

# Darwin<sup>TM</sup> Python SDK Manual

## A SparkCognition<sup>TM</sup> Education Document

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# **About this guide**

This manual describes using the Darwin<sup>TM</sup> 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, 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 (...).

## **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<sup>®</sup> tool that automates model building processes to solve specific problems. This tool enhances data scientist potential because it automates various tasks that are often manually performed. These tasks include data cleaning, latent relationship extraction, and optimal model determination. Darwin promotes rapid and accurate feature generation through both automated windowing and risk generation. Darwin quickly creates highly-accurate, dynamic models using both supervised and unsupervised learning methods.

The general workflow for simple modeling includes:

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

See the SDK example for a modeling example.

For additional information on Darwin, see the white paper titled: *Darwin - A Neurogenesis Platform*.

## **Accessing the API**

This document describes the 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, see the SparkCognition Darwin API Guide.

#### Notes:

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

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

## **Darwin SDK interface**

## Connect to the Darwin interface

1. Obtain an api key.

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

'RsJ74ZS5AvwznbHh0AfVSgrchhS9KxACDy3jefaQnxb9f6QTSDBFmhnGa0c0IWtNAIFRAG9ToOTpi0mnEo3zFA'

## 2. Register the api key.

The purpose of this method is to set a password for an api\_key. Each api\_key is synonymous with a service. This method must be invoked once for each api\_key to establish a password for that key.

#### **Notes:**

- After successful registration, the service uses auth\_login() to login as a service.
- In version 1, the password cannot be changed.

### Example

```
>>> from amb_sdk.sdk import DarwinSdk
>>> s = DarwinSdk()
>>> s.auth_register('asdf', 'iRgwut4kGs0ymULiuKtMd0WFvBYLMWSj16q2uysQeteqH9ssc+\
EETUvcysnPojRpfycLVHa2IlN1IlrfEk1YMA')
(True,'Bearer eyJ0eXAi0iJKV1QiLCJhbGci0iJIUzI1NiJ9.eyJleHAi0jE1MTU1MzM4NjEsImlh\
dCI6MTUxNTUzMDI2MS ... F56xZQiBT-89nrRz1nIXD5LfawHIj_MlUHQqM36vU')
```

## 3. Login.

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

The following explains how to log in as a *service*.

#### **Notes:**

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

```
>>> s.auth_login('asdf', 'iRgwut4kGs0ymULiuKtMd0WFvBYLMWSj16q2uysQeteqH9ssc+EET\
UvcysnPojRpfycLVHa2IlN1IlrfEk1YMA')
(True,'Bearer eyJ0eXAi0iJKV1QiLCJhbGci0iJIUzI1NiJ9.eyJleHAi0jE1MTU1MzQxNzIsImlh\
dCI6MTUxNTUzMD ... UQQfoXqYFKJSoRXXDNPE985-a08cE6_o')
```

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

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

### **Example**

```
>>> s.auth_register_user('atestuser', 'apassword')
(True,
    'Bearer eyJhbGciOiJIUzI1NiIsInR5cCI6IkpXVCJ9.eyJqdGkiOiJkNjY0MmJjOC1iMmU5LTQxO\
DctODF1NS00YjI2MD ... 5zMp_1FfxU')
>>> s.auth_login_user('atestuser', 'apassword')
(True,
    'Bearer eyJhbGciOiJIUzI1NiIsInR5cCI6IkpXVCJ9.eyJqdGkiOiI3NGYzYmUxZS0yOTlmLTRhN\
zMtODU5ZC01NGRmM2F ... u1zGCeCONA')
```

4. Verify the connection.

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

## **Darwin SDK methods**

## **URL Get/Set methods**

DarwinSdk.get url()

Get Darwin service url.

Parameters: None

**Returns:** 

(True, <url-string>)

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

Set Darwin service url and version.

#### Parameters:

- url URL to the Darwin service
- version (optional) defaults to 'v1'

#### **Returns:**

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

### **Example**

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

## **Authentication methods**

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

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

**Note**: In version 1, the password cannot be changed.

#### **Parameters:**

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

#### **Returns**:

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

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

The SDK remember the auth token for the AmkSdk object.

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

```
eyJ0eXAiOiJKV1QiLCJhbGciOiJIUzI1NiJ9.eyJleHAiOjE1MTU1Mz....')
```

DarwinSdk.auth login(password, api key)

Login as a service.

