get_url()

Get Darwin service url.

Parameters: None

Returns:

```
(True, <url-string>)
```

Example

```
In [10]: s.get_url()
Out[10]: (True, 'https://darwin-api.sparkcognition.com/v1/')
```

DarwinSdk.set_url(url, version='v1')

Set Darwin service url and version.

Parameters:

- url URL to the Darwin service
- version Set to 'v1'

Returns:

```
(True, <url>) or (False, 'invalid url')
```

Example

```
In [9]: s.set_url('https://darwin-api.sparkcognition.com/v1/')
Out[9]: (True, 'https://darwin-api.sparkcognition.com/v1/')
```

Authentication methods

DarwinSdk.auth_register(password, api_key, email)

Register as a service. The purpose of this method is to set a password for an api_key. Each api_key is synonymous with a service. This method is invoked only once for each api_key to establish a password for that key. After registration, the service can use *auth_login()* to login as a service.

Parameters:



- password The service level password
- api_key The api key for the service
- email Email address

Returns:

```
(True, 'Bearer <auth-token>') or (False, <error-message>)
```

Bearer <auth-token> is used in subsequent calls to validate authenticity.

The SDK remembers the auth token for the DarwinSdk object.

Note: Although an auth token is currently valid for 1 hour, if the DarwinSdk object session life time exceeds 1 hour, the SDK will request another auth token until the session ends.

Example

```
In [4]: s.auth_register('asdf', 'RsJ74ZS5AvwznbHh0AfVSgrchhS9KxACDy\
3jefaQnxb9f6QTSDBFmhnGa0cOIWtNAIFRAG9ToOTpi0mnEo3zFA', 'email')
Out[4]:
(True,
   'Bearer eyJhbGciOiJIUzI1NiIsInR5cCI6IkpXVCJ9.eyJleHAiO...iSdU8xlF4yJk')
```

DarwinSdk.auth_login(password, api_key)

Login as a service.

Note: A service must have a password set using *auth_register()* to login successfully.

Parameters:

- password The service level password
- api_key The api key for the service

Returns:

```
(True, 'Bearer <auth-token>') or (False, <error-message>)
```

Bearer <auth-token> is used in subsequent calls to validate authenticity. The SDK remembers the auth token for the DarwinSdk object.

Note: Although an auth token is currently valid for 1 hour, if the DarwinSdk object session life time exceeds the 1 hour limit, the SDK will acquire another auth token until the session ends.

```
In [5]: s.auth_login('asdf',
   'iRgwut4kGs0ymULiuKtMd0WFvBYLMWSj16q2uysQeteqH9ssc+EETUvcysnPojRpfyc\
LVHa2IlN1IlrfEk1YMA')
Out[5]:
(True,
```



```
'Bearer
eyJ0eXAiOiJKV1QiLCJhbGciOiJIUzI1NiJ9.eyJleHAiOjE1MTU1MzQxN....')
```

DarwinSdk.auth_register_user(username, password, email)

Register a user. This method registers a new user.

Note: You must be logged in as a service to create a user.

Parameters:

- username The new end user's username
- password The new end user's password
- email The new end user's email address

Returns:

```
(True, 'Bearer <auth-token>') or (False, <error-message>)
```

Bearer <auth-token> is used in subsequent calls to validate authenticity. The SDK remembers the auth token for the DarwinSdk object.

Note: Although an auth token is currently valid for 1 hour, if the DarwinSdk object session life time exceeds the 1 hour limit, the SDK will acquire another auth token until the session ends.

Example

```
In [8]: s.auth_register_user('user1', 'user1-password', 'user-email',
   'iRgwut4kGs0ymULiuKtMd0WFvBYLMWSj16q2uysQeteqH9ssc+EETUvcysnPojRpfycLV\
Ha2IlN1IlrfEk1YMA')
Out[8]:
(True,
   'Bearer
eyJ0eXAiOiJKV1QiLCJhbGciOiJIUzI1NiJ9.eyJleHAiOjE1MTU1MzQyN....')
```

DarwinSdk.auth_set_email(username, email)

Add or change a user's email address.

Parameter:

- username The end user's username
- email The end user's email address

Returns:



```
(True, None) or (False, <error-message>)
```

User must be logged in to add or change an email address. For cloud installations, this email address will be used for password resets and other notifications. For on-prem installations, password resets are executed using a new default password, there is no email sent. If you do not know the default password, contact Darwin support.

Example

```
In [9]: s.auth_set_email('user1', 'user1@company.com')
Out [9]: (True, None)
```

DarwinSdk.auth login user(username, password)

Login as a user.

Note: A user must have a username and password set using auth_register_user() to successfully login.

Parameters:

- username The end user's username
- password The end user's password

Returns:

```
(True, 'Bearer <auth-token>') or (False, <error-message>)
```

Bearer <auth-token> is used in subsequent calls to validate authenticity. The SDK remembers the auth token for the DarwinSdk object.

Note: Although an auth token is currently valid for 1 hour, if the DarwinSdk object session life time exceeds the 1 hour limit, the SDK will acquire another auth token until the session ends.

Example

```
In [9]: s.auth_login_user('user1', 'user1-password',
'iRgwut4kGs0ymULiuKtMd0WFvBYLMWSj16q2uysQeteqH9ssc+EETUvcysnPojRpfycLV\
Ha2IlN1IlrfEk1YMA')
Out[9]:
(True,
'Bearer
eyJ0eXAiOiJKV1QiLCJhbGciOiJIUzI1NiJ9.eyJleHAiOjE1MTU1MzQzM....')
```

DarwinSdk.auth_change_password(curpass, newpass)

Change the current user's password.



Parameters::

- curpass User's current password
- newpass User's new password

Returns:

```
(True, None) or (False, <error-message>)
```

User must be logged in to change password. If the current password is forgotten, use the following **DarwinSdk.auth_reset_password (username)** method to reset it. For cloud installations, an email will be generated with a temporary password. For on-prem installations, password resets are executed using a new default password, there is no email sent. If you do not know the default password, contact Darwin support.

Example

```
In [10]: s.auth_change_password('mypassword', 'newpassword')
Out[10]: (True, None)
```

DarwinSdk.auth_reset_password(username)

Reset a user's password. Any user can reset another user's password. You do not have to be an admin to execute this function. For cloud installation, a temporary password will be sent to the user's email address. For on-prem installations, password resets are executed using a new default password, there is no email sent. If you do not know the default password, contact Darwin support.

Parameter:

• username - Username to reset password for.

Returns:

```
(True, None) or (False, <error-message>)
```

Example

```
In [8]: s.auth_reset_password('username')
Out[8]: (True, None)
```

DarwinSdk.auth_delete_user(username)

Remove/Unregister a user. This can only be performed by an admin account.

Parameter:

• username - Username of the user to be deleted.

Returns:



```
(True, <deleted-user-id>) or (False, <error-message>)
```

You must be logged in with your api_key and password to delete a user.

