

# Darwin TM Python SDK Guide

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# SparkCognition Darwin Python SDK Guide

## **Contents**

About this guide	3
Expectation	3
Darwin overview	3
Accessing the API	4
Darwin SDK interface	4
Setup Darwin SDK	4
Set up Users	5
Set up Admin account	5
Set up User accounts	5
Darwin SDK methods	7
URL Get/Set methods	7
DarwinSdk. <b>get_info()</b>	7
DarwinSdk.get_sdk_version()	7
DarwinSdk. <b>get_url</b> ()	8
DarwinSdk.set_url(url, version='v1')	8
Authentication methods	9
DarwinSdk.auth_register(password, api_key, email)	9
DarwinSdk.auth_login(password, api_key)	9
DarwinSdk.auth_register_user(username, password, email)	10
DarwinSdk. <b>auth_set_email</b> (username, email)	10
DarwinSdk.auth_login_user(username, password)	11
DarwinSdk.auth_change_password(curpass, newpass)	11
DarwinSdk.auth_reset_password(username)	12
DarwinSdk.auth_delete_user(username)	12
DarwinSdk.disable_ssl_cert_check()	13
DarwinSdk.enable_ssl_cert_check()	13
Job status methods	14
DarwinSdk.lookup_job_status(age=None, status=None)	14
DarwinSdk.lookup_job_status_name(job_name)	14
DarwinSdk. <b>delete_job</b> (job_name)	15
DarwinSdk. <b>stop_job</b> (job_name)	16
Lookup methods	16



DarwinSdk. <b>lookup_artifact</b> (type=None)	16
DarwinSdk.lookup_artifact_name(artifact_name)	17
DarwinSdk.lookup_limits()	17
DarwinSdk.lookup_dataset()	18
DarwinSdk.lookup_dataset_name(dataset_name)	19
DarwinSdk.lookup_model()	19
DarwinSdk.lookup_model_name(model_name)	20
DarwinSdk.lookup_tier()	20
DarwinSdk.lookup_tier_num(tier_num)	21
DarwinSdk.lookup_user()	21
DarwinSdk.lookup_username(username)	22
DarwinSdk.display_population(model_name)	23
Datasets and artifact methods	24
DarwinSdk. <b>upload_dataset</b> (dataset, dataset_name=None)	24
DarwinSdk.download_dataset(dataset_name)	25
DarwinSdk.delete_dataset(dataset_name)	25
DarwinSdk.download_model(model_name)	25
DarwinSdk.download_artifact(artifact_name, artifact_path=None)	26
DarwinSdk.delete_artifact(artifact_name)	30
Data Analysis and Data Cleaning methods	31
DarwinSdk.analyze_data(dataset_name, **kwargs)	31
DarwinSdk.clean_data(dataset_name, **kwargs)	34
Modeling and analysis methods	36
DarwinSdk.create_model(dataset_names, **kwargs)	36
DarwinSdk.delete_model(model_name)	39
DarwinSdk. <b>resume_training_model</b> (model_name, dataset_names, **kwargs)	39
DarwinSdk.analyze_model(model_name, job_name=None, artifact_name=None)	40
$Darwin Sdk. \textbf{analyze\_predictions} (model\_name,  dataset\_name,  job\_name=None,  article (a) article (a) article (b) article (b) article (b) article (b) article (c) artic$	
fact_name=None)	40
$Darwin Sdk. \textbf{run\_model} (dataset\_name, model\_name, job\_name=None, artifact\_name=None) \; .$	
Convenience methods	42
DarwinSdk.delete_all_datasets()	42
DarwinSdk.delete_all_models()	42
DarwinSdk.delete_all_artifacts()	43
DarwinSdk.wait_for_job(job_name, time_limit=600)	43
DarwinSdk. <b>help</b> ()	43
Reference	44
SDK modeling examples	
Supervised modeling example	
Unsupervised modeling example	
NBM modeling example	
Contact Support	
Revision Table	



## About this guide

This guide describes using the  $Darwin^{T}$  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 (...).

The Darwin SDK has an independent version number to allow for release outside of the normal Darwin product release window. As of this printing, the Darwin SDK is at version 1.45.0.

The documentation for this version of Darwin includes:

- The Darwin Release Notes, version 2.0.4
- The Darwin User Interface Guide, version 2.0.4
- The Darwin API User Guide, version 1.36.0
- The Darwin Python SDK User Guide, version 1.45.0
- The Darwin RTE User Guide, version 2.0.4

All of these documents are available for download from the Darwin support portal.

## **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 $^{\text{M}}$  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
- Analyze training data
- · Clean training data
- Create model
- Wait for job to complete
- Upload test data
- · Clean test data
- · Run the model
- Wait for job to complete
- · Download the 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 (no more than 4000) and data samples that span rows. Plan your data file so it fits this expectation to help prevent errors.

See the SDK examples for modeling examples of supervised, unsupervised, and normal behavior modeling (NBM) problems.

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 set up the Darwin SDK, unless you have already set up your service and created users using the API.
  - 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 darwin-sdk
```



**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/

**Note:** Ensure you have a trailing slash (/) on the production URL.

**ON-PREM ONLY**: For on-prem installations, the product URL will be in the form:

```
https://customerdomainname.customerdomain.com:8000/v1/
```

Note: On-prem installation must add port 8000 to the product URL.

8. Verify the connection.

Use get\_url() and set\_url() to verify connection to the correct Darwin service. See the URL Get/Set methods below for more information.

## Set up Users

Before you can set up any user accounts, you need to know your api key, also known as an admin key. This key can be obtained from SparkCognition support or your IT manager. The api key is a long string, for example:

## Set up Admin account

Register the api key using the auth\_register() method.

The purpose of this method is to create a password and an email address for the Darwin admin account. This method must be invoked once for each api key to establish an admin account for that key.

## Example

```
>>> from amb_sdk.sdk import DarwinSdk
>>> s = DarwinSdk()
>>> s.auth_register('adminpassword', 'iRgwut4kGs0ymULiuKtMd0WFvBYLMWSj16q2uysQeteq\
UvcysnPojRpfycLVHa2IlN1IlrfEk1YMA', 'admin@company.com')
(True,'Bearer eyJ0eXAiOiJKV1QiLCJhbGciOiJIUzI1NiJ9.eyJleHAiOjE1MTU1MzM4NjEsImlh\
dCI6MTUxNTUzMDI2MS ... F56xZQiBT-89nrRz1nIXD5LfawHIj MlUHQqM36vU')
```

## Set up User accounts

While you can use the SDK as an admin, it is more convenient to create additional user accounts so that you can have certain datasets/models be owned by specific users. Perform the following to create additional user accounts:

<sup>&</sup>quot;RsJ74ZS5AvwznbHh0AfVSgrchhS9KxACDy3jefaQnxb9f6QTSDBFmhnGa0cOIWtNAIFRAG9ToOTpi0mn"



Log in to the *service* as an admin. In the following example, you need to enter your admin password and the api key.

## Example

```
>>> s.auth_login('adminpassword', 'iRgwut4kGs0ymULiuKtMd0WFvBYLMWSj16q2uysQeteq\
UvcysnPojRpfycLVHa2IlN1IlrfEk1YMA')
(True,'Bearer iLCJhbGciOeyJ0eXAiOiJKV1QiJIUzI1NiJ9.eyJleHAiOjE1MTU1MzQxNzIsImlh\
dUxNTUCI6MTzMD ... UQQfoXqYFKJSoRXXDNPE985-a08cE6_o')
```

#### 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 2 hours, if the DarwinSdk object session life time exceeds 2 hours, the SDK will request another auth token until the session ends.

Register a new user by calling the auth\_register\_user() method. You need to input the username, password, and email address for the new user.

## Example

```
>>> s.auth_register_user('user1', 'user1-password', 'user1@company.com')
(True,
   'Bearer eyJhbGciOiJIUzI1NiIsInR5cCI6IkpXVCJ9.eyJqdGkiOiJkNjY0MmJjOC1iMmU5LTQxO\
DctODFlNS00YjI2MD ... 5zMp_1FfxU')
```

You can repeat this procedure for additional users.

The user can now log in by using the auth\_login\_user() method. The user needs to input the username and password.

#### Example

```
>>> s.auth_login_user('userl', 'userl-password')
(True,
   'Bearer eyJhbGciOiJIUzI1NiIsInR5cCI6IkpXVCJ9.eyJqdGkiOiI3NGYzYmUxZS0yOTlmLTRhN\
zMtODU5ZC01NGRmM2F ... u1zGCeCONA')
```

The user is now logged in and can perform other functions. See the following sections for other SDK methods.



## Darwin SDK methods

## **URL Get/Set methods**

## DarwinSdk.get\_info()

Get info on the routes available and the API version. The local flag will return True for an on-prem installation.

Parameters: None

#### Returns:

```
(True, {available_routes: {}, local: False})
```

## Example

```
In [29]: s.get_info()
Out[29]: (True,
{'available_routes': {'Info': True,
'Auth': True,
'Job': True,
'Metadata': True,
'Train': True,
'Risk': True,
'Upload': True,
'Download': True,
'Analyze': True,
'Run': True,
'Admin': True,
'Clean': True,
'Model': True},
'local': False, 'api_version': '1.34.0'})
```

## DarwinSdk.get\_sdk\_version()

Get the version of the SDK.

Parameters: None

## Returns:

```
(True, '1.44.0')
```

```
In [8]: s.get_sdk_version()
Out[8]: (True, '1.44.0')
```



## 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 for SaaS**

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

## **Example for On-prem**

```
In [9]: s.set_url('https://customerdomainname.customerdomain.com:8000/v1/')
Out[9]: (True, 'https://customerdomainname.customerdomain.com:8000/v1/')
```



#### **Authentication methods**

## DarwinSdk.auth\_register(password, api\_key, email)

Register the api key, also known as an admin key, as a service and establish an admin account. The purpose of this method is to set a password and an email address for the Darwin Admin account. This method is invoked only once for each api key to establish a password and Admin account. After registration, the admin can log in to the service using the *auth\_login()* method.

#### Parameters:

- password The service level password for the admin
- 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 2 hours, if the DarwinSdk object session life time exceeds 2 hours, the SDK will request another auth token until the session ends.

#### Example