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

#### **Parameters:**

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

#### Returns:

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

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

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

## **Example**

```
In [6]: s.auth_login('asdf',
  'iRgwut4kGs0ymULiuKtMd0WFvBYLMWSj16q2uysQeteqH9ssc+EETUvcysnPojRpfyc\
LVHa2IlN1IlrfEk1YMA')
Out[6]:
(True,
  'Bearer
eyJ0eXAiOiJKV1QiLCJhbGciOiJIUzI1NiJ9.eyJleHAiOjE1MTU1MzQxN....')
```

DarwinSdk.auth register user(username, password)

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 new 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 AmkSdk object.

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

## **Example**

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

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 service level 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 AmkSdk object.

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

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

## Job status methods

DarwinSdk.job status(status=None)

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

#### **Parameters**:

- status (optional) If not specified, returns all jobs.
- Running (optional)
- Requested (optional)
- Complete (optional)
- Failed (optional)

#### **Returns:**

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

## **Example**

```
In [9]: s.job_status()
Out[9]:
(True,
 [{'accuracy': 0.6167756021022797,
   'dataset_name': 'unittest-cancer-testdataset',
   'generations': 17,
   'id': '4b5207ec-f318-40f1-9713-a04dcb566cd4',
   'model_name': 'unittest-cancer-model',
   'percent_complete': 100,
   'status': 'Complete'},
  {'accuracy': 0.6167756021022797,
   'dataset_name': None,
   'generations': 17,
   'id': '57e3deb6-2c69-44b8-bbbe-da54ed14098e',
   'model_name': 'unittest-cancer-model',
   'percent_complete': 100,
   'status': 'Running'}])
```

DarwinSdk.auth job status id(job id)

Get job status information for a job id.

#### **Parameters:**

• *job id* - The job ID.

### Returns:

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

```
In [6]: s.job_status_id('f403dc53-7330-4a8f-a0d5-690822c0349f')
Out[6]:
```

```
(True,
    {'dataset_name': 'unittest-cancer-dataset',
    'model_name': 'unittest-cancer-model',
    'percent_complete': 100,
    'status': 'Complete'})
```

## **Lookup methods**

DarwinSdk.lookup client()

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

Parameters: None

**Returns:** 

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

## Example

```
In [21]: s.lookup_client()
Out[21]:
(True,
    {'job_limit': None,
    'model_limit': None,
    'tier': 0,
    'upload_limit': None,
    'user_limit': None,
    'username': None})
```

## 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,
    'last_updated': '2018-01-08T16:41:36.860513',
    'mbytes': 0.02019977569580078,
    'name': 'unittest-cancer-dataset',
```

```
'sequential': None},
{'categorical': None,
  'imbalanced': None,
  'last_updated': '2018-01-08T16:43:29.031108',
  'mbytes': 0.0034189224243164062,
  'name': 'unittest-cancer-testdataset',
  'sequential': None}])
```

## DarwinSdk.lookup dataset name(dataset name)

Get a specific dataset's metadata.

#### Parameters:

• *dataset\_name* - The name of the dataset. The name of a dataset is established in the **upload\_dataset**() method.

#### **Returns**:

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

## Example

```
In [6]: s.lookup_dataset_name('unittest-cancer-dataset')
Out[6]:
(True,
    {'categorical': None,
    'imbalanced': None,
    'last_updated': '2018-01-08T16:41:36.860513',
    'mbytes': 0.02019977569580078,
    'sequential': None})
```

## 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>)
```

```
In [7]: s.lookup_model()
Out[7]:
(True,
  [{'accuracy': 0.5278194844722748,
    'last_trained': '2018-01-08T16:42:54.293154',
    'name': 'unittest-cancer-model',
    'parameters': {'target': 'Diagnosis'},
```

```
'trained_on': ['unittest-cancer-dataset']}])
```

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**

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>)
```

## **Example**

```
In [22]: s.lookup_tier()
Out[22]:
(True,
  [{'job_limit': None,
    'model_limit': None,
    'tier': 0,
    'upload_limit': None,
    'user_limit': None}])
```

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 name of the model

## **Returns:**

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

## **Example**

```
In [25]: s.lookup_tier_num(0)
Out[25]:
(True,
    {'job_limit': None,
    'model_limit': None,
    'tier': 0,
    'upload_limit': None,
    'user_limit': None})
```

## Datasets and artifact methods

DarwinSdk.upload dataset(dataset, dataset name=NONE)

Upload a dataset, model, or a figure.

#### **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>)
```

## Example

```
In [5]: s.upload_dataset('sets/cancer_train.csv', 'unittest-cancer-dataset')
Out[5]:
(True,
    {'dataset_name': 'unittest-cancer-dataset'})
```

DarwinSdk.delete dataset(dataset name)

Delete the named dataset.