Example

```
In [8]: s.lookup_username('testuser2')
Out[8]:
(True,
 [{'client_api_key': 'RsJ74ZS5AvwznbHh0AfVSgrch...Eo3zFA',
   'created_at': '2018-01-03T12:54:30.653478',
   'parent_id': '2516b462-df85-11e7-bdd1-e37424c63ea4',
   'tier': 0,
   'username': 'testuser2'}])
In [9]: s.auth_delete_user('testuser2')
Out[9]: (True, '87d721fc-f0b7-11e7-b58d-a3441423b160')
In [10]: s.auth_delete_user('testuser2')
Out[10]:
(False,
'404: NOT FOUND - {\n
                          "message": "Username not found. \
You have requested this URI [/v1/auth/register/user/testuser2] \
but did you mean /v1/auth/register/user/ <username> or /v1/auth/register/user \
 or /v1/auth/register ?"\n}\n')
```

Job status methods

DarwinSdk.lookup_job_status(age=None, status=None)

Get status information for all jobs belonging to the current user or service.

Parameters:

- age (optional) Filter jobs that are less than X units old, for example 3w, 2d, or 1h.
- Optional parameters:
 - status If not specified, returns all jobs.
 - running (Note that only 2 jobs can be running concurrently.)
 - requested
 - complete
 - failed

Returns:

```
(True, <list-of-jobs>) or (False, <error-message>)
```



```
In [6]: s.lookup_job_status(status='Complete')
Out[6]:
(True,
    [{'artifact_names': None,
        'dataset_names': ['cancer-train'],
        'endtime': '2018-02-01T10:53:50.451598',
        'generations': 0,
        'job_name': 'eeef500d629e4a2185eb8af6e18a83b4',
        'job_type': 'TrainModel',
        'loss': 2.0,
        'model_name': 'cancer-model',
        'percent_complete': 100,
        'starttime': '2018-02-01T10:52:42.280929',
        'status': 'Complete'}])
```

DarwinSdk.lookup_job_status_name(job_name)

Get job status information for a job by its name.

Parameters:

• *job_name* - The name of the job you want status on

Returns:

```
(True, <job-info>) or (False, <error-message>)
```

```
In [19]: s.lookup_job_status_name('eeef500d629e4a2185eb8af6e18a83b4')
Out [19]:
(True,
 { 'artifact_names': None,
  'dataset_names': ['cancer-train'],
  'endtime': None,
  'generations': 0,
  'job_error': "MultipleDateColumns: multiple date columns \
    - ['Date' 'PeakMonth' 'PeakQuarter']",
  'job_type': 'TrainModel',
  'loss': None,
  'model_name': 'cancer-model',
  'percent_complete': 0,
  'starttime': '2018-02-01T10:52:42.280929',
  'status': 'Running'})
In [20]: s.lookup_job_status('Running')
```



DarwinSdk.delete_job(job_name)

Delete a job.

Parameter:

• job_name - The name of the job you want to delete

Returns:

```
(True, None) or (False, <error-message>)
```

Example

```
In [17]: s.lookup_job_status_name('7df54dfddfa046d581522f7540e3256c')
Out [17]:
(True,
 {'artifact_names': ['7a245119ca3b42efadc27006e75a225d'],
  'dataset_names': ['market-train'],
  'endtime': '2018-03-06T14:23:59.975793',
  'generations': None,
  'job_error': '',
  'job_type': 'AnalyzeData',
  'loss': None,
  'model_name': None,
  'percent_complete': 100,
  'starttime': '2018-03-06T14:23:57.18095',
  'status': 'Complete'})
In [18]: s.delete_job('7df54dfddfa046d581522f7540e3256c')
Out[18]: (True, None)
In [19]: s.lookup_job_status_name('7df54dfddfa046d581522f7540e3256c')
Out[19]: (False, '404: NOT FOUND - {\n "message": "Job name not found"\n}\n')
```

DarwinSdk.stop_job(job_name)

Stop a job.

Parameter:

• *job_name* - The name of the job.

Returns:

```
(True, 'Job is scheduled to stop') or (False, <error-message>)
```



```
In [21]: s.stop_job('34787793a48b42b48a319bbbf68f13ea')
Out[21]: (True, 'Job is scheduled to stop')
```

Lookup methods

DarwinSdk.lookup_artifact(type=None)

Get a list of artifacts belonging to the current user or service.

Parameter:

• type - (optional) specifies the type of artifact. Values can be 'Model', 'Dataset', 'Run'.

Returns:

```
(True, <artifact-list>) or (False, <error-message>)
```

Example:

```
In [30]: s.lookup_artifact('Run')
http://localhost:5000/v1/lookup/artifact
Out[30]:
(True,
   [{'created_at': '2018-02-01T11:09:55.731040',
    'id': 'b9a9205a-0772-11e8-a003-3b1c8766dad0',
    'mbytes': 0.0,
    'name': '8a63e21030d1483abb0f892963c1728f',
    'type': 'Run'},
   {'created_at': '2018-02-01T11:11:17.560360',
    'id': 'ea6f3f80-0772-11e8-9abe-77bc32e350c5',
    'mbytes': 0.0,
    'name': 'artifact-1',
    'type': 'Run'}]
```

DarwinSdk.lookup_artifact_name(artifact_name)

Get information for an artifact specified by its name.

Parameter:

• artifact - specifies an artifact by its name

Returns:

```
(True, <job-info>) or (False, <error-message>)
```



DarwinSdk.lookup_limits()

Get a client's metadata. A client is the current user or service in context.

Parameters: None

Returns:

```
(True, <client-info>) or (False, <error-message>)
```

Example

DarwinSdk.lookup_dataset()

Get the dataset(s) metadata for a user. The user is the current user or service in the current context. This is useful for listing all created datasets.

Parameters: None

Returns:

```
(True, <list-of-dataset-info>) or (False, <error-message>)
```

```
In [4]: s.lookup_dataset()
Out[4]:
(True,
  [{'categorical': None,
```



```
'imbalanced': None,
'mbytes': 0.02019977569580078,
'minimum_recommended_train_time': "string"
'name': 'unittest-cancer-dataset2',
'sequential': None,
'updated_at': '2018-01-31T15:37:28.310994'},
{'categorical': None,
'imbalanced': None,
'mbytes': 0.02019977569580078,
'minimum_recommended_train_time': "string"
'name': 'cancer-train',
'sequential': None,
'updated_at': '2018-02-01T10:52:06.076279'}])
```

DarwinSdk.lookup_dataset_name(dataset_name)

Get a specific dataset's metadata.

Parameters:

• dataset_name - The name of the dataset. The name of a dataset is established in the **up-load_dataset()** method.

Returns:

```
(True, <dataset-info>) or (False, <error-message>)
```

Example

DarwinSdk.lookup_model()

Get the model(s) metadata for a user. The user is the current user or service in the current context. This is useful for listing all models.

Parameters: None



Returns:

```
(True, <list-of-model-info>) or (False, <error-message>)
```

Example

DarwinSdk.lookup_model_name(model_name)

Get a specific model's metadata. The name of a model is established in the create_model() method.

Parameters:

• model_name - The name of the model

Returns:

```
(True, <model-info>) or (False, <error-message>)
```

```
In [40]: s.lookup_model_name('cancer-model')
Out[40]:
(True,
    [{'generations': 0,
        'loss': 2.0,
        'parameters': {'target': 'Diagnosis'},
        'trained_on': ['cancer-train'],
        'updated_at': '2018-02-01T10:53:50.443166',
        'description': {"best_genome": "DeepNet(\n (10): LSTM(20, 18, num_layers=2)\n (11): Linear(in_features=18, out_features=1, bias=True)\n)", "recurrent": True}
}]
)
```



DarwinSdk.lookup_tier()

Get metadata for all tiers. A tier specifies certain usage limits such as number of models and datasets.

