# Package 'solvebio'

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Description R language bindings for SolveBio's API.  SolveBio is a biomedical knowledge hub that enables life science organizations to collect and harmonize the complex, disparate  ``multi-omic" data essential for today's R&D and BI needs.
<pre>URL https://github.com/solvebio/solvebio-r</pre>
Imports httr, jsonlite, mime
License MIT + file LICENSE
Collate solvebio.R annotation.R application.R beacon.R beacon_set.R dataset.R dataset_commit.R dataset_export.R dataset_field.R dataset_import.R dataset_migration.R dataset_template.R global_search.R object.R saved_query.R shiny.R task.R user.R utils.R vault.R zzz.R
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Contents
Annotator.annotate

2 Contents

Application.delete	6
Application.retrieve	7
Application.update	7
Beacon.all	8
Beacon.create	9
Beacon.delete	10
Beacon.query	10
Beacon.retrieve	11
Beacon.update	12
BeaconSet.all	12
BeaconSet.create	13
BeaconSet.delete	14
BeaconSet.query	14
BeaconSet.retrieve	15
BeaconSet.update	16
createEnv	16
Dataset.activity	17
Dataset.all	18
Dataset.count	18
Dataset.create	19
Dataset.data	20
Dataset.delete	20
Dataset.disable_global_beacon	21
Dataset.enable_global_beacon	22
Dataset.facets	22
Dataset.fields	23
Dataset.get_by_full_path	24
Dataset.get_global_beacon_status	24
Dataset.get_or_create_by_full_path	25
Dataset.query	26
Dataset.retrieve	27
Dataset.template	27
Dataset.update	28
DatasetCommit.all	29
DatasetCommit.delete	29
DatasetCommit.retrieve	30
DatasetExport.all	31
DatasetExport.create	31
DatasetExport.delete	32
DatasetExport.get_download_url	33
DatasetExport.retrieve	33
DatasetField.all	34
DatasetField.create	35
DatasetField.facets	35
DatasetField.retrieve	36
DatasetField.update	37
DatasetImport.all	37
DatasetImport create	38

Contents 3

DatasetImport.delete	. 39
DatasetImport.retrieve	
DatasetMigration.all	
DatasetMigration.create	
DatasetMigration.delete	
DatasetMigration.retrieve	
DatasetTemplate.all	
DatasetTemplate.create	
DatasetTemplate.delete	
DatasetTemplate.retrieve	
DatasetTemplate.update	
Expression.evaluate	
GlobalSearch.facets	
GlobalSearch.request	
GlobalSearch.search	
GlobalSearch.subjects	
GlobalSearch.subjects_count	
login	
Object.all	
Object.create	
Object.data	
Object.delete	
Object.disable_global_beacon	
Object.enable_global_beacon	
Object.fields	
Object.get_by_full_path	
Object.get_by_path	
Object.get_download_url	
Object.get_global_beacon_status	
Object.get_or_upload_file	
Object.query	
Object.retrieve	
Object.update	
Object.upload_file	
protectedServer	
protectedServerJS	
SavedQuery.all	
SavedQuery.create	
SavedQuery.retrieve	
Task.all	
Task.follow	69
Task.retrieve	69
User.retrieve	. 70 71
	. 71
Vault create dataset	. 71 72
VALUE E DE AUGUSTA DE LA CANTA DEL CANTA DE LA CANTA DEL CANTA DE LA CANTA DEL CANTA DEL CANTA DEL CANTA DE LA CANTA DEL CANTA DEL CANTA DE LA CANTA DEL CANTA DEL CANTA DEL CANTA DE LA CANTA DE LA CANTA DEL CANTA DEL CANTA DEL CANTA DEL CANTA DEL	, ,

4 Annotator.annotate

	Vault.create_folder	73
	Vault.datasets	73
	Vault.delete	74
	Vault.files	75
	Vault.folders	75
	Vault.get_by_full_path	76
	Vault.get_or_create_by_full_path	77
	Vault.get_personal_vault	77
	Vault.objects	78
	Vault.retrieve	79
	Vault.search	79
	Vault.update	80
ndex		81

Annotator.annotate

Annotator. annotate

# Description

Annotate a data table/frame with additional fields.

## Usage

```
Annotator.annotate(
  records,
  fields,
  include_errors = FALSE,
  raw = FALSE,
  env = .solveEnv
)
```

## **Arguments**

records The data table or data frame to annotate.

fields The fields to add.

include\_errors Set to TRUE to include errors in the output (default: FALSE).

raw Set to TRUE to return the raw response (default: FALSE).

env (optional) Custom client environment.