```
In [4]: s.auth_register('adminpassword', 'RsJ74ZS5AvwznbHh0AfVSgrchhS9KxACDy\
3jefaQnxb9f6QTSDBFmhnGa0cOIWtNAIFRAG9ToOTpi0mnEo3zFA', 'admin@company.com')
Out[4]:
(True,
   'Bearer eyJhbGciOiJIUzI1NiIsInR5cCI6IkpXVCJ9.eyJleHAiO...iSdU8x1F4yJk')
```

## DarwinSdk.auth\_login(password, api\_key)

Log in to the service as an admin.

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

#### Parameters:

- password The service level password for the admin
- $api\_key$  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 2 hours, if the DarwinSdk object session life time exceeds the 2 hour limit, the SDK will acquire another auth token until the session ends.

## Example

## 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 2 hours, if the DarwinSdk object session life time exceeds the 2 hour limit, the SDK will acquire another auth token until the session ends.

## Example

## 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 2 hours, if the DarwinSdk object session life time exceeds the 2 hour limit, the SDK will acquire another auth token until the session ends.

## Example

## 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 <code>DarwinSdk.auth\_reset\_password</code> (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('userl-password', 'userl-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('user1')
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>)
```



## 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] \setminus
but did you mean /v1/auth/register/user/ <username> or /v1/auth/register/user \
 or /v1/auth/register ?"\n}\n')
```

## DarwinSdk.disable\_ssl\_cert\_check()

For on-prem customers, this disables the SSL certificate check when calling auth\_login\_user(<username>, <password>

If you want to re-enable the SSL certificate check, call the enable\_ssl\_cert\_check() method.

Parameters: None

## DarwinSdk.enable\_ssl\_cert\_check()

For on-prem customers, this enables the SSL certificate check when calling auth\_login\_user(<username>, <password>

If you want to disable the SSL certificate check, call the disable\_ssl\_cert\_check() method.

Parameters: None



#### 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>)
```

## Example

```
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>)
```



```
'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 running job. The job will not stop right away, but it will stop when the current generation is complete.

#### Parameter:

• *job\_name* - The name of the job you want to stop.

#### Returns:

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

## Example

```
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>)
```

```
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>)
```

## Example:

## 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>)
```



## 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 dataset name. The dataset name is established in the **upload\_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>)
```



```
)
} ]
```

## 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>)
```

## Example

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

```
In [41]: s.lookup_tier()
Out[41]:
(True,
  [{'job_limit': None,
```



```
'model_limit': None,
'tier': 0,
'upload_limit': None,
'user_limit': None},
{'job_limit': 10000,
'model_limit': 10000,
'tier': 1,
'upload_limit': 10000,
'user_limit': 10000])
```

## 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>)
```

## Example

## 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>)
```



```
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
                   }
                ],
                "loss_function": "CrossEntropy",
                "fitness": 1.9667300770467946
              },
              "RandomForest": {
                "model_description": {
                   "type": "RandomForestClassifier",
```



## Datasets and artifact methods

## DarwinSdk.upload\_dataset(dataset, dataset\_name=None)

Upload a dataset.

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

**Note:** For .csv files, ensure they are encoded to one of the following standards:

- utf-8
- us-ascii
- iso-8859-1
- iso-8859-2
- ascii

Note: The maximum size that can be uploaded is 10GB due to only supporting uploading data via http.

## 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

```
In [5]: s.download_dataset('cancer-cleandata3', \
    artifact_path='/Users/username/Downloads/artifacts')

Out[5]:
(True,
    {'filename': \
    '/Users/username/Downloads/artifacts/cancer-cleandata3-cleaned-8m38g07j.csv'})
```

## 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*. The ONNX format is only available for neural network models.

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

## Example analyze\_data() artifact

```
In [97]: s.download_artifact('1a38f1af934c4cbabb9136ee94f72718')
Out[97]:
(True,
                                       drop is_date low_samples max \
                                count
name
                                599 False
                                              False
                                                         [] 8233704.0
Code
                                                                 10.0
Clump Thickness
                                599 False
                                             False
                                                         []
Uniformity of Cell Size
                                599 False
                                             False
                                                                  10.0
                                                         []
 Uniformity of Cell Shape
                                599 False
                                             False
                                                         []
                                                                 10.0
Marginal Adhesion
                                599 False
                                             False
                                                         []
                                                                 10.0
 Single Epithelial Cell Size
                                599 False
                                             False
                                                         []
                                                                 10.0
Bare Nuclei
                                599 False
                                             False
                                                         []
                                                                 10.0
Bland Chromatin
                                599 False
                                             False
                                                         []
                                                                  10.0
Normal Nucleoli
                                599 False
                                             False
                                                         []
                                                                  10.0
                                599 False
                                                         []
Mitoses
                                              False
                                                                  10.0
                                599 False
                                                         []
Diagnosis
                                              False
                                                                  NaN
```



```
mean
               min
                              missing num_uniques
                                                           stddev \
                                               557
1.044171e+06 6.163400e+04
                                 0.0
                                                       4.140964e+05
4.555927e+00 1.000000e+00
                                                       2.887488e+00
                                 0.0
                                                10
3.215359e+00
             1.000000e+00
                                 0.0
                                                10
                                                       3.044601e+00
3.287145e+00 1.000000e+00
                                                       2.971045e+00
                                 0.0
                                                10
2.859766e+00 1.000000e+00
                                 0.0
                                                       2.873655e+00
                                                10
3.290484e+00 1.000000e+00
                                 0.0
                                                       2.275159e+00
                                                10
-2.309692e+17 -9.223372e+18
                                 0.0
                                                       1.442374e+18
                                                11
3.520868e+00 1.000000e+00
                                                       2.369500e+00
                                 0.0
                                                10
             1.000000e+00
2.966611e+00
                                 0.0
                                                10
                                                       3.084466e+00
1.607679e+00 1.000000e+00
                                 0.0
                                                 9
                                                       1.734369e+00
NaN
              NaN
                                 0.0
                                                 2
                                                             NaN
top_unique_values
                                    treatment
                                                         type_note \
                                                  type
                                                  int64
    []
                                     numeric
                                                  int64
    []
                                     numeric
    []
                                                  int64
                                     numeric
    []
                                     numeric
                                                  int64
                                                  int64
    []
                                     numeric
                                                  int64
    []
                                     numeric
                                                  int64
    []
                                     numeric
    []
                                                  int64
                                     numeric
    []
                                                  int64
                                     numeric
                                     numeric
                                                  int64
    []
[[BENIGN, 379], [MALIGNANT, 220]] categorical
                                                  object
uniques
[61634, 63375, 76389, 95719, 128059, 142932, 1...
[1, 2, 3, 4, 5, 6, 7, 8, 9, 10]
[1, 2, 3, 4, 5, 6, 7, 8, 9, 10]
[1, 2, 3, 4, 5, 6, 7, 8, 9, 10]
[1, 2, 3, 4, 5, 6, 7, 8, 9, 10]
[1, 2, 3, 4, 5, 6, 7, 8, 9, 10]
[-9223372036854775808, 1, 2, 3, 4, 5, 6, 7, 8, ...
[1, 2, 3, 4, 5, 6, 7, 8, 9, 10]
[1, 2, 3, 4, 5, 6, 7, 8, 9, 10]
   [1, 2, 3, 4, 5, 6, 7, 8, 10]
            [BENIGN, MALIGNANT])
```

#### Example analyze\_model() or prediction artifact

```
In [5]: s.download_artifact('6e4861de29424cb7ad09e467d1869c17',\
   'path_to_download_dir/')
Out[5]:
```



```
True RM
                      0.216088
               0.141956
CRIM
              0.134069
LSTAT
              0.104101
DIS
PTRATIO
              0.089905
AGE
              0.078864
NOX
               0.074132
В
              0.067823
TAX
              0.045741
INDUS
              0.023659
              0.011041
ZN
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()
         ##Sample return for regression, has predicted_value column
Out[9]:
       AGE
                    B CHAS = 1.0
                                       CRIM
                                                  DIS
                                                          INDUS
                                                                    LSTAT \
0 0.000000 0.000000
                       0.000000 -0.664664 -0.923219 -0.720941 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 0.632119 0.079678
                       0.000000 0.076080 -0.488128 -0.016690 -0.102031
            PTRATIO RAD = 2.0
                                                   RAD = 4.0 RAD = 5.0 \setminus
       NOX
0 -0.342404 0.224360
                             0.0
                                                   -0.641678 -0.570788
                                                   0.000000 -0.741561
1 -0.556636 -2.168356
                             0.0
2 0.000000 1.458677
                                                    0.000000 - 0.340486
                             0.0
                                       . . .
3 -0.945060 -1.068743
                             0.0
                                                    0.000000
                                                              0.217991
                                       . . .
4 0.309544 0.298940
                             0.0
                                                    0.000000
                                                              -0.047708
                                       . . .
  RAD = 6.0
            RAD = 7.0 RAD = 8.0
                                                   TAX
                                          RM
                                                              ZN base_value
0
         0.0
                    0.0
                               0.0 - 1.835851 - 0.563795 - 0.600155
                                                                    21.63455
1
         0.0
                    0.0
                               0.0 -1.016655 -0.699813 -0.727181
```



```
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
4
         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
            0.292612
1
                               0.315358
                                                   0.019236
                                                                     -0.014442
2
            0.325615
                               0.329229
                                                   0.003208
                                                                      0.016954
3
            0.232265
                               0.410938
                                                   0.043014
                                                                      0.004154
            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
     0.324994
2
                       1.000000
                                         setosa
3
     0.309628
                       1.000000
                                      virginica
     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>)
```

```
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.

**Note:** *upload\_dataset()* is currently artificially limited to 10GB due to only supporting uploading data via http. 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.

**Note**: You can only analyze a dataset once. If you try to analyze the dataset a second time, you will get a 400: BAD REQUEST error.

#### Parameters:

dataset\_name - (required) 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.
- target: String denoting target prediction column in input data.
- 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.
- ullet 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,
                                                  drop is_cat
                         col name
                                      col_type
                                                                        max
 0
                              PID
                                    StringType
                                                  True False
                                                               2205663001_
 1
                           ST_NUM
                                    StringType
                                                  True False
                                                                        999
 2
                          ST_NAME
                                    StringType
                                                  True False
                                                                     ZELLER
 3
                                    StringType
                     ST_NAME_SUF
                                                  True False
                                                                         ΧT
 4
                          ZIPCODE
                                    StringType
                                                  True
                                                        False
                                                                     02467_
 5
                  Assessed_Value
                                         int64
                                                  True
                                                       False
                                                                  23095700
                                         int64
 6
                         Lot_Area
                                                  True False
                                                                     107158
 7
                      Gross_Area
                                         int64
                                                  True
                                                        False
                                                                      23335
 8
                     Living_Area
                                         int64
                                                                      21711
                                                  True
                                                       False
 9
                  Owner_Occupied
                                  categorical
                                                False
                                                         True
                                                                      None
                      Year Built
 10
                                         int64
                                                  True False
                                                                       2016
                Number_of_Floors
                                       float64
                                               False
 11
                                                       False
                                                                        5.0
 12
           Total_Number_of_Rooms
                                         int64
                                                  True
                                                       False
                                                                         27
              Number of Bedrooms
 13
                                   categorical
                                                False
                                                         True
                                                                       None
 14
            Number_of_Full_Baths
                                  categorical
                                                False
                                                         True
                                                                       None
 15
            Number_of_Half_Baths
                                  categorical
                                                False
                                                         True
                                                                       None
              Number_of_Kitchens
 16
                                   categorical
                                                False
                                                         True
                                                                       None
 17
                           Has_AC
                                   categorical
                                                False
                                                         True
                                                                       None
 18
            Number_of_Fireplaces
                                   categorical
                                                False
                                                         True
                                                                       None
 19
     Year_Since_Remodel_or_Build
                                         int64
                                                  True False
                                                                        307
 20
                  Year_Remodeled
                                    StringType
                                                  True
                                                        False
                                                               Unremodeled
 21
                  Structure_Type categorical
                                                False
                                                         True
                                                                       None
 22
                  Building_Style
                                    StringType
                                                  True
                                                        False
                                                                 Victorian
 2.3
                        Roof_Type
                                  categorical
                                               False
                                                         True
                                                                       None
 24
                 Exterior Finish
                                   categorical
                                                False
                                                         True
                                                                       None
 25
             Main_Bathroom_Style
                                   categorical
                                                False
                                                         True
                                                                       None
 26
              Main_Kitchen_Style
                                   categorical
                                               False
                                                         True
                                                                       None
 27
                    Heating_type
                                  categorical False
                                                         True
                                                                       None
 28
              Exterior_Condition
                                   categorical False
                                                         True
                                                                       None
 29
               Overall_Condition
                                   categorical
                                               False
                                                         True
                                                                       None
 30
              Interior_Condition
                                  categorical
                                               False
                                                         True
                                                                       None
 31
                 Interior_Finish
                                   categorical
                                               False
                                                         True
                                                                       None
 32
                             View
                                   categorical False
                                                         True
                                                                       None
```