Parameters: None

Returns:

```
(True, st-of-tier-info>) or (False, <error-message>)
```

Example

DarwinSdk.lookup_tier_num(tier_num)

Get a specific tier's metadata. A tier specifies certain usage limits such as the *number of models* or *datasets*.

Parameters:

• *tier_num* - The number of the tier

Returns:

```
(True, <tier-info>) or (False, <error-message>)
```



DarwinSdk.lookup_user()

Returns information for users that were created with the current api_key.

Note: Each customer site is assigned a *unique api_key*. All users from that site have the same api_key.

Parameters: None

Returns:

```
(True, <list-of-user-info>) or (False, <error-message>)
```

Example

```
In [25]: s.lookup_user()
Out[25]:
(True,
 [{'client_api_key': 'RsJ74ZS5AvwznbHh0AfVSgrchhS9...3zFA',
   'created_at': '2018-01-03T12:54:30.653478',
   'parent_id': '2516b462-df85-11e7-bdd1-e37424c63ea4',
   'tier': 0,
   'username': 'testuser2'},
  {'client_api_key': 'RsJ74ZS5AvwznbHh0AfVSgrchhS9...3zFA',
   'created_at': '2018-01-03T13:14:36.188371',
   'parent_id': '2516b462-df85-11e7-bdd1-e37424c63ea4',
   'tier': 0,
   'username': 'testuser5'},
  {'client_api_key': 'RsJ74ZS5AvwznbHh0AfVSgrchhS9...3zFA',
   'created_at': '2018-01-03T13:21:21.099148',
   'parent_id': '2516b462-df85-11e7-bdd1-e37424c63ea4',
   'tier': 0,
   'username': 'testuser6'}])
```

DarwinSdk.lookup_username(username)

Returns information for a user.

Notes:

- The user in question should have been created using the current api_key.
- Each customer site is assigned a unique api_key. All users from that site have the same api_key.

Parameters: None

Returns:

```
(True, <user-info>) or (False, <error-message>)
```

```
In [26]: s.lookup_username('testuser2')
Out[26]:
```



```
(True,
    [{'client_api_key': 'RsJ74ZS5AvwznbHh0AfVSgrchhS9...3zFA',
    'created_at': '2018-01-03T12:54:30.653478',
    'parent_id': '2516b462-df85-11e7-bdd1-e37424c63ea4',
    'tier': 0,
    'username': 'testuser2'}])
```

DarwinSdk.display_population(model_name)

Get a specific model's population data. The name of the model is established in the **create_model()** method.

Parameters:

• model_name - The name of the model

Returns:

```
(True, <population-info>) or (False, <error-message>)
```

```
In [40]: s.display_population('cancer-model')
Out [40]:
        (True,
          "population": {
            "model_types": {
              "DeepNeuralNetwork": {
                 "model_description": [
                     "layer 1": {
                       "type": "LinearLayer",
                       "parameters": {
                         "activation": "leakyrelu",
                         "numunits": 221
                    }
                   },
                     "layer 2": {
                       "type": "LinearLayer",
                       "parameters": {
                         "activation": "relu",
                         "numunits": 2
```



Datasets and artifact methods

DarwinSdk.upload_dataset(dataset, dataset_name=None)

Upload a dataset.

Note: Supported file formats are .csv and .h5.

Note: The maximum size that can be uploaded is 10GB. Files larger than ~2GB can be processed by analyze_data() only. Model creation might not be successful for files larger than ~2GB until Big Data is fully supported. Analyze_data() is the only method that supports Big Data.

Parameters:

- dataset- Path to dataset
- dataset_name Name to be given to dataset, or defaults to filename

Returns:

```
(True, {dataset_name: <name-given-to-dataset>}) or (False, <error-message>)
```



DarwinSdk.download_dataset(dataset_name)

Download a dataset artifact given its name.

Parameters:

• dataset_name - Name of the dataset to be downloaded.

Returns:

```
(True, <path-to-file>) or (True, <python-list>) or (False, <error-message>)
```

Example

DarwinSdk.delete_dataset(dataset_name)

Delete the named dataset.

Parameters:

• dataset name - Name of the dataset to be deleted.

Returns:

```
(True, None) or (False, <error-message>)
```

Example

```
In [6]: s.delete_dataset('unittest-cancer-dataset')
Out[6]:
(True, None)
```

DarwinSdk.download_model(model_name)

Download a supervised model given its name.

Parameters:

- model_name Name of the model to be downloaded.
- *path* (optional) Relative or absolute path of the directory to download the model to. This directory must already exist prior to model download. If the path is not specified, the current directory is used. There are two files associated with a model: *'model'* and *'data_profiler'*.



- model_type (optional) Model type of the model to be downloaded. Possible values include the following: DeepNeuralNetwork, RandomForest, GradientBoosted.
- *model_format* (optional) Format in which the model is to be downloaded. Possible values include: *json, onnx*.

Returns:

```
(True, None) or (False, <error-message>)
```

Example

```
In [6]: s.download_model('my-model-name', path='Users/auser/Downloads/mymodel')
Out[6]:
(True, None)
% ls -l ~/Downloads/mymodel
total 272
-rw-r--r- 1 auser staff 58609 Oct 10 15:55 data_profiler
-rw-r--r- 1 auser staff 75507 Oct 10 15:55 model
```

DarwinSdk.download_artifact(artifact_name, artifact_path=None)

Download artifact given its name. The methods that return artifacts are:

- analyze_data()
- analyze_model()
- analyze_predictions()
- run_model()

Note: The artifact for *analyze_model()* is a pandas Series. The artifact displays a two-column series where the name of the feature is in the first column and the second column is a number between 0 and 1 indicating how much that feature influenced the model's predictions over the entire dataset that the model was trained on.

Note: The artifact for *analyze_predictions* is a pandas DataFrame. The artifact has one column for each feature that indicates how much that feature influenced the model's prediction, plus additional columns for the average model prediction ("base_value"), and the model prediction for each row ("predicted_value" for regression or "predicted-class" and "predicted_probability" for classification).

Parameters:

- artifact_name Name of the artifact to download.
- artifact_path: (optional) Relative path of the directory to download the artifact to (only applicable for the artifacts where a temporary file is created). This directory must already exist prior to artifact download.