#### References

https://docs.solvebio.com/

Application.all 5

# **Examples**

```
## Not run:
Annotator.annotate(records=tbl, fields=fields)
## End(Not run)
```

Application.all

Application.all

# Description

Retrieves the metadata about all application on SolveBio available to the current user.

# Usage

```
Application.all(env = .solveEnv, ...)
```

# **Arguments**

```
env (optional) Custom client environment.
... (optional) Additional query parameters.
```

## References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
Application.all()
## End(Not run)
```

Application.create

Application.create

# Description

Create a new SolveBio application.

# Usage

```
Application.create(name, redirect_uris, env = .solveEnv, ...)
```

6 Application.delete

### **Arguments**

name The name of the application.

redirect\_uris A list of space-separated OAuth2 redirect URIs.

env (optional) Custom client environment.
... (optional) Additional application attributes.

#### References

```
https://docs.solvebio.com/
```

# **Examples**

Application.delete

Application.delete

# **Description**

Delete a specific application from SolveBio.

#### **Usage**

```
Application.delete(client_id, env = .solveEnv)
```

## **Arguments**

client\_id The client ID for the application.
env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

```
## Not run:
Application.delete("abcd1234")
## End(Not run)
```

Application.retrieve 7

```
Application.retrieve Application.retrieve
```

## **Description**

Retrieves the metadata about a specific application SolveBio.

## Usage

```
Application.retrieve(client_id, env = .solveEnv)
```

## Arguments

client\_id The client ID for the application.
env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

# **Examples**

```
## Not run:
Application.retrieve("abcd1234")
## End(Not run)
```

Application.update

Application.update

# Description

Updates the attributes of an existing application.

## Usage

```
Application.update(client_id, env = .solveEnv, ...)
```

#### **Arguments**

```
client_id The client ID for the application.
env (optional) Custom client environment.
... Application attributes to change.
```

Beacon.all

## References

```
https://docs.solvebio.com/
```

# **Examples**

Beacon.all

Beacon.all

# Description

Retrieves the metadata about all beacons on SolveBio accessible to the current user.

# Usage

```
Beacon.all(env = .solveEnv, ...)
```

# **Arguments**

```
env (optional) Custom client environment.
... (optional) Additional query parameters.
```

#### References

```
https://docs.solvebio.com/
```

```
## Not run:
Beacon.all()
## End(Not run)
```

Beacon.create 9

Beacon.create Beacon.create

## **Description**

Add a new beacon to an existing beacon set. The beacon set must already exist in order to add beacons.

## Usage

```
Beacon.create(beacon_set_id, vault_object_id, title, env = .solveEnv, ...)
```

## **Arguments**

```
beacon_set_id The ID of the parent beacon set.

vault_object_id

The ID of the vault object (i.e. dataset) queried by the beacon.

title The title displayed for the beacon.

env (optional) Custom client environment.

... (optional) Additional beacon attributes (such as description and params).
```

## References

```
https://docs.solvebio.com/
```

Beacon.query

Beacon.delete

Beacon.delete

# Description

Delete a specific beacon from SolveBio.

## Usage

```
Beacon.delete(id, env = .solveEnv)
```

## **Arguments**

id The ID of the beacon.

env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

# Examples

```
## Not run:
Beacon.delete("1234567890")
## End(Not run)
```

Beacon.query

Beacon.query

# Description

Query an individual beacon.

# Usage

```
Beacon.query(id, query, entity_type, env = .solveEnv, ...)
```

## **Arguments**

id The ID of the beacon.

query The entity ID or query string.

entity\_type (optional) A valid SolveBio entity type.
env (optional) Custom client environment.
... (optional) Additional query parameters.

Beacon.retrieve 11

## References

```
https://docs.solvebio.com/
```

# **Examples**

Beacon.retrieve

Beacon.retrieve

# Description

Retrieves the metadata about a specific beacon on SolveBio.

# Usage

```
Beacon.retrieve(id, env = .solveEnv)
```

# Arguments

id The ID of the beacon.

env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

```
## Not run:
Beacon.retrieve("1234")
## End(Not run)
```

12 BeaconSet.all

Beacon.update

Beacon.update

## **Description**

Updates the attributes of an existing beacon.

## Usage

```
Beacon.update(id, env = .solveEnv, ...)
```

# Arguments

```
id The ID of the beacon to update.env (optional) Custom client environment.... Beacon attributes to change.
```

#### References

```
https://docs.solvebio.com/
```

# **Examples**

BeaconSet.all

BeaconSet.all

## **Description**

Retrieves the metadata about all beacon sets on SolveBio accessible to the current user.

# Usage

```
BeaconSet.all(env = .solveEnv, ...)
```

## **Arguments**

```
env (optional) Custom client environment.
... (optional) Additional query parameters.
```

BeaconSet.create 13

# References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
BeaconSet.all()
## End(Not run)
```

BeaconSet.create

BeaconSet.create

# Description

Create a new beacon set.