	mean	min	missing	num_uniques	scalable	\
0	None	0100021000_	0.000000	28578	True	
1	122.09705524787249	1005R	0.010223	1922	True	
2	None	ABBOTSFORD	0.000000	2246	True	
3	None	ST	0.003015	21	True	
4	None	02108_	0.000000	28	True	
5	534716.6815977456	101300	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	
1	0 1926.970935185792	1710	0.000000	225	True	
1	1 1.8748115866046269	1.0	0.000000	9	True	
1	2 7.233632610262796	2	0.000000	26	True	
1	3 3.3851169801428664	None	0.000000	12	False	
1	4 1.4273543482534898	None	0.000000	10	False	
1	5 0.5716953928828888	None	0.000000	7	False	
1	6 1.0287043711907726	None	0.000000	4	False	
1	7 0.18733206632151517	None	0.000000	2	False	
1	8 0.590995478078511	None	0.000000	13	False	
1	9 60.88419948882627	1	0.000000	190	True	
2	0 2000.3376960831488	1890	0.000000	82	True	
2	1 None	None	0.000000	5	False	
2	2 None	Bi-Level	0.000000	17	True	
2	None	None	0.000000	7	False	
2	4 None	None	0.000000	13	False	
2	5 None	None	0.000000	4	False	
2	6 None	None	0.000000	4	False	
2	7 None	None	0.000000	6	False	
2	8 None	None	0.000000	5	False	
2	9 None	None	0.000000	5	False	
3	0 None	None	0.000000	5	False	
3	1 None	None	0.000000	3	False	
3	None 2	None	0.000000	5	False	
	stddev				u	ıniques
0	None					None
1	294.1511958893473					None
2	None					None
3	None					None
4	None					None
5	634750.7826113638					None
6	3218.286557124007					None
7	1069.3847598444354					None
8	758.9874732061347					None



```
9
     0.3658237412175791
                                                                        [0, 1]
10
       34.9170355483078
                                                                         None
11
     0.5737101635770085
                                                                         None
12
     1.8082562295656077
                                                                          None
13
     1.0095185504254367
                                     [12, 9, 1, 5, 2, 6, 3, 10, 7, 4, 11, 8]
14
     0.6850264359951297
                                             [12, 9, 1, 5, 2, 6, 3, 7, 4, 8]
                                                        [0, 1, 5, 2, 6, 3, 4]
15
     0.5645602408681473
                                                                 [0, 1, 2, 3]
    0.17162236936210065
16
17
     0.3901842537872663
                                                                        [0, 1]
18
     0.8584446055814273
                                  [0, 12, 9, 1, 5, 2, 6, 3, 10, 7, 4, 11, 8]
     43.323487380439225
19
                                                                         None
20
     13.578956800881818
                                                                         None
21
                          ['Residential', 'Wood/Frame', 'Unknown', 'Bric...
                    None
2.2
                    None
23
                          ['Shed', 'Gambrel', 'Flat', 'Other', 'Mansard'...
                    None
24
                          ['Cement Board', 'Frame/Clapboard', 'Wood Shak...
                    None
25
                          ['Semi-Modern', 'Luxury', 'No Remodeling', 'Mo...
                    None
                          ['Semi-Modern', 'Luxury', 'No Remodeling', 'Mo...
2.6
                    None
27
                          ['Electric', 'Other', 'None', 'Hot Water', 'Sp...
                    None
28
                           ['Poor', 'Good', 'Excellent', 'Average', 'Fair']
                    None
                           ['Poor', 'Good', 'Excellent', 'Average', 'Fair']
29
                    None
                           ['Poor', 'Good', 'Excellent', 'Average', 'Fair']
30
                    None
                                      ['Elaborate', 'Normal', 'Substandard']
31
                    None
32
                    None
                           ['Poor', 'Good', 'Excellent', 'Average', 'Fair'] )
```

## 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 <code>clean\_data()</code> before creating a model, you must specify a dataset\_name and a target. When you run <code>clean\_data()</code> 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 (required) 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.



- index: String denoting the date/time column name to use as an index.
- 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.	
'median'	Median Fill: Computes the median value of all non-missing	Linear
	examples in a column to fill in missing examples. While the result	Fast
	is interpretable in terms of the input space for categorical variables,	
	the approach might not be appropriate for non-ordinal data.	
'Linear'	Linear Interpolation Fill: Interpolation using a Linear function.	Linear
	Useful for timeseries or sequential data.	Fast

- 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.

### Example



# 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 - (required) 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.

fit\_profile\_name: (required) This is the profile\_name that is generated from the clean\_data call.

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

### parameters -

- *val\_size*: Portion of the dataset to be used as a validation set during training, expressed as a decimal that is greater than 0 and less than 1. Default value is 0.2 (i.e., 20%).
- *cv\_kfold*: k-fold cross-validation, where k is the number of groups that a given data sample is to be split into for training/validation. Default is 1 for non-timeseries data or 3 for timeseries data. Maximum value allowed is 10.
- *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*: 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 can result in slower model evolution. If you want to see the LSTM and TCN models used during training or if you want to treat your problem as a time series problem, you must set recurrent=True.
- *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.
- *forecast\_horizon* (forecasting only): Integer indicating how long in the future you want to forecast predictions. For example, if you have 6 months of time-series data and each row represents a 1 day interval and you want to predict the next week of data, you should set <code>forecast\_horizon=7</code>. If each row is a 1 hour interval, then you should set <code>forecast\_horizon=168</code>. (168 = 7\*24)



**Note**: For best results, be sure that the minimum length to run the model is 5 times of the *forecast\_horizon* specified.

- *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.
- *class\_weights*: A string to indicate how relatively important each class is for predictive correctness. This is done by providing a numeric value to each class. Note that the class name is case-sensitive. The following is an example *class\_weights* setting:

```
class_weights = "{'BENIGN': 4, 'MALIGNANT': 6}"
```

• loss\_fn\_name: Specify the loss function. Possible values include: "CrossEntropy", "MSE", "BCE", "L1", "NLL", "BCEWithLogits", "SmoothL1".

"CrossEntropy", "BCE", and "BCEWithLogits" 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.

For classification problems, possible values include:

- average\_precision (Average Precision) Measures the average precision across the spectrum of all recall values from 0 to 1. Average precision is a good metric to use for imbalanced problems, and only works on binary target columns, that is, there are two class labels being predicted.
- roc\_auc (ROC Area Under Curve) Measures the area under the Receiver Operating Characteristics curve, which plots the relationship between precision and recall for a model. ROC area under curve only works on binary target columns, that is, there are two class labels being predicted.
- accuracy (Accuracy) Measures the total number of correct predictions divided by the total number of predictions made.
- f1\_weighted (F1 Weighted) (default) Measures the F1 score for each label and finds their average, which is weighted by the number of true instances for each label. This alters 'macro' to account for label imbalance.
- f1\_macro (F1 Macro) Measures the F1 score, but calculates metrics for each label, and finds their unweighted mean. This is recommended for imbalanced problems.
- f1\_micro (F1 Micro) Measures the F1 metrics globally by counting the total true positives, false negatives, and false positives.
- balanced\_accuracy (Balanced Accuracy) Measures the proportion correct of each class individually and then averages those values. This is a good metric to use for imbalanced problems.
- neg\_log\_loss (Log Loss) Measures the prediction probability of each output and how closely that maps to the actual label. In binary classification, if the actual label was 0 and the prediction probability was 0.01, the prediction would be 0.49 better than a prediction probability of 0.5. This is a very harsh penalty mechanism and will result in a model that tries to find a very defined boundary between classes.
- precision\_macro (Precision Macro) Measures precision for each label and finds their un-



weighted mean. This is recommended for imbalanced problems.