Returns:

```
(True, <path-to-file>) or (True, <python-list>) or (False, <error-message>)
```

Example run_model() or prediction artifact



```
In [16]: s.download_artifact('5da17d64be9c4441899316edb9afd403')
Out[16]:
(True,
         Diagnosis prob_BENIGN prob_MALIGNANT
      BENIGN 0.999400 6.002134e-04
1
      BENIGN
               1.000000 3.600000e-09
2
      BENIGN
               0.999999 8.689000e-07
             1.000000 2.500000e-09
3
      BENIGN
4 MALIGNANT
                          9.958413e-01
               0.004159
             0.002674 9.973264e-01
5 MALIGNANT
92 MALIGNANT
              0.002499
                         9.975013e-01
93
    BENIGN
                          5.250000e-08
               1.000000
               1.000000
                          3.100000e-08
94
    BENIGN
                         9.866350e-05
95
    BENIGN
                0.999901
96
    BENIGN
               1.000000
                         9.230000e-08
                0.003884 9.961160e-01
97 MALIGNANT
                          9.972232e-01
98 MALIGNANT
                0.002777
99 MALIGNANT
                         9.963139e-01
                0.003686
[100 rows x 3 columns])
```

Example analyze_data() artifact

In [19]: s.download_artifact('9233	38b7512f4770k	o239e1b5	3406cfa	6 ')		
Out[19]:						
(Tru	e, col	_name co.	l_type	drop i	s_cat	max	\
0	Code	int64	True	False	8233704		
1	Clump Thickness	categorical	False	True	None		
2	Uniformity of Cell Size	categorical	False	True	None		
3	Uniformity of Cell Shape	categorical	False	True	None		
4	Marginal Adhesion	categorical	False	True	None		
5	Single Epithelial Cell Size	categorical	False	True	None		
6	Bare Nuclei	LongType	False	False	10		
7	Bland Chromatin	categorical	False	True	None		
8	Normal Nucleoli	categorical	False	True	None		
9	Mitoses	categorical	False	True	None		
10	Diagnosis	categorical	False	True	None		
	mean		min mi	ssing n	um_uniques	\	
0	1044171.0667779633	61	1634	0.0	559		
1	4.555926544240401	1	None	0.0	10		
2	3.2153589315525877	1	None	0.0	10		
3	3.287145242070117	1	None	0.0	10		
4	2.8597662771285477	1	None	0.0	10		
5	3.290484140233723	1	None	0.0	10		



6	-2.30969	249670820768E17	-922337	2036	854	775	808			0.0			1	.1
7	3.5	208681135225377				No	one			0.0			1	. 0
8	2	.96661101836394				No	one			0.0			1	. 0
9	1.6	076794657762938				No	one			0.0				9
10		None				No	one			0.0				2
	scalable		stddev								u:	niq	ues	
0	True	414096.368	7689267									No	one	
1	False	2.88748784	4960718	[9,	1,	5,	2,	6,	3,	10,	7,	4,	8]	
2	False	3.04460120	2894244	[9,	1,	5,	2,	6,	3,	10,	7,	4,	8]	
3	False	2.971045056	2657416	[9,	1,	5,	2,	6,	3,	10,	7,	4,	8]	
4	False	2.87365509	2520189	[9,	1,	5,	2,	6,	3,	10,	7,	4,	8]	
5	False	2.275158768	9827613	[9,	1,	5,	2,	6,	3,	10,	7,	4,	8]	
6	True	1.4423736395283	3229E18									No	one	
7	False	2.36950002	0847775	[9,	1,	5,	2,	6,	3,	10,	7,	4,	8]	
8	False	3.084466482	0475916	[9,	1,	5,	2,	6,	3,	10,	7,	4,	8]	
9	False	1.734368638	0557295		[1,	5,	2,	6,	3,	10,	7,	4,	8]	
10	False		None			['BEI	NIGI	Ν',	'MA	LIG	NAN'	Γ'])

Example analyze_model() or prediction artifact

```
In [5]: s.download_artifact('6e4861de29424cb7ad09e467d1869c17',\
   'path_to_download_dir/')
 Out[5]:
 True RM
                     0.216088
CRIM
            0.141956
LSTAT
             0.134069
             0.104101
DIS
              0.089905
PTRATIO
              0.078864
AGE
              0.074132
NOX
              0.067823
В
              0.045741
TAX
INDUS
              0.023659
ZN
              0.011041
RAD = 4.0
             0.009464
RAD = 5.0
           0.001577
RAD = 6.0
             0.001577
RAD = 24.0
             0.000000
RAD = 3.0
            0.000000
RAD = 7.0
              0.000000
CHAS = 1.0
              0.000000
RAD = 8.0
              0.000000
RAD = 2.0
              0.000000
dtype: float64
```

Example analyze_predictions() artifact



```
In [8]: (code, fis) = s.download_artifact('34b461c7a52a48318e982068f87e6562',\
 'path_to_download_dir/')
In [9]: fis.head()
Out[9]: ##Sample return for regression, has predicted_value column
        AGE
                    B CHAS = 1.0
                                      CRIM
                                                  DIS
                                                           INDUS
                                                                     LSTAT \
0.000000 \quad 0.000000 \quad 0.000000 \quad -0.664664 \quad -0.923219 \quad -0.720941 \quad 2.328635
1 -1.220243 -0.648893
                        0.000000 0.000000 1.187539 -0.630767 3.506132
2 - 0.456561 - 0.226880 - 0.424802 0.000000 - 0.077616 - 0.333270 - 0.292705
3 -0.195096 0.352712
                        0.000000 -1.867664 -0.152037 0.273082 -3.583178
4 \quad 0.632119 \quad 0.079678 \quad 0.000000 \quad 0.076080 \quad -0.488128 \quad -0.016690 \quad -0.102031
        NOX
            PTRATIO RAD = 2.0
                                                    RAD = 4.0 RAD = 5.0 \setminus
0 -0.342404 0.224360
                             0.0
                                                    -0.641678 -0.570788
1 -0.556636 -2.168356
                             0.0
                                                     0.000000 - 0.741561
                                        . . .
2 0.000000 1.458677
                              0.0
                                                     0.000000 -0.340486
3 - 0.945060 - 1.068743
                              0.0
                                                     0.000000 0.217991
                                        . . .
4 0.309544 0.298940
                                                     0.000000 - 0.047708
                              0.0
                                        . . .
   RAD = 6.0 \quad RAD = 7.0 \quad RAD = 8.0
                                          RM
                                                                ZN base_value \
                                                    TAX
0
                   0.0
                              0.0 -1.835851 -0.563795 -0.600155
         0.0
                                                                      21.63455
1
         0.0
                    0.0
                                0.0 -1.016655 -0.699813 -0.727181
                                                                      21.63455
2
         0.0
                    0.0
                               0.0 -1.137559 0.000000 -0.310209
                                                                      21.63455
3
         0.0
                    0.0
                              0.0 -1.220045 0.156790 0.256763
                                                                     21.63455
4
         0.0
                    0.0
                               0.0 -0.999328 -0.149627 -0.045493
                                                                      21.63455
  predicted_value
0
        24.620939
         26.128595
1
2
         24.200972
3
        11.255393
        21.982929
[5 rows x 22 columns]
Out[9]: ##Sample return for classification, returns predicted_class as well
   petal length (cm) petal width (cm) sepal length (cm) sepal width (cm) \
0
            0.217699
                             0.424209
                                                  0.026237
                                                                    0.005834
1
            0.292612
                               0.315358
                                                  0.019236
                                                                   -0.014442
2
            0.325615
                               0.329229
                                                  0.003208
                                                                     0.016954
3
            0.232265
                               0.410938
                                                  0.043014
                                                                     0.004154
4
            0.317190
                               0.339065
                                                   0.015227
                                                                     0.003523
   base_value predicted_value predicted_class
```



0	0.309628	0.983607	virginica
1	0.365378	0.978142	versicolor
2	0.324994	1.000000	setosa
3	0.309628	1.000000	virginica
4	0.324994	1.000000	setosa

DarwinSdk.delete_artifact(artifact_name)

Delete the artifact given its name.

Parameters:

• artifact_name - Name of the artifact to be deleted.