## Usage

```
BeaconSet.create(title, description, is_shared = FALSE, env = .solveEnv, ...)
```

# Arguments

title The title displayed for the beacon set.

description (optional) An optional description for the new beacon set.

is\_shared If TRUE, everyone else in your account will be able to see and query the beacon

set, but will not be able to edit it. (Default: FALSE)

env (optional) Custom client environment.

... (optional) Additional beacon set attributes.

#### References

```
https://docs.solvebio.com/
```

14 BeaconSet.query

BeaconSet.delete

BeaconSet.delete

## **Description**

Delete a specific beacon set (including all its beacons) from SolveBio.

## Usage

```
BeaconSet.delete(id, env = .solveEnv)
```

## **Arguments**

id The ID of the beacon set.

env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

# **Examples**

```
## Not run:
BeaconSet.delete("1234")
## End(Not run)
```

BeaconSet.query

BeaconSet.query

# Description

Query a beacon set (i.e. all the beacons within a beacon set).

# Usage

```
BeaconSet.query(id, query, entity_type, env = .solveEnv, ...)
```

## **Arguments**

```
id The ID of the beacon set.
query The entity ID or query string.
```

entity\_type (optional) A valid SolveBio entity type.
env (optional) Custom client environment.
... (optional) Additional query parameters.

BeaconSet.retrieve 15

## References

```
https://docs.solvebio.com/
```

# **Examples**

BeaconSet.retrieve

BeaconSet.retrieve

# Description

Retrieves the metadata about a specific beacon set on SolveBio.

# Usage

```
BeaconSet.retrieve(id, env = .solveEnv)
```

# Arguments

id The ID of the beacon set.env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

```
## Not run:
BeaconSet.retrieve("1234")
## End(Not run)
```

16 createEnv

BeaconSet.update

BeaconSet.update

# Description

Updates the attributes of an existing beacon set.

# Usage

```
BeaconSet.update(id, env = .solveEnv, ...)
```

# Arguments

```
id The ID of the beacon set to update.env (optional) Custom client environment.... Beacon set attributes to change.
```

#### References

```
https://docs.solvebio.com/
```

## **Examples**

createEnv

createEnv

# Description

Create a new SolveBio environment.

## Usage

```
createEnv(token, token_type = "Token", host = .solveEnv$host)
```

Dataset.activity 17

## **Arguments**

token A SolveBio API key or OAuth2 token token\_type SolveBio token type (default: Token)

host (optional) The SolveBio API host (default: https://api.solvebio.com)

#### References

```
https://docs.solvebio.com/
```

#### **Examples**

```
## Not run:
env <- createEnv("MyAPIkey")
User.retrieve(env = myEnv)
## End(Not run)</pre>
```

Dataset.activity

Dataset.activity

## **Description**

A helper function to get or follow the current activity on a dataset.

## Usage

```
Dataset.activity(id, follow = TRUE, env = .solveEnv)
```

## **Arguments**

id String The ID of a SolveBio datasetfollow Follow active tasks until they complete.env (optional) Custom client environment.

## References

```
https://docs.solvebio.com/
```

```
## Not run:
Dataset.activity("1234567890")
## End(Not run)
```

18 Dataset.count

Dataset.all

Dataset.all

# Description

Retrieves the metadata about datasets on SolveBio.

# Usage

```
Dataset.all(env = .solveEnv, ...)
```

## **Arguments**

```
env (optional) Custom client environment.
... (optional) Additional query parameters (e.g. page).
```

#### References

```
https://docs.solvebio.com/
```

# **Examples**

```
## Not run:
Dataset.all()
## End(Not run)
```

Dataset.count

Dataset.count

## **Description**

Returns the total number of records for a given SolveBio dataset.

# Usage

```
Dataset.count(id, env = .solveEnv, ...)
```

#### **Arguments**

```
    id The ID of a SolveBio dataset, or a Dataset object.
    env (optional) Custom client environment.
    ... (optional) Additional query parameters (e.g. filters, limit, offset).
```

Dataset.create 19

#### References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
dataset <- Dataset.get_by_full_path("solvebio:public:/ClinVar/3.7.4-2017-01-30/Variants-GRCh37")
Dataset.count(dataset)
Dataset.count(dataset, filters='[["gene_symbol", "BRCA2"]]')
## End(Not run)</pre>
```

Dataset.create

Dataset.create

## Description

Create an empty SolveBio dataset.

## Usage

```
Dataset.create(vault_id, vault_parent_object_id, name, env = .solveEnv, ...)
```

# **Arguments**

#### References