- precision\_micro (Precision Micro) Measures the precision metrics globally by counting the total true positives predicted.
- precision\_weighted (Precision Weighted) Measures the precision score for each label, and
  then finds their average weighted by the number of true instances for each label. This alters
  'macro' to account for label imbalance.
- recall\_macro (Recall Macro) Measures recall for each label and finds their unweighted mean.
   This is recommended for imbalanced problems.
- recall\_micro (Recall Micro) Measures the recall metrics globally by counting the total true positives predicted.
- recall\_weighted (Recall Weighted) Measures the recall score for each label and finds their average weighted by the number of true instances for each label. This alters 'macro' to account for label imbalance.

For regression problems, possible values include:

- r2 (R<sup>2</sup>) (default) Measures how closely the data maps to the fitted regression line. It is also known as the coefficient of determination and is useful for mapping the relationships that exist in data.
- neg\_mean\_absolute\_error (Mean Absolute Error) Measures the average error for each
  predicted data point versus the expected value. This is useful as a good baseline metric or for
  capturing general trends.
- neg\_mse (Mean Squared Error) Measures the square of the average error for each predicted data point versus the expected value. This is useful if you want to penalize large errors more harshly.
- neg\_median\_absolute\_error (Median Absolute Error) Measures the median error for the
  predicted data point versus the expected value. This is useful if your dataset has biases toward
  certain values.
- neg\_rmse (Root Mean Squared Error) Measures the square root Mean Squared Error values. This is useful if there are not a lot of outliers in your data.
- neg\_rmsle (Root Mean Squared Logarithmic Error) Measures the ratio between the actual and predicted values by calculating the square root of the Mean Squared Error values in which a logarithmic transform is performed on predicted and actual values. This is useful for targets with very large numbers or that contain outliers. An error will be generated if a negative target value is encounted. This fitness function should only be used for positive datasets.
- *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

# DarwinSdk.analyze\_model(model\_name, job\_name=None, artifact\_name=None)

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

**Note**: This method is supported for clustering and NBM models. It does not support forecasting or unsupervised anomaly detection.

#### Parameters:

- *model\_name* (required) 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.
- *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'})
```

### 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 100 MB.

**Note**: This method is not supported in forecasting models or for clustering/anomaly detection, however it does support NBM modeling.



#### Parameters:

- *dataset\_name* (required) 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 (required) 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) User can specify a model type to use for their prediction. If nothing is defined, the SDK will use the best model type. Possible values include:
  - DeepNeuralNetwork: The run\_model command will pick the best performing neural network to use when running the prediction.
  - RandomForest: The run\_model command will pick the best performing sklearn random forest to use when running the prediction.
  - GradientBoosted: The run\_model command will pick the best performing sklearn gradient boosted model to use when running the prediction.

### Returns:

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

# Example

### 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. If the *time\_limit* is reached, your job will continue to run but **wait\_for\_job** will discontinue monitoring it. You can re-run **wait\_for\_job** or modify the *time\_limit* parameter.

#### Parameters:

- job\_name The id for the job
- time\_limit (optional) defaults to 600 seconds

### **Returns:**

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

# DarwinSdk.help()

Shows all the methods available.

Parameters: None

### Example

```
In [5]: s.help()
Out [5]:
analyze_data (self, dataset_name, **kwargs)
analyze_model (self, model_name, job_name=None, artifact_name=None, \
    category_name=None, model_type=None)
analyze_predictions (self, model_name, dataset_name, job_name=None, \
    artifact_name=None, model_type=None)
auth_change_password (self, curpass, newpass)
auth_delete_user (self, username)
auth_login (self, password, api_key)
auth_login_user (self, username, password)
auth_register (self, password, api_key, email)
auth_register_user (self, username, password, email)
```



```
auth_reset_password (self, username)
auth_set_email (self, username, email)
clean_data (self, dataset_name, **kwargs)
create_model (self, dataset_names, **kwargs)
delete_all_artifacts (self)
delete_all_datasets (self)
delete all models (self)
delete_artifact (self, artifact_name)
delete_dataset (self, dataset_name)
delete_job (self, job_name)
delete_model (self, model_name)
disable_ssl_cert_check (self)
display_population (self, model_name)
download_artifact (self, artifact_name, artifact_path=None)
download_dataset (self, dataset_name, file_part=None, artifact_path=None)
download_model (self, model_name, path=None, model_type=None, model_format=None)
enable_ssl_cert_check (self)
get_info (self)
get_url (self)
lookup_artifact (self, type=None)
lookup_artifact_name (self, artifact_name)
lookup dataset (self)
lookup_dataset_name (self, dataset_name)
lookup_job_status (self, age=None, status=None)
lookup_job_status_name (self, job_name)
lookup_limits (self)
lookup_model (self)
lookup_model_name (self, model_name)
lookup_tier (self)
lookup_tier_num (self, tier_num)
lookup_user (self)
lookup_username (self, username)
resume_training_model (self, model_name, dataset_names, **kwargs)
run_model (self, dataset_name, model_name, **kwargs)
set_url (self, url, version='v1')
stop_job (self, job_name)
upload_dataset (self, dataset_path, dataset_name=None, has_header=True)
wait_for_job (self, job_name, time_limit=600)
```

# Reference

- SDK modeling example
- Revision table



# SDK modeling examples

The following section details modeling examples for the following types of problems:

- Supervised
- Unsupervised
- NBM

# Supervised modeling example

```
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 eyJ0eXAiOiJK...A8sj4pAzX1FpMMscwY_rMJbnGo0YQ_4')
In [14]: s.upload_dataset('sets/cancer_train.csv', 'pacman-cancer')
Out[14]: (True, {'dataset_name': 'pacman-cancer'})
In [15]: s.analyze_data('pacman-cancer')
Out[15]:
(True,
{'job_name': '3b3a54324a68427583ccae1194822fdd',
 'job_id': '688d5aba-a63b-11e9-b969-d3fc6b14b182',
 'artifact_name': '5b38f0d797cd45c5a7081a0c1b02ccad'})
In [16]: s.clean_data('pacman-cancer', target='Diagnosis')
Out[16]:
(True,
{'job_name': '77b47b391b0d483699fe0741cc52e6ad',
 'job_id': '72756658-a63b-11e9-824c-c75a5cadd55f',
 'profile_name': '60f9429a0a2141d9ad98c3d073b8425f',
 'profile_id': '7276d1d2-a63b-11e9-824c-6be79b2f3967'})
In [18]: s.create_model('pacman-cancer', \
fit_profile_name='60f9429a0a2141d9ad98c3d073b8425f', max_train_time='00:02')
Out[18]:
(True,
{'job_name': 'b29d680547f94d1e87bf0e6ae6913ae0',
 'job_id': '81a68bde-a63b-11e9-aa27-2ba9c50e485f',
 'model_name': '1f871bd9b0b3405680a2bce0c3b2b226'})
```



```
In [19]: s.wait_for_job('b29d680547f94d1e87bf0e6ae6913ae0')
{'status': 'Running', 'starttime': '2019-07-14T08:30:11.995459', \
 'endtime': None, 'percent_complete': 0, 'job_type': 'TrainModel', \
 'loss': None, 'generations': 0, 'dataset_names': ['pacman-cancer'], \
 'artifact_names': None, 'model_name': '1f871bd9b0b3405680a2bce0c3b2b226', \
 'job_error': ''}
{'status': 'Running', 'starttime': '2019-07-14T08:30:11.995459', \
'endtime': None, 'percent_complete': 50, 'job_type': 'TrainModel', \
'loss': 0.20505395531654358, 'generations': 1, 'dataset_names': ['pacman-cancer'], \
'artifact_names': None, 'model_name': '1f871bd9b0b3405680a2bce0c3b2b226',\
 'job error': ''}
. . .
'generations': 29, 'dataset_names': ['pacman-cancer'], 'artifact_names': None, \
'model_name': '1f871bd9b0b3405680a2bce0c3b2b226', 'job_error': ''}
{'status': 'Complete', 'starttime': '2019-07-14T08:30:11.995459', \
'endtime': '2019-07-14T08:34:40.40471', 'percent_complete': 100, \
'job_type': 'TrainModel', 'loss': 0.20505395531654358, 'generations': 31, \
'dataset_names': ['pacman-cancer'], 'artifact_names': None, \
'model_name': '1f871bd9b0b3405680a2bce0c3b2b226', 'job_error': ''}
Out[19]: (True, 'Job completed')
In [21]: s.run model('pacman-cancer', '1f871bd9b0b3405680a2bce0c3b2b226')
Out [21]:
(True,
{'job name': 'dfd758d43bda429cb19b4d9460db689d',
 'job_id': '612b7b2a-a63c-11e9-acd6-7b9b48091c5f',
 'artifact_name': '2f8502b4e3494ba3b5ce2133e066ec1a'})
In [22]: s.wait_for_job('dfd758d43bda429cb19b4d9460db689d')
{'status': 'Complete', 'starttime': '2019-07-14T08:36:26.997233', \
 'endtime': '2019-07-14T08:36:40.418987', 'percent_complete': 100, \
 'job_type': 'RunModel', 'loss': 0.20505395531654358, 'generations': 31, \
 'dataset_names': ['pacman-cancer'], \
 'artifact_names': ['2f8502b4e3494ba3b5ce2133e066ec1a'], \
 'model_name': '1f871bd9b0b3405680a2bce0c3b2b226', 'job_error': ''}
Out[22]: (True, 'Job completed')
In [23]: s.download_artifact('2f8502b4e3494ba3b5ce2133e066ec1a')
Out [23]:
(True,
           Diagnosis prob_BENIGN prob_MALIGNANT
     0
            BENIGN
                      0.896724
                                        0.103276
     1
         MALIGNANT
                       0.313960
                                        0.686040
    . . .
                       0.886773
                                        0.113227
     597
             BENIGN
                                        0.068040
     598
             BENIGN
                        0.931960
```