Returns:

```
(True, None) or (False, <error-message>)
```

Example

```
In [8]: s.delete_artifact('6c482eac9f894cdb9b0e1e487e41730a')
Out[8]:
(True, None)
```

Data Analysis and Data Cleaning methods

DarwinSdk.analyze_data(dataset_name, **kwargs)

Analyze the dataset given its *name*. Basic statistics about the data are returned. This method supports Big Data (greater than 2GB) although *upload_dataset()* is artificially limited to 10GB for version 1.4.

Note:Please contact us if you have data greater than 10GB. We would like to see a sampling of the large datasets that you'd like to see supported.

Parameters:

dataset_name - The name of the dataset to be analyzed.

**kwargs - variable number of keyword arguments, described below:

- job_name (optional) If not specified, a uuid will be created as the job_name.
- artifact_name: (optional) If not specified, a unid will be created as the artifact_name.
- max_unique_values: Expected input/type: integer. Default value of 15. Threshold for automatic pruning of categorical columns prior to one hot encoding based on the number of unique values.
 Note: If a categorical column contains at least max_unique_values, it is dropped during preprocessing prior to one hot encoding.

Returns:

```
(True, {"job_name": <string>, "artifact_name": <string>}) or (False, <error-message>)
```



Statistics included in the artifact:

- col_name name of the column (any periods ('.') in the column name will be replaced by underscores
 ('_'))
- *col_type* type of column
- *drop* returns True if column is dropped for modeling. Also returns True if the number of unique values is greater than the number defined in *max_unique_values* (default of 15) or if it has more than 80% missing values or has a standard deviation of 0.
- *is_cat* returns True for categorical otherwise returns False
- max column maximum
- mean column mean
- *min* column minimum
- missing percentage of missing values
- num_uniques number of unique values if the distinct count is less than the number defined in max_unique_values (default of 15), otherwise the value is the approximate number of unique values.
- scalable returns True if column is scalable
- stddev column standard deviation
- *uniques* actual unique values if there are less than the number defined in *max_unique_values* (default of 15). Otherwise, nothing is returned, see *num_uniques* for the approximate number of unique values.

```
In [6]: s.analyze_data('boston')
Out[6]:
(True,
 { 'artifact_name': 'db968d77d2c4444ab731777d01e5e0c0',
  'job_name': '8c12f0df4c39485f9a488fa63196e00c'})
In [8]: s.download_artifact('db968d77d2c4444ab731777d01e5e0c0')
Out[8]:
(True,
                        col_name
                                   col_type
                                                drop is_cat
                                                                     max
 0
                             PID
                                   StringType
                                                True False 2205663001
 1
                          ST NUM
                                   StringType
                                                True False
                                                                     999
 2
                         ST NAME
                                   StringType
                                                True False
                                                                  ZELLER
 3
                     ST_NAME_SUF
                                   StringType
                                                True False
                                                                      ΧT
 4
                         ZIPCODE
                                   StringType
                                                True False
                                                                  02467
                                        int64
                                                                23095700
 5
                  Assessed Value
                                                True False
 6
                       Lot Area
                                        int64
                                                True False
                                                                  107158
 7
                     Gross_Area
                                        int64
                                                True False
                                                                   23335
 8
                     Living_Area
                                        int64
                                               True False
                                                                   21711
 9
                  Owner_Occupied categorical False
                                                     True
                                                                    None
 10
                      Year_Built
                                        int64
                                               True False
                                                                    2016
 11
               Number_of_Floors
                                      float64 False False
                                                                     5.0
 12
           Total_Number_of_Rooms
                                        int64
                                                                      2.7
                                               True False
 13
              Number_of_Bedrooms categorical False
                                                      True
                                                                    None
```



14	Number_of_Full				False	True		lone
15	Number_of_Half		_		False	True		lone
16	Number_of_Ki	tchens	-		False	True	Ŋ	lone
17		Has_AC	-	,	False	True	Ŋ	lone
18	Number_of_Fire		cate		False	True	N	lone
19	Year_Since_Remodel_or			int64	True	False		307
20	Year_Rem			ngType	True	False	Unremode	
21	Structur		_		False	True		lone
22	Building	_Style	Stri	ngType	True	False	Victor	rian
23		f_Type	-		False	True	N	lone
24	Exterior_	Finish	cate	gorical	False	True	N	lone
25	Main_Bathroom	_Style	cate	gorical	False	True	N	lone
26	Main_Kitchen			,	False	True	N	lone
27	Heatin		cate	gorical	False	True	N	lone
28	Exterior_Con		-	,	False	True	N	lone
29	Overall_Con		cate	gorical	False	True	N	lone
30	Interior_Con	dition	cate	gorical	False	True	N	lone
31	Interior_	Finish	cate	gorical	False	True	N	lone
32		View	cate	gorical	False	True	N	lone
			!] .].] .	\
0	mean	0100001	min				scalable	\
0		0100021		0.000000		28578	True	
1	122.09705524787249		.005R	0.010223		1922	True	
2	None	ABBOTS		0.000000		2246	True	
3	None	0.0	ST	0.003015		21	True	
4	None		108_	0.000000		28	True	
5	534716.6815977456	10	1300	0.000000		7737	True	
6	5116.273150271971		375	0.000000		8342	True	
7	2931.1126220591127		510	0.000000		4472	True	
8	1752.7717084999017		332	0.000000		3169	True	
9	0.8408480241169146		None	0.000000		2	False	
10	1926.970935185792		1710	0.000000		225	True	
11	1.8748115866046269		1.0	0.000000		9	True	
12	7.233632610262796		2 Nana	0.000000		26	True	
13	3.3851169801428664		None	0.000000		12	False	
14	1.4273543482534898		None	0.000000		10	False	
15	0.5716953928828888		None	0.000000		7	False	
16	1.0287043711907726		None	0.000000		4	False	
17	0.18733206632151517		None	0.000000		2	False	
18	0.590995478078511		None	0.000000		13	False	
1.0	60.88419948882627		1	0.000000		190	True	
19			1000			0.0		
20	2000.3376960831488		1890	0.000000		82	True	
	2000.3376960831488 None		None	0.000000		82 5	False	
20								