```
https://docs.solvebio.com/
```

```
## Not run:
Dataset.create(vault_id=vault$id, vault_parent_object_id=NULL, name="My Dataset")
## End(Not run)
```

20 Dataset.delete

Dataset.data

Dataset.data

## **Description**

Returns one page of documents from a SolveBio dataset and processes the response.

## Usage

```
Dataset.data(id, filters, env = .solveEnv, ...)
```

## **Arguments**

id The ID of a SolveBio dataset, or a Dataset object.

filters (optional) Query filters.

env (optional) Custom client environment.

... (optional) Additional query parameters (e.g. limit, offset).

#### References

```
https://docs.solvebio.com/
```

# **Examples**

```
## Not run:
Dataset.data("1234567890")
## End(Not run)
```

Dataset.delete

Dataset.delete

## **Description**

Delete a specific dataset from SolveBio.

# Usage

```
Dataset.delete(id, env = .solveEnv)
```

## Arguments

id	String The ID of a SolveBio dataset
env	(optional) Custom client environment.

## References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
Dataset.delete("1234567890")
## End(Not run)
```

```
\label{lem:disable_global_beacon} Dataset.disable\_global\_beacon
```

# **Description**

Disables Global Beacon for the dataset.

# Usage

```
Dataset.disable_global_beacon(id, env = .solveEnv)
```

## **Arguments**

id The ID of a SolveBio dataset.

env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

```
## Not run:
Dataset.disable_global_beacon("1234567890")
## End(Not run)
```

22 Dataset.facets

```
Dataset.enable_global_beacon
```

Dataset.enable\_global\_beacon

### **Description**

Enables Global Beacon for the the dataset.

#### Usage

```
Dataset.enable_global_beacon(id, env = .solveEnv)
```

## **Arguments**

id The ID of a SolveBio dataset.

env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
Dataset.enable_global_beacon("1234567890")
## End(Not run)
```

Dataset.facets

Dataset.facets

# **Description**

Retrieves aggregated statistics or term counts for one or more fields in a SolveBio dataset. Returns a list of data frames, one for each requested facet.

#### Usage

```
Dataset.facets(id, facets, env = .solveEnv, ...)
```

## **Arguments**

id The ID of a SolveBio dataset, or a Dataset object.

facets A list of one or more field facets.
env (optional) Custom client environment.

... (optional) Additional query parameters (e.g. filters, limit, offset).

Dataset.fields 23

## References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
Dataset.facets("1234567890", list("clinical_significance", "gene_symbol"))
## End(Not run)
```

Dataset.fields

Dataset.fields

## **Description**

Retrieves the list of fields and field metadata for a dataset.

# Usage

```
Dataset.fields(id, env = .solveEnv, ...)
```

## **Arguments**

```
    id The ID of a SolveBio dataset, or a Dataset object.
    env (optional) Custom client environment.
    ... (optional) Additional query parameters (e.g. limit, offset).
```

#### References

```
https://docs.solvebio.com/
```

```
## Not run:
Dataset.fields("1234567890")
## End(Not run)
```

```
Dataset.get_by_full_path

Dataset.get_by_full_path
```

## **Description**

A helper function to get a dataset by its full path.

## Usage

```
Dataset.get_by_full_path(full_path, env = .solveEnv)
```

## **Arguments**

```
full_path A valid full path to a dataset.
env (optional) Custom client environment.
```

#### References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
Dataset.get_by_full_path("solvebio:public:/ClinVar/3.7.4-2017-01-30/Variants-GRCh37")
## End(Not run)
```

```
Dataset.get_global_beacon_status 
 Dataset.get_global_beacon_status
```

## **Description**

Retrieves the global beacon status for the dataset.

# Usage

```
Dataset.get_global_beacon_status(
   id,
   raise_on_disabled = FALSE,
   env = .solveEnv
)
```

#### **Arguments**

```
id The ID of a SolveBio dataset.

raise_on_disabled

Whether to raise an exception if Global Beacon is disabled or to return NULL.

env (optional) Custom client environment.
```

#### References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
Dataset.get_global_beacon_status("1234567890")
## End(Not run)
```

```
\label{lem:decomposition} Dataset.get\_or\_create\_by\_full\_path \label{lem:decomposition} Dataset.get\_or\_create\_by\_full\_path
```

## **Description**

A helper function to get or create a dataset by its full path.

#### Usage

```
Dataset.get_or_create_by_full_path(full_path, env = .solveEnv, ...)
```

#### **Arguments**

full\_path A valid full path to a dataset.

env (optional) Custom client environment.

... Additional dataset creation parameters.