#### Parameters:

• *dataset name* - The 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 artifact(artifact id)

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

- analyze\_data()
- analyze\_model()
- test model()
- run model()
- create risk info()

**Note**: The artifact for *analyze model()* is a png file.

#### Parameters:

• articfact id - The id of the artifact to be downloaded.

#### **Returns**:

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

#### Example run model() or prediction artifact

```
In [5]: s.download_artifact('6e4861de29424cb7ad09e467d1869c17')
Out[5]:
(True,
    {'filename': '/var/folders/9r/_wwbjsd17_5b74dww69mtg9h0000gp/T/artifact\
    -o9fz9gwe.csv'})

$ head /var/folders/9r/_wwbjsd17_5b74dww69mtg9h0000gp/T/artifact-o9fz9gwe.csv
x,actual,predicted
-0.5,0.0,0.0
0.5,0.0,0.0
0.5,0.0,0.0
1.5,0.0,0.0
1.5,0.0,0.0
```

## **Example test\_model() artifact**

```
In [8]: s.download_artifact('4c4b0833fc1b4edeb689cba86c94a58d')
Out[8]:
(True,
 ['Diagnosis',
    'BENIGN',
```

```
'BENIGN',
'BENIGN',
'MALIGNANT',
...
]
```

## **Example analyze\_data() artifact**

```
In [9]:
         s.download_artifact('97b0ea5d0c5a478b9a09202be61740fb')
Out[9]:
(True,
 [{'categories': None,
    col_name': 'Code',
   'col_type': 'numerical',
   'max': 8233704.0,
   'mean': 1044171.0667779633,
   'min': 61634.0,
   'num_categories': 557.0,
   'std': 414096.3687689267},
  {'categories': '1,2,3,4,5,6,7,8,9,10',
   'col name': ' Clump Thickness',
   'col_type': 'categorical',
   'max': None,
   'mean': None,
   'min': None,
   'num_categories': 10.0,
   'std': None},
  {'categories': '1,2,3,4,5,6,7,8,9,10',
   'col_name': ' Uniformity of Cell Size',
   'col_type': 'categorical',
   'max': None,
   'mean': None,
   'min': None,
   'num_categories': 10.0,
   'std': None},
)
```

### Example analyze model() or prediction artifact

```
In [5]: s.download_artifact('6e4861de29424cb7ad09e467d1869c17')
Out[5]:
(True,
   {'filename': '/var/folders/9r/_wwbjsd17_5b74dww69mtg9h0000gp/T/artifact-\
   0nq69erf.png'})
```

## **Example create\_risk\_info() artifact**

```
In [5]: s.download_artifact('6ec2c1bc6c1244ccb3b03f25ebac5850')
Out[5]:
(True,
    {'filename': '/var/folders/9r/_wwbjsd17_5b74dww69mtg9h0000gp/T/tmp\
```

```
jawjkj4f'})

$ head tmpjawjkj4f
risk
0.0
0.0
1.0
0.0
1.0
0.0
```

## DarwinSdk.delete artifact(artifact id)

Delete the artifact given its id.

## Parameters:

• artifact id - The id of the artifact to be deleted.

#### **Returns:**

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

## **Example**

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

## Modeling and analysis methods

DarwinSdk.create\_model(dataset name, parameters)

Create a model trained on the dataset identified by dataset\_name. The name of a model is specified in a parameter setting.

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

## Parameters:

- dataset name Name of the dataset to be used for training
- *parameters* json format parameters.

## Parameter descriptions:

- target: String denoting target prediction column in input data.
- *Model name*: The string identifier of the model to be trained.
- max\_train\_time: Sets the training time for the model in 'HH:MM' format. **Note**: This overrides any values set for max generation.

- max\_generation: Expected input/type: numeric. Sets the training time for the model in generations. If max train time is set, this parameter is ignored.
- *recurrent*: Expected input/type: *true/false*. Enables recurrent connections to be evolved in the model. This option can be useful for timeseries or sequential data.