e	False	13	0.000000	None	None	24
e	False	4	0.000000	None	None	25
Э	False	4	0.00000	None	None	26
e	False	6	0.00000	None	None	27
e	False	5	0.00000	None	None	28
e	False	5	0.00000	None	None	29
Э	False	5	0.000000	None	None	30
Э	False	3	0.000000	None	None	31
e	False	5	0.00000	None	None	32
uniques	ι				stddev	
None					None	0
None					294.1511958893473	1
None					None	2
None					None	3
None					None	4
None					634750.7826113638	5
None					3218.286557124007	6
None					1069.3847598444354	7
None					758.9874732061347	8
[0, 1]					0.3658237412175791	9
None					34.9170355483078	10
None					0.5737101635770085	11
None					1.8082562295656077	12
, 11, 8]	10, 7, 4,	5, 2, 6, 3, 1	2, 9, 1, 5	[1	1.0095185504254367	13
7, 4, 8]	, 6, 3, 7,	9, 1, 5, 2,	[12,		0.6850264359951297	14
6, 3, 4]	, 5, 2, 6,	[0, 1,			0.5645602408681473	15
1, 2, 3]	[0, 1,				0.17162236936210065	16
[0, 1]					0.3901842537872663	17
, 11, 8]	10, 7, 4,	5, 2, 6, 3, 1	2, 9, 1, 5	[0, 1	0.8584446055814273	18
None					43.323487380439225	19
None					13.578956800881818	20
'Bric	known', 'E	'Frame ', ' Unk	1', 'Wood/	['Residentia	None	21
None					None	22
sard'	r', 'Mansa	lat', 'Other	mbrel', 'F	['Shed', 'Ga	None	23
Shak	', 'Wood S	ne/Clapboard'	rd', 'Fram	['Cement Boa	None	24
, 'Mo	odeling',	ry', 'No Remo	n', 'Luxur	['Semi-Mode	None	25
, 'Mo	odeling',	ry', 'No Remo	n', 'Luxur	['Semi-Moder	None	26
, 'Sp	t Water',	'None', 'Hot	'Other',	['Electric',	None	27
'Fair']	verage',	cellent', 'Av	Good', 'Exc	['Poor', '(None	28
'Fair']	verage',	cellent', 'Av	Good', 'Exc	['Poor', '(None	29
'Fair']	verage',	cellent', 'Av	Good', 'Exc	['Poor', '(None	30
andard']	, 'Substar	e', 'Normal',	'Elaborate		None	31
'Fair'])	verage',	cellent', 'Av	Good', 'Exc	['Poor', '(None	32



DarwinSdk.clean_data(dataset_name, **kwargs)

Clean the dataset given its name. The output is the cleaned dataset which is scaled and one-hot-encoded based on parameters in <code>analyze_data()</code>. Use <code>download_dataset()</code> to retrieve the cleaned dataset. <code>clean_data()</code> needs to be performed prior to creating a model and again before running a model. When you run clean_data() before creating a model, you must specify a dataset_name and a target. When you run clean_data() before running a model, you must specify a dataset_name and a model_name. <code>clean_data()</code> can also be used for visualizing what Darwin would do with the dataset or for when you want to use the cleaned data outside of Darwin.

Parameters:

- dataset name The name of the dataset to be analyzed.
- **kwargs variable number of keyword arguments, described below:
 - job_name : (optional) If not specified, a uuid will be created as the job_name.
 - artifact_name: (optional) If not specified, a unid will be created as the artifact_name.
 - *model_name*: (Mandatory for running a model) Specify the model name when you clean data before running a model.
 - target: (Mandatory for Supervised Model Building) String denoting target prediction column in input data.
 - impute: String alias that indicates how to fill in missing values in input data.

ALIAS	DESCRIPTION	COMPLEXITY
'ffill'	(Default) Forward Fill: Propagate values forward from one example	Linear
	into the missing cell of the next example. Might be useful for	Fast
	timeseries data, but also applicable for both numerical and	
	categorical data.	
'bfill'	Backward Fill: Propagate values backward from one example into	Linear
	the missing cell of the previous example. Might be useful for	Fast
	timeseries data, but also applicable for both numerical and	
	categorical data.	
'mean'	Mean Fill: Computes the mean value of all non-missing examples	Linear
	in a column to fill in missing examples. The result may or might	Fast
	not be interpretable in terms of the input space for categorical	
	variables.	

- max_int_uniques: Expected input/type: integer. Threshold for automatic encoding of categorical variables. If a column contains less than max_int_uniques unique values, it is treated as categorical and one hot encoded during preprocessing. Note: If the target has more numeric values than the max_int_uniques set point, the problem is treated as a regression and will use MSE.
- max_unique_values: Expected input/type: integer. Threshold for automatic pruning of categorical columns prior to one hot encoding based on the number of unique values.

Note: If a categorical column contains at least *max_unique_values*, it is dropped during preprocessing prior to one hot encoding.



Modeling and analysis methods

DarwinSdk.create_model(dataset_names, **kwargs)

Create a model trained on the dataset identified by dataset_names. You must clean the data using *clean_data()* The name of a model is specified in a parameter in kwargs.

Note: If no name is specified, the model is named with a *uuid-like* name.

Parameters:

dataset_names - A single dataset name as a string or a list of dataset string names to be used for training. The maximum file size is 500 MB for unsupervised and NBM and 10 GB for supervised.

**kwargs - variable number of keyword arguments, described in parameters.

parameters -

- *model_name*: The string identifier of the model to be trained. If no name is specified, the model is named with a *uuid-like* name.
- job_name: If no name is specified, the job is named with a uuid-like name.
- *max_train_time* (supervised only): Sets the training time for the model in 'HH:MM' format. Default value is 00:01.
- *max_epochs* (unsupervised only): Expected input/type: *numeric*. Sets the training time for the model in epochs. Default value is 10.
- *recurrent*: Expected input/type: *True/False*. Enables recurrent connections to be evolved in the model. This option can be useful for timeseries or sequential data.

Note: This option is automatically enabled if a *datetime* column is detected in the input data. This can result in slower model evolution.

• impute: String alias that indicates how to fill in missing values in input data.



ALIAS	DESCRIPTION	COMPLEXITY
'ffill'	(Default) Forward Fill: Propagate values forward from one example	Linear
	into the missing cell of the next example. Might be useful for	Fast
	timeseries data, but also applicable for both numerical and	
	categorical data.	
'bfill'	Backward Fill: Propagate values backward from one example into	Linear
	the missing cell of the previous example. Might be useful for	Fast
	timeseries data, but also applicable for both numerical and	
	categorical data.	
'mean'	Mean Fill: Computes the mean value of all non-missing examples	Linear
	in a column to fill in missing examples. The result may or might	Fast
	not be interpretable in terms of the input space for categorical	
	variables.	

- *anomaly*: Setting this parameter to **True** indicates that an isolation forest should be built for anomaly detection. If set to **True**, clustering will automatically be interpreted as **False**.
- *clustering* (unsupervised only): Enables clustering for unsupervised problems. If False, detects outliers.
- *n_clusters* (unsupervised only): Expected input/type: *integer*. Specifies the number of clusters. **Note**: If this value is not provided, the number of clusters will be heuristically determined.
- *anomaly_prior* (unsupervised only): Expected input/type: *between* [0,1]. Significance level at which a point is defined as anomalous. This is only used for unsupervised problems if *clustering* is disabled.
- loss_fn_name: Specify the loss function. Possible values include: "CrossEntropy", "MSE", "BCE", "L1", "NLL", "BCEWithLogits", "SmoothL1". "CrossEntropy" can be used for classification data, while all others can be used for regression data. The default value is CrossEntropy if this field is left empty.
- *fitness_fn_name*: Specify the fitness function. This represents the name of the fitness function used for evolution of the model population during training. Possible values include: "Accuracy", "F1", "R2", "MSE". "F1" is the default for classification and "R2" is the default for regression problems. "Accuracy" and "F1" are for classification only. "R2" and "MSE" are for regression only.
- *lead_time_days* (*nbm* only): Expected input/type: *integer*. Default value is 60. The number of days prior to failure when the behavior starts trending toward either abnormal behavior or failure.
- *nbm_window_size* (*nbm* only): Expected input/type: *integer*. Default value is 256. The number of sample points to consider for each failure detection.
- *nbm* (*nbm* only): Expected input/type: *True/False*. Default value is False. Set value to True for a normal behavioral model (NBM).
- failure_dates (nbm only): Expected input/type: string. List of failure dates to use for the calculation. Currently, only a list of one date can be used in the query. Example date format: "07/01/2015"
- recovery_dates (nbm only): Expected input/type: string. List of recovery dates to use for the



calculation. Currently, only a list of one date can be used in the query. Example date format: "11/01/2015"

Returns:

```
(True, {'job_id': <uuid1>, model_name: <model_name>}) or (False, <error-message>)
```

Example

DarwinSdk.delete_model(model_name)

Delete a model named by model_name.