#### References

```
https://docs.solvebio.com/
```

```
## Not run:
Dataset.get_or_create_by_full_path("MyVault:/folder/sub-folder/dataset")
## End(Not run)
```

26 Dataset.query

Dataset.query

Dataset.query

## **Description**

Queries a SolveBio dataset and returns an R data frame containing all records. Returns a single page of results otherwise (default).

#### Usage

```
Dataset.query(
   id,
   paginate = FALSE,
   use_field_titles = TRUE,
   env = .solveEnv,
   ...
)
```

## **Arguments**

```
id The ID of a SolveBio dataset, or a Dataset object.

paginate When set to TRUE, retrieves all records (memory permitting).

use_field_titles

(optional) Use field title instead of field name for query.

env (optional) Custom client environment.

... (optional) Additional query parameters (e.g. filters, limit, offset).
```

#### References

```
https://docs.solvebio.com/
```

```
## Not run:
Dataset.query("12345678790", paginate=TRUE)
## End(Not run)
```

Dataset.retrieve 27

Dataset.retrieve

Dataset.retrieve

# Description

Retrieves the metadata about a specific dataset from SolveBio.

## Usage

```
Dataset.retrieve(id, env = .solveEnv)
```

## **Arguments**

id String The ID of a SolveBio dataset env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

# **Examples**

```
## Not run:
Dataset.retrieve("1234567890")
## End(Not run)
```

Dataset.template

Dataset.template

# Description

Retrieves the template for a dataset.

# Usage

```
Dataset.template(id, env = .solveEnv)
```

## **Arguments**

id String The ID of a SolveBio dataset env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

28 Dataset.update

# **Examples**

```
## Not run:
Dataset.template("1234567890")
## End(Not run)
```

Dataset.update

Dataset.update

# Description

Updates the attributes of an existing dataset.

# Usage

```
Dataset.update(id, env = .solveEnv, ...)
```

# Arguments

id The ID of the dataset to update.env (optional) Custom client environment.... Dataset attributes to change.

# References

```
https://docs.solvebio.com/
```

DatasetCommit.all 29

DatasetCommit.all

DatasetCommit.all

## **Description**

Retrieves the metadata about all dataset commits on SolveBio.

## Usage

```
DatasetCommit.all(env = .solveEnv, ...)
```

## **Arguments**

```
env
                   (optional) Custom client environment.
                   (optional) Additional query parameters (e.g. page).
. . .
```

## References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
DatasetCommit.all()
## End(Not run)
```

DatasetCommit.delete
DatasetCommit.delete

# Description

Deletes a specific dataset commit on SolveBio.

# Usage

```
DatasetCommit.delete(id, env = .solveEnv)
```

## **Arguments**

String The ID or full name of a SolveBio dataset commit. id

(optional) Custom client environment. env

## References

```
https://docs.solvebio.com/
```

30 DatasetCommit.retrieve

# Examples

```
## Not run:
DatasetCommit.delete(<ID>)
## End(Not run)
```

DatasetCommit.retrieve

DatasetCommit.retrieve

# Description

Retrieves the metadata about a specific dataset commit on SolveBio.

# Usage

```
DatasetCommit.retrieve(id, env = .solveEnv)
```

# Arguments

id String The ID of a SolveBio dataset commit.

env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

```
## Not run:
DatasetCommit.retrieve(<ID>)
## End(Not run)
```

DatasetExport.all 31

DatasetExport.all DatasetExport.all

# Description

Retrieves the metadata about all dataset exports on SolveBio.

# Usage

```
DatasetExport.all(env = .solveEnv, ...)
```

## **Arguments**

```
env (optional) Custom client environment.
... (optional) Additional query parameters (e.g. page).
```

#### References

```
https://docs.solvebio.com/
```

# **Examples**

```
## Not run:
DatasetExport.all()
## End(Not run)
```

DatasetExport.create
DatasetExport.create

# Description

Create a new dataset export.

# Usage

```
DatasetExport.create(
  dataset_id,
  format = "json",
  params = list(),
  follow = FALSE,
  env = .solveEnv,
  ...
)
```

32 DatasetExport.delete

#### **Arguments**

dataset\_id The target dataset ID.

format (optional) The export format (default: json).
params (optional) Query parameters for the export.

follow (default: FALSE) Follow the export task until it completes.

env (optional) Custom client environment.

... (optional) Additional dataset export parameters.

#### References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
DatasetExport.create(dataset_id=<ID>, format='json', params=list(fields=c("field_1"), limit=100))
## End(Not run)
```

 ${\tt DatasetExport.delete} \quad \textit{DatasetExport.delete}$ 

# Description

Deletes a specific dataset export on SolveBio.