```
[599 rows x 3 columns])
In [26]: s.upload_dataset('sets/cancer_test.csv', 'pacman-cancertest')
Out[26]: (True, {'dataset_name': 'pacman-cancertest'})
In [27]: s.clean_data('pacman-cancertest', \
model_name='1f871bd9b0b3405680a2bce0c3b2b226')
Out [27]:
(True,
{'job_name': '462ade77431d4fde92b780ddc00573d9',
 'job_id': 'a868dd5c-a63c-11e9-94d1-8b9399879e43',
 'artifact_name': 'd8d4e61a4c624e70abb8bf66fde42e45',
 'artifact id': 'a86a8a30-a63c-11e9-94d1-93c25fc085f5'})
In [28]: s.run_model('pacman-cancertest', \
'1f871bd9b0b3405680a2bce0c3b2b226')
Out [28]:
(True,
    {'job_name': '015bc7fa826c4f36b5ca2b4e8b27dba0',
     'job_id': 'b3eff6c4-a63c-11e9-bfd4-2387a41a7f36',
     'artifact_name': 'f5e5de3ac8bc413385d94ff9203ed919'})
In [29]: s.wait_for_job('015bc7fa826c4f36b5ca2b4e8b27dba0')
{'status': 'Complete', 'starttime': '2019-07-15T12:53:34.483154', \
'endtime': '2019-07-15T12:53:42.266985', 'percent_complete': 100, \
'job_type': 'RunModel', 'loss': 0.4324471354484558, 'generations': 21, \
'dataset_names': ['pacman-cancer'], \
'artifact_names': ['f5e5de3ac8bc413385d94ff9203ed919'], \
'model_name': '1f871bd9b0b3405680a2bce0c3b2b226', 'job_error': ''}
Out[29]: (True, 'Job completed')
In [30]: s.download_artifact('f5e5de3ac8bc413385d94ff9203ed919')
Out[30]:
(True,
           Diagnosis prob_BENIGN prob_MALIGNANT
     0
                      0.816393
                                        0.183607
            BENIGN
     1
                      0.947398
                                        0.052602
            BENIGN
     2
            BENIGN
                      0.947646
                                        0.052354
     3
            BENIGN
                      0.947398
                                        0.052602
     4
         MALIGNANT
                      0.189687
                                        0.810313
     5
         MALIGNANT
                       0.256924
                                        0.743076
    . . .
         MALIGNANT
                       0.225788
                                        0.774212
     98
     99
         MALIGNANT
                        0.202293
                                        0.797707
     100
            BENIGN
                        0.816393
                                        0.183607
     101
          BENIGN
                        0.816393
                                        0.183607
```



```
[102 rows x 3 columns])
In [31]: s.analyze_model('1f871bd9b0b3405680a2bce0c3b2b226')
Out[31]:
   (True,
     {'job_name': '173bc091175c41c3a03d2cef9b4344fb',
      'job_id': 'eef624aa-a63c-11e9-94d1-2bccd93a7796',
      'artifact_name': '4d4684c1956844df8b412119637e890b'})
In [32]: s.wait_for_job('173bc091175c41c3a03d2cef9b4344fb')
{'status': 'Complete', 'starttime': '2019-07-14T08:40:24.885663', \
'endtime': '2019-07-14T08:40:26.360908', 'percent_complete': 100, \
'job_type': 'AnalyzeModel', 'loss': 0.20505395531654358, 'generations': 31, \
'dataset_names': None, 'artifact_names': ['4d4684c1956844df8b412119637e890b'], \
'model_name': '1f871bd9b0b3405680a2bce0c3b2b226', 'job_error': ''}
Out[32]: (True, 'Job completed')
In [33]: s.download_artifact('4d4684c1956844df8b412119637e890b')
Out [33]:
(True, Diagnosis = BENIGN
                                     0.143460
      Single Epithelial Cell Size 0.141226
     Bland Chromatin
                                     0.114085
     Normal Nucleoli
                                     0.112659
     Mitoses
                                     0.100054
     Marginal Adhesion
                                     0.091280
     Uniformity of Cell Size
                                     0.070170
     Uniformity of Cell Shape
                                     0.066997
     Code
                                     0.058587
     Clump Thickness
                                     0.053780
    Diagnosis = MALIGNANT
                                     0.046033
     Bare Nuclei
                                     0.001669
     dtype: float64)
In [34]: s.analyze_predictions('1f871bd9b0b3405680a2bce0c3b2b226', \
         'pacman-cancertest')
Out[34]:
    (True,
     {'job_name': 'b8e93f64e00f41819886cccbd2cad488',
      'job_id': 'f9f081a0-a729-11e9-8035-c3c7e048165a',
      'artifact_name': 'cbb36947ba5e4511846a80668207c77c'})
In [35]: s.wait_for_job('b8e93f64e00f41819886cccbd2cad488')
{'status': 'Running', 'starttime': '2019-07-15T12:57:14.029058', \
'endtime': None, 'percent_complete': 0, 'job_type': 'AnalyzePredictions', \
```



```
'loss': 0.4324471354484558, 'generations': 21, 'dataset_names': None, \
'artifact_names': ['cbb36947ba5e4511846a80668207c77c'], \
'model_name': '9f3ffa24162448158bdb1c0f3fe8c21e', 'job_error': ''}
{'status': 'Running', 'starttime': '2019-07-15T12:57:14.029058', \
'endtime': None, 'percent_complete': 0, 'job_type': 'AnalyzePredictions', \
'loss': 0.4324471354484558, 'generations': 21, 'dataset_names': None, \
'artifact names': ['cbb36947ba5e4511846a80668207c77c'], \
'model_name': '9f3ffa24162448158bdb1c0f3fe8c21e', 'job_error': ''}
{'status': 'Complete', 'starttime': '2019-07-15T12:57:14.029058', \
'endtime': '2019-07-15T13:05:26.715496', 'percent_complete': 100, \
'job_type': 'AnalyzePredictions', 'loss': 0.4324471354484558, 'generations': 21, \
'dataset_names': None, 'artifact_names': ['cbb36947ba5e4511846a80668207c77c'], \
'model_name': '9f3ffa24162448158bdb1c0f3fe8c21e', 'job_error': ''}
Out[35]: (True, 'Job completed')
In [36]: s.download_artifact('cbb36947ba5e4511846a80668207c77c')
Out [36]:
(True,
Code_shap Clump Thickness_shap Uniformity of Cell Size_shap
                                                                 Uniformity of ... \
     0
         0.001568
                                  0.029792
                                                                 0.068253
    1
         0.000836
                                  0.039068
                                                                 0.052454
     2.
         0.000584
                                  0.039056
                                                                 0.052201
     3
         0.002181
                                  0.037539
                                                                 0.052797
     4
         -0.001198
                                  0.012781
                                                                 0.124094
     98
         -0.001878
                                  0.000495
                                                                 0.131678
                                                                 0.141285
     99 -0.000813
                                  0.000495
    100 0.001568
                                 0.029792
                                                                 0.068253
     101 0.001568
                                  0.029792
                                                                 0.068253
     [102 rows x 13 columns])
In [39]: s.resume\_training\_model('1f871bd9b0b3405680a2bce0c3b2b226', \
'pacman-cancer', max_train_time='00:01')
Out[39]:
    (True,
     {'job_name': '30ba83c61814459f95b668a8316e647f',
      'job_id': '66e3ef92-a63d-11e9-9792-a38b3282194d',
      'model_name': '1f871bd9b0b3405680a2bce0c3b2b226'})
In [40]: s.wait_for_job('30ba83c61814459f95b668a8316e647f')
{'status': 'Running', 'starttime': '2019-07-14T08:43:46.090825', 'endtime': None, \
'percent_complete': 0, 'job_type': 'UpdateModel', 'loss': 0.20505395531654358, \
'generations': 31, 'dataset_names': ['pacman-cancer'], 'artifact_names': None, \
```



```
'model_name': '1f871bd9b0b3405680a2bce0c3b2b226', 'job_error': ''}
{'status': 'Running', 'starttime': '2019-07-14T08:43:46.090825', 'endtime': None, \
'percent_complete': 13, 'job_type': 'UpdateModel', 'loss': 0.20505395531654358, \
'generations': 40, 'dataset_names': ['pacman-cancer'], 'artifact_names': None, \
'model_name': '1f871bd9b0b3405680a2bce0c3b2b226', 'job_error': ''}
{'status': 'Running', 'starttime': '2019-07-14T08:43:46.090825', 'endtime': None, \
'percent_complete': 35, 'job_type': 'UpdateModel', 'loss': 0.20505395531654358, \
'generations': 43, 'dataset_names': ['pacman-cancer'], 'artifact_names': None, \
'model_name': '1f871bd9b0b3405680a2bce0c3b2b226', 'job_error': ''}
{'status': 'Running', 'starttime': '2019-07-14T08:43:46.090825', 'endtime': None, \
'percent_complete': 61, 'job_type': 'UpdateModel', 'loss': 0.20505395531654358, \
'generations': 47, 'dataset_names': ['pacman-cancer'], 'artifact_names': None, \
'model_name': '1f871bd9b0b3405680a2bce0c3b2b226', 'job_error': ''}
{'status': 'Running', 'starttime': '2019-07-14T08:43:46.090825', 'endtime': None, \
'percent_complete': 95, 'job_type': 'UpdateModel', 'loss': 0.20505395531654358, \
'generations': 52, 'dataset_names': ['pacman-cancer'], 'artifact_names': None, \
'model_name': '1f871bd9b0b3405680a2bce0c3b2b226', 'job_error': ''}
{'status': 'Complete', 'starttime': '2019-07-14T08:43:46.090825', 'endtime': \
'2019-07-14T08:46:07.389587', 'percent_complete': 100, 'job_type': 'UpdateModel', \
'loss': 0.20505395531654358, 'generations': 52, 'dataset_names': ['pacman-cancer'], \
'artifact_names': None, 'model_name': '1f871bd9b0b3405680a2bce0c3b2b226', \
'job error': ''}
Out[40]: (True, 'Job completed')
In [41]: s.run_model('pacman-cancer', '1f871bd9b0b3405680a2bce0c3b2b226')
Out[41]:
    (True,
     {'job_name': 'cc623240c5b1453eb73cdcdc93777f55',
      'job_id': 'c4380d18-a63d-11e9-85df-2337b425c294',
      'artifact_name': '48d74e4277944412ac032879fc23c5ba'})
In [45]: s.download_artifact('48d74e4277944412ac032879fc23c5ba')
Out[45]:
(True,
           Diagnosis prob_BENIGN prob_MALIGNANT
     0
            BENIGN
                      0.920420
                                        0.079580
     1
         MALIGNANT
                       0.461758
                                        0.538242
    1 1 1
     597
            BENIGN
                       0.834559
                                        0.165441
     598
            BENIGN
                       0.949016
                                        0.050984
    [599 \text{ rows x 3 columns}])
```



## Unsupervised modeling example