Parameters:

• model_name - Name of the model to be deleted.

Returns:

```
(True, None) or (False, <error-message>)
```

Example

```
In [5]: s.delete_model('unittest-cancer-model')
Out[5]: (True, None)
```

DarwinSdk.resume_training_model(model_name, dataset_names, **kwargs)

Resume training for a model on the dataset(s) identified by dataset_names.

Parameters:

- model_name Name of the model to be trained.
- dataset_name- Name of dataset(s) used for training.
- **kwargs variable number of keyword arguments, described below:.
 - job_name If not specified, a unid is created as the job_name.
 - max_train_time If not specified, the *default* is used.

Returns:

```
(True, {"job_id""<uuid>", "model_name": "<model_name>"}) or (False, <error-message>)
```



Example

```
In [8]: s.resume_training_model('unittest-cancer-model', 'unittest-cancer-\
dataset', max_train_time="00:01")
Out[8]:
(True, {"job_id": "4e59ffc425e047e1a3b872f1e7396976", "model_name": "unittest-\
cancer-model"})
```

DarwinSdk.analyze_model(model_name, job_name=None, artifact_name=None)

Analyze the universal feature importances for a particular model given the model name.

Parameters:

- model_name The name of the model to be analyzed.
- job_name (optional) If not specified, a uuid is created as the job_name.
- artifact_name (optional) If not specified, a uuid is created as the artifact_name.
- *category_name* (optional) The name of the class for supervised or cluster for unsupervised to get feature importance for. If this is not specified, the feature importance will be over all classes/clusters.
- model_type: (optional) Model type from the population. Possible values include: *DeepNeuralNetwork*, *RandomForest*. *GradientBoosted*.

Returns:

```
(True, {"job_name": <string>, "artifact_name": <string>}) or (False, <error-message>)
```

Example

```
In [5]: s.analyze_model('unittest-cancer-model')
Out [5]:
(True, {'artifact_id': '71a8ae55f2934014b45c13a3975f419c', 'job_id': \
'4e59ffc425e047e1a3b872f1e7396976'})
```

${\bf DarwinSdk.analyze_predictions} (model_name,\ dataset_name,\ job_name=None,\ artifact_name=None)$

Analyze specific feature importances for a particular sample or samples given the model name and sample data. Analyze predictions cannot be used if you trained your model with a dataset that is larger than 500 MB.

Parameters:

- *dataset_name* The name of the dataset containing the data to analyze predictions for. This is a new dataset that was not used during training for which you want feature importance scores for each row of this dataset. This dataset has a limit of 500 rows. There is no limit for columns.
- model_name The name of the model to be analyzed.



- job_name (optional) If not specified, a uuid is created as the job_name.
- artifact_name (optional) If not specified, a unid is created as the artifact_name.
- *start_index* (optional) Index to start at in the dataset when analyzing model predictions. All numeric and datetime data types can be indexes. When specifying an index as a datetime, the preferred timestamp format is 2019-02-15 19:46:48.
- *end_index* (optional) Index to stop at in the dataset when analyzing model predictions. All numeric and datetime data types can be indexes. When specifying an index as a datetime, the preferred timestamp format is 2019-02-15 19:46:48.
- model_type: (optional) Model type from the population. Possible values include: DeepNeuralNetwork, RandomForest, GradientBoosted.

Returns:

```
(True, {"job_name": <string>, "artifact_name": <string>}) or (False, <error-message>)
```

Example

```
In [5]: s.analyze_predictions('model_name', 'dataset_name')
Out [5]:
(True, {'artifact_name': '71a8ae55f2934014b45c13a3975f419c', 'job_name': \
'4e59ffc425e047e1a3b872f1e7396976'})
```

DarwinSdk.run_model(dataset_name, model_name, job_name=None, artifact_name=None)

Run the model given its name and a dataset to use. Use upload_dataset() to upload a data set.

Parameters:

- dataset_name The name of a dataset to use for running the model.
- model_name The name of the model to run.
- *anomaly*: Setting this parameter to **True** indicates that an isolation forest should be built for anomaly detection. If set to **True**, clustering will automatically be interpreted as **False**.
- *supervised* (**Deprecated**: This argument exists only for backward compatibility.) (optional) A boolean (True/False) indicating whether the model is supervised or not, for example, set this to *False* for *unsupervised*.
- job_name (optional) If not specified, a uuid is created as the job_name.
- artifact_name (optional) If not specified, a unid is created as the artifact_name.
- *model_type* (optional) Model type of the model to be downloaded. Possible values include the following: *DeepNeuralNetwork*, *RandomForest*, *GradientBoosted*.

Returns:

```
(True, {"job_name": <string>, "artifact_name": <string>}) or (False, <error-message>)
```



```
[In [9]: s.run_model('unittest-cancer-testdataset', 'unittest-cancer-model')
Out [9]:
(True, {'artifact_id': '6c482eac9f894cdb9b0e1e487e41730a', 'job_id': \
'1696e03c8165404c8e05685ea68baa3c'})
```

Convenience methods

DarwinSdk.delete_all_datasets()

Deletes user datasets. This method deletes all datasets in the current user or service context.

Note: Use *lookup_dataset()* to view/verify the datasets for deletion.

Parameters: None

Returns:

```
(True, None) or (False, <error-message>)
```

DarwinSdk.delete_all_models()

Delete all models for a user. This method will delete all models in the current user's or service's context. **Note**: Use *lookup_model()* to review and verify that you want to delete all listed models.

Parameters: None

Returns:

```
(True, None) or (False, <error-message>)
```

DarwinSdk.delete_all_artifacts()

Delete all artifacts for a user. This method will delete all artifacts in the current user's or service's context. **Note**: Use *lookup_artifact()* to review and verify that you want to delete all listed artifacts.

Parameters: None

Returns:

```
(True, None) or (False, <error-message>)
```



DarwinSdk.wait_for_job(job_name, time_limit=600)

Synchronously wait for a job to complete, limited by time_limit that defaults to 600 seconds.