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

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

ALIAS	DESCRIPTION	COMPLEXITY
'genetic'	Genetic Fill: Automatically determines the most appropriate fast imputation method using evolutionary methods.	Linear
'ffill'	(Default) Forward Fill: Propagate values forward from one example into the missing cell of the next example. Might be useful for timeseries data, but also applicable for both numerical and categorical data.	Linear Fast
'bfill'	Backward Fill: Propagate values backward from one example into the missing cell of the previous example. Might be useful for timeseries data, but also applicable for both numerical and categorical data.	Linear Fast
'mean'	Mean Fill: Computes the mean value of all non-missing examples in a column to fill in missing examples. The result may or might not be interpretable in terms of the input space for categorical variables.	Linear Fast
'median'	Median Fill: Computes the median value of all non-missing examples in a column to fill in missing examples. While the result is interpretable in terms of the input space for categorical variables, the approach might not be appropriate for non-ordinal data.	
'mode'	Mode Fill: Uses the most common value on a column-by-column basis to fill in missing examples. The result is interpretable for both numerical and categorical variables.	Linear Fast
'spline'	Spline Fill: Interpolation using a spline (piecewise function). Might be useful for timeseries or sequential data.	Linear Fast
'Linear'	Linear Interpolation Fill: Interpolation using a Linear function. Might be useful for timeseries or sequential data.	Linear Fast
'knn'	K-Nearest Neighbors Fill: Fills in missing values by averaging the cell values of the k nearest neighbors in the reduced feature space defined by all non-missing columns.	Polynomial Slow
'rmf'	Robust Matrix Factorization Fill: Computes low-rank matrices L (observations x rank), R (features x rank), and E where X is input data, and $X = LR^T + E$ .	Polynomial Slow
'mice'	Multiple Imputation by Chained Equations: First imputes missing values using <i>Forward Fill</i> . Then, column-by-column, missing values are reintroduced and regressed upon using the other (non-missing) columns. Continues iteratively.	Polynomial, Iterative Very Slow

- *drop*: Expected input/type: *true/false*. Enables automatic pruning of input columns based on different criteria such as amount of missing data, number of unique values, and standard deviation.

  Note: This automatically drops identifier columns (unique value for each sample) and columns that do not contain sufficient data to aid prediction.
- max\_int\_uniques: Expected input/type: integer. Threshold for automatic encoding of categorical variables. If a column contains at least max\_int\_uniques unique values, it is treated as categorical and

one hot encoded during preprocessing.

- 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.
- feature\_eng: Enables automatic feature generation. Identifies an appropriate time window and augments input with new features derived in the frequency and time domains.

  Note: Can only be applied to timeseries data. String aliases specify methods for window computation.

ALIAS	DESCRIPTION	
None	No feature generation will be applied.	
ʻmi'	Uses mutual information to estimate the window length.	
'auc'	(Default) Uses autocorrelation to estimate the window length	
'user'	User specified window length: see* window len*.	

- window\_len: Expected input/type: integer. User specified window length for feature generation.

  Note: This parameter is used only in the case that user is provided for the feature eng parameter.
- feature\_select: A number in [0,1] specifying the percentage of numerical features to maintain based on their dependency to the target. Ranks all features using mutual information and drops (1 feature select)% of the lowest-ranking features. Default is 1 (keep all features).
- *outlier*: A string alias that indicates the outlier detection to apply during preprocessing. **Note**: Outliers are removed and later filled using imputation.

ALIAS	DESCRIPTION		
None	( <b>Default</b> ) No outlier detection will be applied.		
'mad'	Uses Median Absolute Deviation to detect outliers.		
'perc'	Uses Percentile-based outlier detection.		
'isol'	Uses an Isolation Forest to detect outliers.		

- auto\_save\_per: Expected input/type: integer. Sets the checkpoint frequency. The model creation progress is recorded after every auto\_save\_per generations.
   Note: If the model is retrained, the model begins from the last recorded checkpoint. The model is automatically saved at the end of evolution.
- *imbalance*: Expected input/type: *true/false*. Enables automatic imbalance correction that selectively applies random oversampling, random undersampling, synthetic minority oversampling (SMOTE), or adaptive synthetic sampling (ADASYN) to the input data depending on problem characteristics.
- *clustering*: Expected input/type: *true/false*. Enables clustering for unsupervised problems. If false, detects outliers.
- *n\_clusters*: Expected input/type: *integer*. Specifies the number of clusters to be used if clustering is enabled.