```
In [47]: s.upload_dataset('sets/pulsars.csv', 'pacman-pulsars')
Out[47]: (True, {'dataset_name': 'pacman-pulsars'})
In [48]: s.analyze_data('pacman-pulsars')
Out [48]:
(True,
 {'job name': '294ed354f3484b2ebebe658033284128',
  'job_id': '80193b9c-a63e-11e9-85df-fba3623db3bc',
  'artifact_name': 'b3d8dacf77fc409b9dd48030b2dda07b'})
In [49]: s.clean_data('pacman-pulsars')
Out [49]:
(True,
 {'job_name': '454f781b2a02403ea74d915a4b6b530c',
  'job_id': '8763baf8-a63e-11e9-900b-4335c8f0f324',
  'artifact_name': 'b9820a205bf140b1ae24bc6b1b133d2d',
  'artifact_id': '876560d8-a63e-11e9-900b-7f0b539a80a9'})
In [50]: s.create model('pacman-pulsars',\
fit_profile_name='b9820a205bf140b1ae24bc6b1b133d2d')
Out [50]:
(True,
 {'job name': '2a9a1e55f8e34ddd828326ecee2b42f4',
  'job_id': '97a92592-a63e-11e9-a08f-cf415e11311a',
  'model_name': 'fa1b82cab28c46cdac3b44c8e8bc1265'})
In [51]: s.wait_for_job('2a9a1e55f8e34ddd828326ecee2b42f4')
{'status': 'Running', 'starttime': '2019-07-14T08:52:17.412985', 'endtime': None, \
 'percent_complete': 0, 'job_type': 'TrainModel', 'loss': None, 'generations': 0, \
 'dataset_names': ['pacman-pulsars'], 'artifact_names': None, \
 'model_name': 'fa1b82cab28c46cdac3b44c8e8bc1265', 'job_error': ''}
{'status': 'Running', 'starttime': '2019-07-14T08:52:17.412985', 'endtime': None,\
 'percent_complete': 0, 'job_type': 'TrainModel', 'loss': None, 'generations': 0, \
  'dataset_names': ['pacman-pulsars'], 'artifact_names': None,\
   'model_name': 'fa1b82cab28c46cdac3b44c8e8bc1265', 'job_error': ''}
{'status': 'Complete', 'starttime': '2019-07-14T08:52:17.412985', \
'endtime': '2019-07-14T08:53:53.736499', 'percent_complete': 100, \
'job_type': 'TrainModel', 'loss': None, 'generations': 0, \setminus
 'dataset_names': ['pacman-pulsars'], 'artifact_names': None, \
 'model_name': 'fa1b82cab28c46cdac3b44c8e8bc1265', 'job_error': ''}
Out[51]: (True, 'Job completed')
```



```
In [53]: s.run_model('pacman-pulsars', 'falb82cab28c46cdac3b44c8e8bc1265')
Out [53]:
(True,
 {'job_name': 'c72ef79400014a3fb9e0a5821adf0826',
  'job_id': '0b58fb52-a63f-11e9-9f7b-53a24e4e1e2a',
  'artifact_name': '05380df9e93c4650ab6f7e5e67e23f72'})
In [54]: s.wait_for_job('c72ef79400014a3fb9e0a5821adf0826')
{'status': 'Complete', 'starttime': '2019-07-14T08:55:31.501449', \
'endtime': '2019-07-14T08:55:34.679054', 'percent_complete': 100, \
'job_type': 'RunModel', 'loss': None, 'generations': 0, \
'dataset_names': ['pacman-pulsars'], \
'artifact_names': ['05380df9e93c4650ab6f7e5e67e23f72'], \
'model_name': 'fa1b82cab28c46cdac3b44c8e8bc1265', 'job_error': ''}
Out[54]: (True, 'Job completed')
In [55]: s.download_artifact('05380df9e93c4650ab6f7e5e67e23f72')
Out [55]:
(True,
{'filename': '/var/folders/wc/w7ktf3392_s7c6t2djlhyb9/T/artifact-6ac2m9yp.csv'})
n [17]: s.analyze model('b7d28ee423d4430fafa51a017be827ac')
Out [17]:
(True,
 {'job_name': 'b36800c1a47f4c559c231d155a12fd85',
  'job_id': '8c5dfd70-a8f5-11e9-88ca-173db83c1239',
  'artifact_name': 'a417260dfabc409a8204fa1ceeae112f'})
In [18]: s.wait_for_job('b36800c1a47f4c559c231d155a12fd85')
{'status': 'Complete', 'starttime': '2019-07-17T19:46:58.691115',\
 'endtime': '2019-07-17T19:47:02.152927', 'percent_complete': 100,\
  'job_type': 'AnalyzeModel', 'loss': None, 'generations': 0,\
   'dataset_names': None, 'artifact_names': ['a417260dfabc409a8204fa1ceeae112f'], \
    'model_name': 'b7d28ee423d4430fafa51a017be827ac', 'job_error': ''}
Out[18]: (True, 'Job completed')
In [19]: s.download_artifact('a417260dfabc409a8204fa1ceeae112f')
Out[19]:
(True, kurt_dmsnr
                      0.0
               0.0
skew_dmsnr
skew_profile
                0.0
kurt_profile
                0.0
 class
                0.0
 std_profile
                 0.0
 mean_dmsnr
                 0.0
```



```
mean profile
                0.0
 std_dmsnr
                 0.0
dtype: float64)
In [22]: s.upload_dataset('sets/pulsars_predict.csv', 'pulsars-test')\
  # Need to trim original dataset to have fewer than 500 rows.
Out[22]: (True, {'dataset_name': 'pulsars-test'})
In [23]: s.clean_data('pulsars-test', model_name='b7d28ee423d4430fafa51a017be827ac')
Out [23]:
(True,
 {'job_name': '8324da8dea734455a73daeeddd3e0b5f',
  'job_id': 'fc2a4974-a8f5-11e9-9074-7fa762e40db7',
  'profile_name': 'bccde471e8514ef59b0b106fa7af6be9',
  'profile_id': 'fc2c0e76-a8f5-11e9-9074-13d39d00d68d'})
In [24]: s.analyze_predictions ('b7d28ee423d4430fafa51a017be827ac', 'pulsars-test')
Out [24]:
(True,
{'job_name': 'def26ef5be3a4d5b822542fd125c8600',
  'job id': '12995024-a8f6-11e9-89b8-cf7654542d20',
  'artifact name': 'd726a45761a1431d8bbe381c4f4f2782'})
In [25]: s.wait_for_job('def26ef5be3a4d5b822542fd125c8600')
{'status': 'Complete', 'starttime': '2019-07-17T19:50:43.895873',\
 'endtime': '2019-07-17T19:50:49.098101', 'percent_complete': 100,\
  'job_type': 'AnalyzePredictions', 'loss': None, 'generations': 0,\
   'dataset_names': None, 'artifact_names': ['d726a45761a1431d8bbe381c4f4f2782'],\
    'model_name': 'b7d28ee423d4430fafa51a017be827ac', 'job_error': ''}
Out[25]: (True, 'Job completed')
In [26]: s.download_artifact('d726a45761a1431d8bbe381c4f4f2782')
Out [26]:
(True,
    std_dmsnr_shap mean_dmsnr_shap kurt_dmsnr_shap mean_profile_shap
 0
               0.0
                                0.0
                                                  0.0
                                                                      0.0
 1
               0.0
                                0.0
                                                  0.0
                                                                      0.0
 2
               0.0
                                0.0
                                                  0.0
                                                                      0.0
 3
               0.0
                                0.0
                                                  0.0
                                                                      0.0
 4
               0.0
                                0.0
                                                  0.0
                                                                      0.0
 5
               0.0
                                0.0
                                                  0.0
                                                                      0.0
                                                                      0.0
 6
               0.0
                                0.0
                                                  0.0
 7
               0.0
                                0.0
                                                  0.0
                                                                      0.0
 8
               0.0
                                0.0
                                                  0.0
                                                                      0.0
 [9 rows x 12 columns])
```



# NBM modeling example

```
In [13]: s.upload_dataset('sets/SmokyT025_full_raw.csv', 'pacman-smoky')
Out[13]: (True, {'dataset_name': 'pacman-smoky'})
In [14]: s.analyze_data('pacman-smoky')
Out[14]:
(True,
 {'job_name': 'af7d01c94e774b3aa5648bf675cf990f',
  'job id': 'fc4e1f30-a895-11e9-be40-8b36979e9d84',
  'artifact_name': 'c267aa0daa43407fbb59f44c3644d2b8'})
In [15]: s.clean_data('pacman-smoky', index='timestamp')
Out[15]:
(True,
 {'job_name': '1950b29aef9a42afb1eb17125cea38a5',
  'job_id': '567ddbe4-a896-11e9-b2c6-c3499b5ebff0',
  'profile_name': '1700ee4ecb854ba8977a06f85efd1644',
  'profile_id': '567f38e0-a896-11e9-b2c6-770cfb81b384'})
In [18]: s.create model('pacman-smoky', \
 fit_profile_name='1700ee4ecb854ba8977a06f85efd1644',\
 max_train_time='00:02', recurrent=False, failure_dates=['08/23/2015'], nbm=True)
Out[18]:
(True,
 {'job_name': '23ce4b284df14f0ca14c0a49f63806fc',
  'job_id': '8b91e172-a896-11e9-be40-df999689405b',
  'model_name': '7e080e51cc15408492d6136e07df2a63'})
In [19]: s.wait_for_job('23ce4b284df14f0ca14c0a49f63806fc')
{'status': 'Running', 'starttime': '2019-07-17T08:26:55.165137', 'endtime': None,\
'percent_complete': 0, 'job_type': 'TrainModel', 'loss': None, 'generations': 0, \
  'dataset_names': ['pacman-smoky'], 'artifact_names': None,\
   'model_name': '7e080e51cc15408492d6136e07df2a63', 'job_error': ''}
{'status': 'Running', 'starttime': '2019-07-17T08:26:55.165137', 'endtime': None,\
 'percent_complete': 0, 'job_type': 'TrainModel', 'loss': None, 'generations': 0, \
  'dataset_names': ['pacman-smoky'], 'artifact_names': None,\
   'model name': '7e080e51cc15408492d6136e07df2a63', 'job error': ''}
{'status': 'Complete', 'starttime': '2019-07-17T08:26:55.165137',\
 'endtime': '2019-07-17T08:29:11.943704', 'percent_complete': 100,\
  'job_type': 'TrainModel', 'loss': 0.054839795631057814, 'generations': 1,\
   'dataset_names': ['pacman-smoky'], 'artifact_names': None,\
    'model_name': '7e080e51cc15408492d6136e07df2a63', 'job_error': ''}
Out[19]: (True, 'Job completed')
```