#### Usage

```
DatasetExport.delete(id, env = .solveEnv)
```

# Arguments

id String The ID of a SolveBio dataset export. env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

```
## Not run:
DatasetExport.delete(<ID>)
## End(Not run)
```

```
{\tt DatasetExport.get\_download\_url}
```

 $DatasetExport.get\_download\_url$ 

## **Description**

Helper method to get the download URL for a dataset export.

## Usage

```
DatasetExport.get_download_url(id, env = .solveEnv)
```

# **Arguments**

id The ID of the dataset export.

env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
DatasetExport.get_download_url("1234567890")
## End(Not run)
```

DatasetExport.retrieve

DatasetExport.retrieve

# Description

Retrieves the metadata about a specific dataset export on SolveBio.

## Usage

```
DatasetExport.retrieve(id, env = .solveEnv)
```

## Arguments

id String The ID of a SolveBio dataset export.
env (optional) Custom client environment.

DatasetField.all

## References

```
https://docs.solvebio.com/
```

# Examples

```
## Not run:
DatasetExport.retrieve(<ID>)
## End(Not run)
```

DatasetField.all

DatasetField.all

# Description

Retrieves the metadata about all dataset fields on SolveBio.

# Usage

```
DatasetField.all(env = .solveEnv, ...)
```

# Arguments

```
env (optional) Custom client environment.... (optional) Additional query parameters (e.g. page).
```

## References

```
https://docs.solvebio.com/
```

```
## Not run:
DatasetField.all()
## End(Not run)
```

DatasetField.create 35

# Description

Create a new dataset field.

# Usage

```
DatasetField.create(dataset_id, name, data_type = "auto", env = .solveEnv, ...)
```

# Arguments

```
dataset_id The dataset ID.
```

name The name of the dataset field.

data\_type (optional) The data type for the field (default: auto).

env (optional) Custom client environment.

... (optional) Additional dataset import attributes.

#### References

```
https://docs.solvebio.com/
```

# **Examples**

```
## Not run:
DatasetField.create(dataset_id=<ID>, name="my_field", title="My Field", data_type="string")
## End(Not run)
```

# Description

Returns the facets for a given dataset field.

## Usage

```
DatasetField.facets(id, env = .solveEnv, ...)
```

36 DatasetField.retrieve

### **Arguments**

id String The ID of a dataset field.

env (optional) Custom client environment.
... (optional) Additional query parameters.

#### References

```
https://docs.solvebio.com/
```

# **Examples**

```
## Not run:
DatasetField.facets(691)
## End(Not run)
```

DatasetField.retrieve DatasetField.retrieve

# Description

Retrieves the metadata about a specific dataset field.

#### Usage

```
DatasetField.retrieve(id, env = .solveEnv)
```

## **Arguments**

id String The ID of a dataset field.

env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

```
## Not run:
DatasetField.retrieve(691)
## End(Not run)
```

DatasetField.update 37

DatasetField.update Da

Dataset Field.update

## **Description**

Updates the attributes of an existing dataset field. NOTE: The data\_type of a field cannot be changed.

## Usage

```
DatasetField.update(id, env = .solveEnv, ...)
```

## **Arguments**

id The ID of the dataset field to update.env (optional) Custom client environment.... Dataset field attributes to change.

#### References

```
https://docs.solvebio.com/
```

## **Examples**

DatasetImport.all

DatasetImport.all

# Description

Retrieves the metadata about all dataset imports on SolveBio.

## Usage

```
DatasetImport.all(env = .solveEnv, ...)
```

38 DatasetImport.create

## **Arguments**

```
env (optional) Custom client environment.
... (optional) Additional query parameters (e.g. page).
```

#### References

```
https://docs.solvebio.com/
```

# **Examples**

```
## Not run:
DatasetImport.all()
## End(Not run)
```

 ${\tt DatasetImport.create} \quad \textit{DatasetImport.create}$ 

## **Description**

Create a new dataset import. Either an object\_id, manifest, or data\_records is required.

# Usage

```
DatasetImport.create(dataset_id, commit_mode = "append", env = .solveEnv, ...)
```

#### **Arguments**

```
dataset_id The target dataset ID.

commit_mode (optional) The commit mode (default: append).

env (optional) Custom client environment.