Parameters:

- job_name The id for the job
- time_limit (optional) defaults to 600 seconds

Returns:

```
(True, None) or (False, <error-message>)
```

Reference

- SDK modeling example
- Revision table

SDK modeling example

The following example shows the Darwin SDK performing a modeling process:

```
In [1]: from amb_sdk.sdk import DarwinSdk
In [2]: s = DarwinSdk()
In [3]: s.auth_login_user('your-username', 'your-password')
Out[3]:
(True,
 'Bearer eyJ0eXAi0iJK...A8sj4pAzX1FpMMscwY_rMJbnGo0YQ_4')
In [4]: s.upload_dataset('sets/cancer_train.csv', 'mydata')
Out[4]: (True, {'dataset_name': 'mydata'})
In [5]: s.clean_data('mydata', target='Diagnosis')
Out[5]:
(True,
 {'job_name': '801ee7e95dfd4380b7be76332ead5036',
  'artifact_name': '97fad4a3598f41068eadd84df26a6eaa'})
In [6]: s.wait_for_job('801ee7e95dfd4380b7be76332ead5036')
{'status': 'Complete', 'starttime': '2019-01-16T11:28:09.779535',\
 'endtime': '2019-01-16T11:28:12.613227', 'percent_complete': 100,\
  'job_type': 'CleanDataTiny', 'loss': None, 'generations': None,\
```



```
'dataset_names': ['mydata'], 'artifact_names': \
   ['97fad4a3598f41068eadd84df26a6eaa'], 'model_name': None,\
   'job_error': ''}
Out[6]: (True, 'Job completed')
In [7]: s.create_model(dataset_names='mydata', model_name='my-model')
Out[7]:
(True,
 {'job_name': '2bbf5dc050b6499a9e19e0c6173a2821',
  'job id': '2fa8953e-19b4-11e9-a52a-1b252aa286fd',
  'model_name': 'my-model'})
In [8]: s.wait_for_job('2bbf5dc050b6499a9e19e0c6173a2821')
{'status': 'Running', 'starttime': '2019-01-16T11:28:49.588621',\
 'endtime': None, 'percent_complete': 0, 'job_type': 'TrainModel',\
  'loss': 0.4303114712238312, 'generations': 2, 'dataset_names': ['mydata'], \
   'artifact_names': None, 'model_name': 'my-model', 'job_error': ''}
{'status': 'Running', 'starttime': '2019-01-16T11:28:49.588621',
 'endtime': None, 'percent_complete': 3, 'job_type': 'TrainModel',\
  'loss': 0.18398252129554749, 'generations': 2, 'dataset_names': ['mydata'], \
   'artifact_names': None, 'model_name': 'my-model', 'job_error': ''}
{'status': 'Running', 'starttime': '2019-01-16T11:28:49.588621',\
 'endtime': None, 'percent_complete': 8, 'job_type': 'TrainModel', \
  'loss': 0.41190358996391296, 'generations': 4, 'dataset_names': ['mydata'],\
   'artifact_names': None, 'model_name': 'my-model', 'job_error': ''}
{'status': 'Complete', 'starttime': '2019-01-16T11:28:49.588621',\
 'endtime': '2019-01-16T11:29:55.502275', 'percent_complete': 100,\
  'job_type': 'TrainModel', 'loss': 0.41190358996391296, 'generations': 4,\
   'dataset_names': ['mydata'], 'artifact_names': None, \
    'model_name': 'my-model', 'job_error': ''}
Out[8]: (True, 'Job completed')
In [9]: s.upload_dataset('sets/cancer_test.csv', 'mytestdata')
Out[9]: (True, {'dataset_name': 'mytestdata'})
In [10]: s.clean_data('mytestdata', model_name='my-model')
Out[10]:
(True,
 {'job_name': '625f2cc5c6e2437d808b158ad66bfefc',
 'artifact_name': '17bad7c5426c4166afa4fe70eb0ff8a1'})
In [11]: s.wait_for_job('625f2cc5c6e2437d808b158ad66bfefc')
{'status': 'Complete', 'starttime': '2019-01-16T11:34:11.567819',\
 'endtime': '2019-01-16T11:34:13.283641', 'percent_complete': 100,\
  'job_type': 'CleanDataTiny', 'loss': None, 'generations': None,\
```



```
'dataset_names': ['mytestdata'], 'artifact_names': ['17bad726...4fe70eb0ff8a1'], \
    'model_name': None, 'job_error': ''}
Out[11]: (True, 'Job completed')
In [12]: s.run_model('mytestdata', 'my-model')
Out[12]:
(True,
 {'job_name': '75d8bf61689346fda84b430f5fe1be58',
 'artifact_name': '20e87cda3ef24cd18f065ccaf87e8ca4'})
In [13]: s.wait_for_job('75d8bf61689346fda84b430f5fe1be58')
{'status': 'Complete', 'starttime': '2019-01-16T11:35:33.1109',\
 'endtime': '2019-01-16T11:35:34.891138', 'percent_complete': 100,\
  'job_type': 'RunModel', 'loss': 0.41190358996391296, 'generations': 4,\
   'dataset_names': ['mytestdata'], 'artifact_names': ['20e87cda3...065ccaf8ca4'],\
    'model_name': 'my-model', 'job_error': ''}
Out[13]: (True, 'Job completed')
In [14]: s.download_artifact('20e87cda3ef24cd18f065ccaf87e8ca4')
Out[14]:
(True,
         Diagnosis prob_BENIGN prob_MALIGNANT
 0
       BENIGN
                 0.665922
                                   0.334078
 1
       BENIGN
                  0.676795
                                   0.323205
 2
       BENIGN
                  0.676795
                                   0.323205
 3
       BENIGN
                   0.676795
                                   0.323205
 4
                 0.107175
                                   0.892825
    MALIGNANT
 5
    MALIGNANT
                  0.049802
                                   0.950199
                                   0.997893
 6
    MALIGNANT
                   0.002107
 7
                   0.676795
                                   0.323205
       BENIGN
 . . .
 95
       BENIGN
                   0.676795
                                   0.323205
 96
       BENIGN
                   0.676795
                                   0.323205
 97 MALIGNANT
                   0.050580
                                   0.949420
 98 MALIGNANT
                   0.032286
                                   0.967714
 99 MALIGNANT
                   0.052880
                                   0.947120
 [100 rows x 3 columns])
```

Revision Table

Version	Date	Notes
v 1.0	05-Feb-2018	Initial Release



Version	Date	Notes
v 1.2	28-Mar-2018	Added:
		DarwinSdk.auth_change_password
		DarwinSdk.delete_job
		DarwinSdk.stop_job
		 DarwinSdk.lookup_user
		 DarwinSdk.lookup_username
		DarwinSdk.auth_delete_user
		Name change: lookup_client to lookup_limits
v 1.3	23-May-2018	Added:
		DarwinSdk.auth_reset_password
		• DarwinSdk.auth_set_email
		 DarwinSdk.analyze_predictions
		Updated endpoints:
		• DarwinSdk.auth_register_user
		• DarwinSdk.analyze_model
v 1.3.1	14-Jun-2018	Fixed issues only. See Release Notes.
v 1.4	31-Jul-2018	 Island Models implemented to allow model types to reproduce
		at their own speeds
		 User selectable loss functions
		 Output model confidence value
		 Specify download paths for artifacts
		Parameter validation
		 Stored data is encrypted
		 DarwinSdk.lookup_model() and
		DarwinSdk.lookup_model_name(model_name) calls
		display model description
v 1.5	15-Oct-2018	New endpoints:
		DarwinSdk.clean_data
		DarwinSdk.download_dataset
		DarwinSdk.download_model
		Updated endpoints:
		DarwinSdk.analyze_data
		 DarwinSdk.download_artifact
		DarwinSdk.create_model
		 DarwinSdk.lookup_model



Version	Date	Notes
v 1.6	16-Jan-2019	New endpoints:
		 DarwinSdk.display_population
		 DarwinSdk.delete_all_artifacts
		Updated endpoints:
		DarwinSdk.analyze_data
		 DarwinSdk.download_artifact
		• DarwinSdk.create_model
		DarwinSdk.clean_data
v 1.6.1	06-Feb-2019	Fixed issues only. See Release Notes. Added on-prem installation notes.