```
In [21]: s.run_model('pacman-smoky', '7e080e51cc15408492d6136e07df2a63')
Out [21]:
(True,
{'job_name': '268b0fb872054ec58c0344053625e69c',
  'job_id': '1351b15a-a897-11e9-a06a-2b396d284046',
  'artifact_name': '4fd95f6ec4c442e586b941f7e2656dbf'})
In [24]: s.wait_for_job('268b0fb872054ec58c0344053625e69c')
{'status': 'Complete', 'starttime': '2019-07-17T08:30:42.913437',\
 'endtime': '2019-07-17T08:31:08.226901', 'percent_complete': 100,\
  'job_type': 'RunModel', 'loss': 0.054839795631057814, 'generations': 1,\
   'dataset_names': ['pacman-smoky'], \
   'artifact names': ['4fd95f6ec4c442e586b941f7e2656dbf'],
   'model_name': '7e080e51cc15408492d6136e07df2a63', 'job_error': ''}
Out[24]: (True, 'Job completed')
In [25]: s.download artifact('4fd95f6ec4c442e586b941f7e2656dbf')
Out [25]:
(True,
                 Risk
 0
      0.000000
      0.000000
1
 2
       0.000000
 3
      0.000000
       0.000000
 4
51401 3.175640
51402 3.153954
 51403 3.131895
51404 3.109017
 51405 3.087970
[51406 rows x 1 columns])
In [27]: s.upload_dataset('sets/SmokyT025_test_raw.csv', 'pacman-smokytest')
Out[27]: (True, {'dataset_name': 'pacman-smokytest'})
In [28]: s.clean_data('pacman-smokytest', \
model name='7e080e51cc15408492d6136e07df2a63')
Out [28]:
(True,
 {'job_name': '229b24bfc46d4601aef1ce8b017e4ff7',
  'job_id': 'bd369cca-a898-11e9-8ea7-2bc9c8ab84c4',
  'profile_name': 'b4f5cc08f980442f983a8137a756dc79',
  'profile id': 'bd38ac40-a898-11e9-8ea7-9fb23d6a2658'})
```



```
In [29]: s.run_model('pacman-smokytest', '7e080e51cc15408492d6136e07df2a63')
Out [29]:
(True,
 {'job_name': '35d9cd3b4a3b45a99eb8dc089ef9b152',
  'job_id': 'c7d0d5ce-a898-11e9-bfef-379c562f822e',
  'artifact_name': 'f750090766f14e82b3713e3f3ef9bcb5'})
In [30]: s.wait_for_job('35d9cd3b4a3b45a99eb8dc089ef9b152')
{'status': 'Complete', 'starttime': '2019-07-17T08:42:55.232948',\
 'endtime': '2019-07-17T08:43:01.540698', 'percent_complete': 100,\
  'job_type': 'RunModel', 'loss': 0.054839795631057814, 'generations': 1,\
   'dataset_names': ['pacman-smokytest'], \
   'artifact_names': ['f750090766f14e82b3713e3f3ef9bcb5'], \
   'model_name': '7e080e51cc15408492d6136e07df2a63', 'job_error': ''}
Out[30]: (True, 'Job completed')
In [31]: s.download_artifact('f750090766f14e82b3713e3f3ef9bcb5')
Out[31]:
(True,
                  Risk
Ω
      0.000000
       0.000000
1
 2
      0.000000
 3
       0.000000
. . .
17276 3.121538
17277 3.098760
17278 3.077858
 [17279 rows x 1 columns])
In [32]: s.analyze_model('7e080e51cc15408492d6136e07df2a63')
Out [32]:
(True,
 {'job_name': 'e54bfd3d620d44b083f5ea65dda12aec',
  'job_id': '605a6300-a899-11e9-bfef-6fedf02afe89',
  'artifact name': 'd8ea781de54c4303ad49e17f0d208db3'})
In [33]: s.wait_for_job('e54bfd3d620d44b083f5ea65dda12aec')
{'status': 'Running', 'starttime': '2019-07-17T08:47:11.149655', 'endtime': None,\
 'percent_complete': 0, 'job_type': 'AnalyzeModel', 'loss': 0.054839795631057814,\
  'generations': 1, 'dataset_names': None, \
  'artifact_names': ['d8ea781de54c4303ad49e17f0d208db3'], \
  'model_name': '7e080e51cc15408492d6136e07df2a63', 'job_error': ''}
{'status': 'Complete', 'starttime': '2019-07-17T08:47:11.149655',\
 'endtime': '2019-07-17T08:47:27.795439', 'percent_complete': 100,\
```



```
'job_type': 'AnalyzeModel', 'loss': 0.054839795631057814, 'generations': 1,\
   'dataset_names': None, 'artifact_names': ['d8ea781de54c4303ad49e17f0d208db3'],\
    'model_name': '7e080e51cc15408492d6136e07df2a63', 'job_error': ''}
Out[33]: (True, 'Job completed')
In [34]: s.download_artifact('d8ea781de54c4303ad49e17f0d208db3')
Out[34]:
(True, ICA
                           0.290290
PSETKW
                     0.095161
GENTMPSLIPDEGC
                     0.075301
AMBTMPDEGC
                     0.062500
NACTMPDEGC
                     0.040816
 XFMRTMPPHCDEGC
                     0.034540
                    0.032311
GENTMPPHADEGC
YAWDIRDEG
                    0.031473
 GENTMPPHCDEGC
                     0.027232
WDRELDEG
                     0.026735
HYDRPRESBAR
                     0.026336
FREQHZ
                     0.022709
XFMRTMPPHADEGC
                    0.020440
GENTMPPHBDEGC
                     0.018139
BRGTMPGENNDEDEGC
                    0.017267
WD10MDEG
                     0.016984
VANV
                     0.015967
ΡF
                     0.012871
                     0.012038
TMPCTRLTOPDEGC
IBA
                     0.011195
TMPSPINNERDEGC
                     0.011191
                     0.010891
BLDANGDEG
WS10MMPS
                     0.010032
VBNV
                     0.009880
VCNV
                     0.008443
XFMRTMPPHBDEGC
                     0.007716
OILTMPGBXDEGC
                     0.007068
GENSPDRPM
                     0.006877
 TMPCTRLHUBDEGC
                     0.006240
PEXPKW
                     0.005889
WSMPS
                     0.005433
BRGTMPGBXADEGC
                     0.004322
 QKVAR
                     0.004213
                     0.003860
HYDRTMPDEGC
P10MACTKW
                     0.003125
 RTRSPDRPM
                     0.002313
                     0.002204
 IAA
 dtype: float64)
```



```
In [39]: s.upload_dataset('sets/SmokyT025_test_small.csv', 'pacman-smokytestsmall')
Out[39]: (True, {'dataset_name': 'pacman-smokytestsmall'})
In [40]: s.clean_data('pacman-smokytestsmall', \
  model_name='7e080e51cc15408492d6136e07df2a63')
Out[40]:
(True,
 {'job_name': 'a9a68568b4f44fe19661f0b5d4e8822d',
  'job_id': 'f68898c4-a899-11e9-9e51-2b93b6669bbd',
  'profile_name': '5c90c01159684d6491d956f035651c6f',
  'profile_id': 'f68a2a5e-a899-11e9-9e51-5bba6c3dc2d1'})
In [41]: s.analyze_predictions ('7e080e51cc15408492d6136e07df2a63',\
   'pacman-smokytestsmall')
Out[41]:
(True,
 {'job name': '04e5d614672a4e598276bd19a06f7df9',
  'job_id': 'fe1c8906-a899-11e9-8ea7-cfc85b8e9384',
  'artifact_name': '96fd4aa1b57043d69b658e72071c35c9'})
In [42]: s.wait_for_job('04e5d614672a4e598276bd19a06f7df9')
{'status': 'Running', 'starttime': '2019-07-17T08:51:35.824573', 'endtime': None,\
 'percent_complete': 0, 'job_type': 'AnalyzePredictions', \
 'loss': 0.054839795631057814, 'generations': 1, 'dataset_names': None, \
 'artifact_names': ['96fd4aa1b57043d69b658e72071c35c9'],\
 'model_name': '7e080e51cc15408492d6136e07df2a63', 'job_error': ''}
{'status': 'Running', 'starttime': '2019-07-17T08:51:35.824573', 'endtime': None,\
 'percent_complete': 0, 'job_type': 'AnalyzePredictions', \
 'loss': 0.054839795631057814, 'generations': 1, 'dataset_names': None, \
 'artifact_names': ['96fd4aa1b57043d69b658e72071c35c9'],\
 'model_name': '7e080e51cc15408492d6136e07df2a63', 'job_error': ''}
...In [47]: s.upload_dataset('sets/pulsars.csv', 'pacman-pulsars')
Out[47]: (True, {'dataset_name': 'pacman-pulsars'})
In [48]: s.analyze_data('pacman-pulsars')
Out [48]:
(True,
 {'job name': '294ed354f3484b2ebebe658033284128',
  'job_id': '80193b9c-a63e-11e9-85df-fba3623db3bc',
  'artifact_name': 'b3d8dacf77fc409b9dd48030b2dda07b'})
In [49]: s.clean_data('pacman-pulsars')
Out [49]:
(True,
 {'job_name': '454f781b2a02403ea74d915a4b6b530c',
```