**Note**: If this value is not provided, the number of clusters will be heuristically determined.

• *anomaly\_prior*: Expected input/type: \*between [0,1]. \*Significance level at which a point is defined as anomalous.

**Note**: This parameter is used only for unsupervised problems if clustering is disabled.

#### **Returns:**

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

## **Example**

```
In [10]: s.create_model('unittest-cancer-data', {"target": \
    "Diagnosis", "model_name": \
    "unittest-cancer-model"})
Out[10]:
(True,
    {'job_id': 'f5124576a4f34e5c9ab3499770455509',
    'model_name': 'unittest-cancer-model'})
```

## DarwinSdk.remove model(model name)

Remove a model named by model name.

#### **Parameters:**

• *model name* - Name of the model to be removed.

#### Returns:

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

#### Example

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

DarwinSdk.resume training model(dataset name, model name, parameters)

Resume training for a model on the dataset identified by *data name*.

#### Parameters:

- dataset name- Name of dataset to be used for training.
- model name Name of the model to be trained.
- parameters json format parameters.

## Payload:

```
{
    "max_train_time": "00:01"
}
```

#### Returns:

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

## **Example**

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

### DarwinSdk.analyze data(dataset name)

Analyze the dataset given its *name*.

### Parameters:

• dataset name - The name of the dataset to be analyzed.

## Returns:

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

### **Example**

```
In [10]: s.analyze_data('unittest-cancer-dataset')
Out [10]:
(True, {'artifact_id': '2c90c1dc402a4a6da37a139dfc2f7871', 'job_id': \
'7871ebb62ad1458da64d800bc73019de'})
```

## DarwinSdk.analyze model(model name)

Analyze the model given its model name.

## **Parameters**:

• *model name* - The name of the model to be analyzed.

#### Returns:

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

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

```
'4e59ffc425e047e1a3b872f1e7396976'})
```

DarwinSdk.test model(dataset name, model name)

Test the model given its name and a test dataset. Use *upload dataset()* to upload a test dataset.

#### Parameters:

- *dataset name* The name of the dataset to use for testing the model.
- *model name* The name of the model to be tested.

## Returns:

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

## Example

```
[In [8]: s.test_model('unittest-cancer-testdataset', 'unittest-cancer-model')
Out [8]:
(True, {'artifact_id': '6691c7eb6e404343be3882baaea88444', 'job_id': \
'a3a083c8d43647959a9cd81d43923125'})
```

DarwinSdk.**run model**(dataset name, model name)

Run the model given its name and a dataset to use. Use **upload\_dataset()** to upload a dataset.

#### **Parameters:**

- dataset name The name of a dataset to use for running the model
- model name The name of the model to run

#### Returns:

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

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

DarwinSdk.create\_risk\_info(failure\_dataset\_name, timeseries\_dataset\_name, parameters)

Create risk information given failure and timeseries data. Use *upload dataset()* to upload datasets.

**Notes concerning risk** - Risk is a value used in calculating future events. Risk is calculated using algorithms based on sliding time frames and associated historical data that projects forward in time to predict the likelihood of the event. The outcome of the calculations is that the likelihood of an event occurring within a particular time frame becomes available for use. Note that the risk values are dependent on the quality and extent of the historical data as well as the scope of the timeframe used for evaluation.

#### **Parameters:**

- failure dataset name The name of a failure dataset
- timeseries dataset name The name of a timeseries dataset
- parameters json format parameters.

## **Parameter Descriptions:**

- *shutdown\_column*: Name of the column in the risk data that denotes the beginning of the predicted event of interest.
- return\_column: Name of the column in the risk data that denotes the end of the predicted event and when all data can again be considered "normal".
- asset\_column: Name of the asset column in the risk data. This parameter is used when the datasets consist of multiple different assets.
- *lead\_time*: Lead time in seconds. This value is half width of the risk function that means. the risk index is 0 prior to 2\* *lead\_time* and increases to 1 at a failure time.
- Functional form: Shape of a risk function, includes:

step: Step functionlinear: Linear functionsigmoid: Sigmoid function

• exponential: Exponential function

#### **Returns:**

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

## Example

```
**NEED 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.wait for job(job id, time limit=600)

Synchronously wait for a job to complete, limited by *time limit* that defaults to 10 minutes (600 seconds).

#### **Parameters:**

- job id The id for the job
- time limit (optional) defaults to 600 seconds

#### Returns::