... (optional) Additional dataset import attributes.
```

## References

```
https://docs.solvebio.com/
```

```
## Not run:
DatasetImport.create(dataset_id=<ID>, upload_id=<ID>)
## End(Not run)
```

DatasetImport.delete 39

 ${\tt DatasetImport.delete} \quad \textit{DatasetImport.delete}$ 

# Description

Deletes a specific dataset import on SolveBio.

## Usage

```
DatasetImport.delete(id, env = .solveEnv)
```

## **Arguments**

id String The ID of a SolveBio dataset import.
env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

# **Examples**

```
## Not run:
DatasetImport.delete(<ID>)
## End(Not run)
```

DatasetImport.retrieve

DatasetImport.retrieve

## **Description**

Retrieves the metadata about a specific dataset import on SolveBio.

## Usage

```
DatasetImport.retrieve(id, env = .solveEnv)
```

# Arguments

id String The ID of a SolveBio dataset import.
env (optional) Custom client environment.

40 DatasetMigration.all

#### References

```
https://docs.solvebio.com/
```

# Examples

```
## Not run:
DatasetImport.retrieve(<ID>)
## End(Not run)
```

 ${\tt DatasetMigration.all} \quad \textit{DatasetMigration.all}$ 

# Description

Retrieves the metadata about all dataset migrations on SolveBio.

# Usage

```
DatasetMigration.all(env = .solveEnv, ...)
```

# Arguments

```
env (optional) Custom client environment.
... (optional) Additional query parameters (e.g. page).
```

## References

```
https://docs.solvebio.com/
```

```
## Not run:
DatasetMigration.all()
## End(Not run)
```

DatasetMigration.create

DatasetMigration.create

Dataset Migration. create

#### **Description**

Create a new dataset migration.

## Usage

```
DatasetMigration.create(
   source_id,
   target_id,
   commit_mode = "append",
   source_params = NULL,
   target_fields = NULL,
   include_errors = FALSE,
   env = .solveEnv,
   ...
)
```

#### **Arguments**

```
The source dataset ID.
source_id
target_id
                  The target dataset ID.
commit_mode
                  (optional) The commit mode (default: append).
                  (optional) The query parameters used on the source dataset.
source_params
target_fields
                  (optional) A list of valid dataset fields to add or override in the target dataset.
include_errors
                  (optional) If TRUE, a new field (_errors) will be added to each record containing
                   expression evaluation errors (default: FALSE).
                   (optional) Custom client environment.
env
                   (optional) Additional dataset migration attributes.
```

#### References

```
https://docs.solvebio.com/
```

```
## Not run:
DatasetMigration.create(dataset_id=<ID>, upload_id=<ID>)
## End(Not run)
```

DatasetMigration.delete

DatasetMigration.delete

## **Description**

Deletes specific dataset migration on SolveBio.

## Usage

```
DatasetMigration.delete(id, env = .solveEnv)
```

# Arguments

id String The ID of a SolveBio dataset migration.

env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
DatasetMigration.delete(<ID>)
## End(Not run)
```

DatasetMigration.retrieve

DatasetMigration.retrieve

# Description

Retrieves the metadata about a specific dataset migration on SolveBio.

## Usage

```
DatasetMigration.retrieve(id, env = .solveEnv)
```

## Arguments

id String The ID of a SolveBio dataset migration.

env (optional) Custom client environment.

DatasetTemplate.all 43

## References

```
https://docs.solvebio.com/
```

# Examples

```
## Not run:
DatasetMigration.retrieve(<ID>)
## End(Not run)
```

DatasetTemplate.all

DatasetTemplate.all

# Description

Retrieves the metadata about all dataset templates on SolveBio.

# Usage

```
DatasetTemplate.all(env = .solveEnv, ...)
```

# Arguments

```
env (optional) Custom client environment.... (optional) Additional query parameters (e.g. page).
```

## References

```
https://docs.solvebio.com/
```

```
## Not run:
DatasetTemplate.all()
## End(Not run)
```

DatasetTemplate.create

Dataset Template.create

## **Description**

Create a SolveBio dataset template.

## Usage

```
DatasetTemplate.create(env = .solveEnv, ...)
```

# **Arguments**

```
env (optional) Custom client environment.
... (optional) Dataset template attributes.
```

#### References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
DatasetTemplate.create(name="My Dataset Template")
## End(Not run)
```

DatasetTemplate.delete

Dataset Template. de lete

# Description

Deletes a specific dataset template on SolveBio.

# Usage

```
DatasetTemplate.delete(id, env = .solveEnv)
```

# Arguments

id String The ID or full name of a SolveBio dataset template.

env (optional) Custom client environment.

## References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
DatasetTemplate.delete(<ID>)
## End(Not run)
```

DatasetTemplate.retrieve

Dataset Template. retrieve

# Description

Retrieves the metadata about a specific dataset template on SolveBio.

# Usage

```
DatasetTemplate.retrieve(id, env = .solveEnv)
```

# Arguments

id String The ID of a SolveBio dataset template.

env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

```
## Not run:
DatasetTemplate.retrieve(<ID>)
## End(Not run)
```

Expression.evaluate

DatasetTemplate.update

Dataset Template.update

# Description

Updates the attributes of an existing dataset template.