```
'job_id': '8763baf8-a63e-11e9-900b-4335c8f0f324',
  'artifact_name': 'b9820a205bf140b1ae24bc6b1b133d2d',
  'artifact_id': '876560d8-a63e-11e9-900b-7f0b539a80a9'})
In [50]: s.create_model('pacman-pulsars', \
fit_profile_name='b9820a205bf140b1ae24bc6b1b133d2d')
Out [50]:
(True,
 {'job_name': '2a9a1e55f8e34ddd828326ecee2b42f4',
  'job_id': '97a92592-a63e-11e9-a08f-cf415e11311a',
  'model name': 'fa1b82cab28c46cdac3b44c8e8bc1265'})
In [51]: s.wait_for_job('2a9a1e55f8e34ddd828326ecee2b42f4')
{'status': 'Running', 'starttime': '2019-07-14T08:52:17.412985', 'endtime': None, \
'percent_complete': 0, 'job_type': 'TrainModel', 'loss': None, 'generations': 0, \
'dataset_names': ['pacman-pulsars'], 'artifact_names': None, \
'model_name': 'fa1b82cab28c46cdac3b44c8e8bc1265', 'job_error': ''}
{'status': 'Running', 'starttime': '2019-07-14T08:52:17.412985', 'endtime': None, \
'percent_complete': 0, 'job_type': 'TrainModel', 'loss': None, 'generations': 0, \
'dataset_names': ['pacman-pulsars'], 'artifact_names': None, \
'model_name': 'fa1b82cab28c46cdac3b44c8e8bc1265', 'job_error': ''}
{'status': 'Complete', 'starttime': '2019-07-14T08:52:17.412985', \
'endtime': '2019-07-14T08:53:53.736499', 'percent_complete': 100, \
'job_type': 'TrainModel', 'loss': None, 'generations': 0, \
'dataset_names': ['pacman-pulsars'], 'artifact_names': None, \
'model_name': 'fa1b82cab28c46cdac3b44c8e8bc1265', 'job_error': ''}
Out[51]: (True, 'Job completed')
In [53]: s.run_model('pacman-pulsars', 'fa1b82cab28c46cdac3b44c8e8bc1265')
Out [53]:
(True,
 {'job_name': 'c72ef79400014a3fb9e0a5821adf0826',
  'job_id': '0b58fb52-a63f-11e9-9f7b-53a24e4e1e2a',
  'artifact_name': '05380df9e93c4650ab6f7e5e67e23f72'})
In [54]: s.wait_for_job('c72ef79400014a3fb9e0a5821adf0826')
{'status': 'Complete', 'starttime': '2019-07-14T08:55:31.501449', \
'endtime': '2019-07-14T08:55:34.679054', 'percent_complete': 100, \
'job_type': 'RunModel', 'loss': None, 'generations': 0, \
'dataset_names': ['pacman-pulsars'], \
'artifact_names': ['05380df9e93c4650ab6f7e5e67e23f72'], \
'model_name': 'fa1b82cab28c46cdac3b44c8e8bc1265', 'job_error': ''}
Out[54]: (True, 'Job completed')
```



```
In [55]: s.download artifact('05380df9e93c4650ab6f7e5e67e23f72')
Out [55]:
(True,
{'filename': '/var/folders/wc/w7ktf3_h8br6t2djlhyb9/T/artifact-6ac2m9yp.csv'})
In [17]: s.analyze_model('b7d28ee423d4430fafa51a017be827ac')
Out [17]:
(True,
 {'job_name': 'b36800c1a47f4c559c231d155a12fd85',
  'job_id': '8c5dfd70-a8f5-11e9-88ca-173db83c1239',
  'artifact name': 'a417260dfabc409a8204fa1ceeae112f'})
In [18]: s.wait_for_job('b36800cla47f4c559c231d155a12fd85')
{'status': 'Complete', 'starttime': '2019-07-17T19:46:58.691115', \
'endtime': '2019-07-17T19:47:02.152927', 'percent_complete': 100, \
'job_type': 'AnalyzeModel', 'loss': None, 'generations': 0, \
'dataset_names': None, 'artifact_names': ['a417260dfabc409a8204fa1ceeae112f'], \
'model_name': 'b7d28ee423d4430fafa51a017be827ac', 'job_error': ''}
Out[18]: (True, 'Job completed')
In [19]: s.download_artifact('a417260dfabc409a8204fa1ceeae112f')
Out[19]:
                     0.0
(True, kurt_dmsnr
skew dmsnr 0.0
skew profile
                0.0
kurt_profile 0.0
class
               0.0
 std_profile
               0.0
mean_dmsnr
               0.0
mean_profile
               0.0
 std_dmsnr
            0.0
dtype: float64)
In [22]: s.upload_dataset('sets/pulsars_predict.csv', 'pulsars-test') \
 # Need to trim original dataset to have fewer than 500 rows.
Out[22]: (True, {'dataset_name': 'pulsars-test'})
In [23]: s.clean_data('pulsars-test', model_name='b7d28ee423d4430fafa51a017be827ac')
Out [23]:
(True,
 {'job_name': '8324da8dea734455a73daeeddd3e0b5f',
  'job_id': 'fc2a4974-a8f5-11e9-9074-7fa762e40db7',
  'profile_name': 'bccde471e8514ef59b0b106fa7af6be9',
  'profile_id': 'fc2c0e76-a8f5-11e9-9074-13d39d00d68d'})
```



```
In [24]: s.analyze_predictions ('b7d28ee423d4430fafa51a017be827ac', \
'pulsars-test')
Out [24]:
(True,
 {'job_name': 'def26ef5be3a4d5b822542fd125c8600',
  'job_id': '12995024-a8f6-11e9-89b8-cf7654542d20',
  'artifact name': 'd726a45761a1431d8bbe381c4f4f2782'})
In [25]: s.wait_for_job('def26ef5be3a4d5b822542fd125c8600')
{'status': 'Complete', 'starttime': '2019-07-17T19:50:43.895873', \
'endtime': '2019-07-17T19:50:49.098101', 'percent_complete': 100, \
'job_type': 'AnalyzePredictions', 'loss': None, 'generations': 0, \
'dataset_names': None, 'artifact_names': ['d726a45761a1431d8bbe381c4f4f2782'], \
'model_name': 'b7d28ee423d4430fafa51a017be827ac', 'job_error': ''}
Out[25]: (True, 'Job completed')
In [26]: s.download_artifact('d726a45761a1431d8bbe381c4f4f2782')
Out [26]:
(True,
    std_dmsnr_shap mean_dmsnr_shap kurt_dmsnr_shap mean_profile_shap ... \
 Λ
               0.0
                                0.0
                                                  0.0
                                                                     0.0
               0.0
                                0.0
                                                                     0.0
 1
                                                  0.0
 2
               0.0
                                0.0
                                                  0.0
                                                                     0.0
 3
                                                                     0.0
               0.0
                                0.0
                                                  0.0
 4
               0.0
                                0.0
                                                  0.0
                                                                     0.0
 5
               0.0
                                0.0
                                                  0.0
                                                                     0.0
 6
               0.0
                                0.0
                                                  0.0
                                                                     0.0
 7
               0.0
                                0.0
                                                  0.0
                                                                     0.0
 8
               0.0
                                0.0
                                                  0.0
                                                                     0.0
 [9 rows x 12 columns])
In [27]:
{'status': 'Complete', 'starttime': '2019-07-17T08:51:35.824573', \
 'endtime': '2019-07-17T08:55:05.619098', 'percent_complete': 100, \
 'job_type': 'AnalyzePredictions', 'loss': 0.054839795631057814, \
 'generations': 1, 'dataset_names': None,\
 'artifact names': ['96fd4aa1b57043d69b658e72071c35c9'],\
 'model_name': '7e080e51cc15408492d6136e07df2a63', 'job_error': ''}
Out[42]: (True, 'Job completed')
In [43]: s.download_artifact('96fd4aa1b57043d69b658e72071c35c9')
Out [43]:
(True,
      GENSPDRPM_shap VANV_shap PSETKW_shap P10MACTKW_shap ... \
 0
           20.060011 5.785717
                                   0.000000
                                                    -1.762059
           -1.002102 -4.857683
                                   -1.833142
                                                     0.000000
 1
```



2	0.641536	0.015167	0.000000	0.000000	
3	0.000000	-1.827971	0.00000	0.000000	
4	-5.824179	6.567310	10.847957	-5.231445	
5	-16.634360	0.000000	-8.437045	-9.929700	
				• • •	
396	-1.932885	-1.780289	-1.361495	0.00000	
397	5.469176	-8.030168	38.908720	-1.336821	
398	-0.598940	-8.100026	57.731217	17.159244	
[399 rows x 39 columns])					

# **Contact Support**

The following methods enable you to research issues, create a support ticket, or contact SparkCognition:

- Use the Darwin support portal Read Frequently Asked Questions (FAQ), download documentation, or log your issue.
- **Email Support** Send email to darwin\_support@sparkcognition.com.
- **Phone Support** The SparkCognition support line is +1-512-400-2001.

# **Revision Table**

Version	Date	Notes
v 1.6	16-Jan-2019	New endpoints:
		<ul> <li>DarwinSdk.display_population</li> </ul>
		<ul> <li>DarwinSdk.delete_all_artifacts</li> </ul>
		Updated endpoints:
		• DarwinSdk.analyze_data
		<ul> <li>DarwinSdk.download_artifact</li> </ul>
		<ul> <li>DarwinSdk.create_model</li> </ul>
		• DarwinSdk.clean_data
v 1.6.1	06-Feb-2019	Fixed issues only. See Release Notes. Added on-prem installation notes.
v 1.6.2	22-Mar-2019	New endpoints:
		• DarwinSdk.get_info
		• DarwinSdk.help
		Added Setup Users section.
		On-prem SDK users need to add port 8000 to the URL.



Version	Date	Notes
v 1.43.0	16-May-2019	Major change to version number to facilitate independent releases
		of the API
		New endpoints:
		<ul><li>DarwinSdk.disable_ssl_cert_check</li></ul>
		<ul> <li>DarwinSdk.enable_ssl_cert_check</li> </ul>
		<ul><li>DarwinSdk.get_sdk_version</li></ul>
		Updated endpoints:
		• DarwinSdk.create_model
v 1.44.0	22-Jul-2019	Updated modeling examples for Supervised, Unsupervised, and
		NBM
		Added Analyze Training Data step to basic workflow
		Updated endpoints:
		DarwinSdk.create_model: Added forecast_horizon,
		class_weights, cv_kfold, fit_profile_name
v 1.44.1	24-Sep-2019	Removed model_type parameter from run_model()
		Removed impute parameter from clean_data()
v 1.45.0	18-Dec-2019	Added model_type parameter to run_model()
		Added impute parameter to clean_data()
		Added fitness_fn_name to create_model()
		Limited to 4000 columns in dataset