```
(True, None) or (False, )
```

## Reference

## SDK example

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

```
In [1]: from amb_sdk.sdk import DarwinSdk
In [2]: s = DarwinSdk()
In [3]: s.auth_login_user('username', 'password')
Out[3]:
```

```
(True,
 'Bearer eyJhbGciOiJIUzI1NiIsInR5cCI6IkpXVCJ9.eyJqdGkiOiI2ZDZkMTI3Mi0wZDAxLTR\
 hYmMtOWYwOC0xYWEwZmYxNDY2NjAiLCJpYXQiOjE1MTYyMTY1MTMsImV4cCI6MTUxNjIy\
MDExMywidHlwZSI6ImFjY2VzcyIsIm5iZiI6MTUxNjIxNjUxMywiaWRlbnRpdHki0iJ\
 jNTc2NzFjNC11NTAwLTExZTctOWY4ZS1iNzk2ODU2ZTcwMGYiLCJmcmVzaCI6ZmFsc\
 2V9.slh1mYPy_M7DqAok-tV1NT0kU4lAgKQoQHk6nYtetg4')
In [4]: s.upload_dataset('sets/cancer_train.csv')
Out[4]: (True, {'dataset_name': 'cancer_train.csv'})
In [5]: s.create_model('cancer_train.csv', {"target": "Diagnosis", \
"model_name": "cancer-model"})
Out[5]:
(True,
 {'job_id': '6ec2c1bc6c1244ccb3b03f25ebac5850', 'model_name': 'cancer-model'})
In [6]: s.wait_for_job('6ec2c1bc6c1244ccb3b03f25ebac5850')
{'status': 'Running', 'model_name': 'cancer-model', 'loss': \
0.4371463656425476, 'percent_complete': 33, 'starttime': \
'generations': 9}
{'status': 'Running', 'model_name': 'cancer-model', 'loss': \
0.41981565952301025, 'percent_complete': 58, 'starttime': \
'generations': 14}
{'status': 'Running', 'model_name': 'cancer-model', 'loss': \
0.4167576730251312, 'percent_complete': 77, 'starttime': \
'2018-01-17T13:17:30.967628', 'dataset_name': 'cancer_train.csv', \
'generations': 17}
{'status': 'Complete', 'model_name': 'cancer-model', 'loss': \
0.4150310754776001, 'percent_complete': 100, 'starttime': \
'2018-01-17T13:17:30.967628', 'dataset_name': 'cancer_train.csv', \
'generations': 19}
Out[6]: (True, 'Job completed')
In [7]: s.upload_dataset('sets/cancer_test.csv')
Out[7]: (True, {'dataset_name': 'cancer_test.csv'})
In [8]: s.test model('cancer train.csv', 'cancer-model')
Out[8]:
(True,
 { 'artifact_id': 'd27466dec9b548b5bd42f0295a23432e',
  'job id': 'a7469b2913504c80934ca2cd72c7fa59'})
In [9]: s.wait_for_job('a7469b2913504c80934ca2cd72c7fa59')
{'status': 'Complete', 'model_name': 'cancer-model', 'loss': \
0.4150310754776001, 'percent_complete': 100, 'starttime': \
'2018-01-17T13:20:32.529977', 'dataset_name': 'cancer_train.csv', \
'generations': 19}
Out[9]: (True, 'Job completed')
In [10]: s.download_artifact('d27466dec9b548b5bd42f0295a23432e')
Out[10]:
(True,
 ['Diagnosis',
  'BENIGN',
  'BENIGN',
  'BENIGN',
  'BENIGN',
  'MALIGNANT',
```

# **Revision Table**

Version	Author	Date	Notes
v 0.1	RL & SCheng	01.11.2018	Initial doc
v 0.2	RL & SCheng	01.17.2018	add info from quick start, update to html
v 0.3	RL & SCheng	01.18.2018	add new info, include example in ref section.
v 0.4	RL & SCheng	01.22.2018	SCh's comments, add back section from getting started.
11/ 11/ 1	RL & SCheng & KM	01.22.2018ь	Completed KMs suggestions, edited section, generated 01.22.2018b version