# Usage

```
DatasetTemplate.update(id, env = .solveEnv, ...)
```

# Arguments

```
id The ID of the dataset template to update.env (optional) Custom client environment.... Dataset template attributes to change.
```

## References

```
https://docs.solvebio.com/
```

# **Examples**

Expression.evaluate

Expression.evaluate

# Description

Evaluate a SolveBio expression.

GlobalSearch.facets 47

#### Usage

```
Expression.evaluate(
  expression,
  data_type = "string",
  is_list = FALSE,
  data = NULL,
  raw = FALSE,
  env = .solveEnv
)
```

#### **Arguments**

expression The SolveBio expression string.

data\_type The data type to cast the expression result (default: string).

is\_list Set to TRUE if the result is expected to be a list (default: FALSE).

data Variables used in the expression (default: NULL).

raw Set to TRUE to return the raw response (default: FALSE).

env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

# **Examples**

```
## Not run:
Expression.evaluate("1 + 1", data_type="integer", is_list=FALSE)
## End(Not run)
```

GlobalSearch.facets GlobalSearch.facets

## **Description**

Performs a Global Search based on provided filters, entities, queries, and returns an R data frame containing the facets results from API response.

#### Usage

```
GlobalSearch.facets(facets, env = .solveEnv, ...)
```

GlobalSearch.request

#### **Arguments**

```
facets Facets list.
```

env (optional) Custom client environment.

... (optional) Additional query parameters (e.g. filters, entities, query).

#### References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
GlobalSearch.facets(facets="study")
## End(Not run)
```

GlobalSearch.request GlobalSearch.request

## **Description**

Performs a single Global Search API request with the provided filters, queries and entities. A single request will only retrieve one page of results (based on the 'limit' parameter). Use 'GlobalSearch.search()' to retrieve all pages of results. Returns the full API response (containing attributes: results, vaults, subjects, subjects\_count, total)

# Usage

```
GlobalSearch.request(query = NULL, filters, entities, env = .solveEnv, ...)
```

(optional) Additional query parameters (e.g. limit, offset).

# Arguments

```
query (optional) Advanced search query.

filters (optional) Low-level filter specification.

entities (optional) Low-level entity specification.

env (optional) Custom client environment.
```

#### References

. . .

https://docs.solvebio.com/

GlobalSearch.search 49

#### **Examples**

```
## Not run:
# No filters are applied
GlobalSearch.request()
# Global Beacon search
GlobalSearch.request(entities = '[["gene","BRCA2"]]')
# Type filter (only vaults)
GlobalSearch.request(filters = '[{"and":[["type__in",["vault"]]]}]')
# Advanced search
GlobalSearch.request(query = "fuji")
# Multiple filters and entities
GlobalSearch.request(
  entities = '[["gene","BRCA2"]]',
  filters = '[{
               "and": [
                      {"and": [
                         ["created_at__range",["2021-11-28","2021-12-28"]]]},
                         ["type__in",["dataset"]]
                     ]
             }]'
)
## End(Not run)
```

GlobalSearch.search

GlobalSearch.search

## **Description**

Performs a Global Search based on provided filters, entities, queries, and returns an R data frame containing results from API response. Returns a single page of results otherwise (default).

## Usage

```
GlobalSearch.search(paginate = FALSE, env = .solveEnv, ...)
```

## **Arguments**

```
paginate When set to TRUE, retrieves all records (memory permitting).
env (optional) Custom client environment.
... (optional) Additional query parameters (e.g. filters, entities, query, limit, offset).
```

#### References

```
https://docs.solvebio.com/
```

#### **Examples**

```
## Not run:
# No filters applied
GlobalSearch.search()

#Global Beacon search
GlobalSearch.search(entities = '[["gene","BRCA2"]]')

GlobalSearch.search(filters = '[{"and":[["type__in",["vault"]]]}]')

# Advanced search
GlobalSearch.search(query = "fuji")

## End(Not run)
```

GlobalSearch.subjects GlobalSearch.subjects

## **Description**

Performs a Global Search based on provided filters, entities, queries, and returns an R data frame containing subjects from API response.

## Usage

```
GlobalSearch.subjects(env = .solveEnv, ...)
```

# Arguments

```
env (optional) Custom client environment.
... (optional) Additional query parameters (e.g. filters, entities, query, limit, offset).
```

#### References

```
https://docs.solvebio.com/
```

```
## Not run:
GlobalSearch.subjects(entities = '[["gene","BRCA2"]]')
## End(Not run)
```

```
GlobalSearch.subjects_count
```

GlobalSearch.subjects\_count

## **Description**

Performs a Global Search based on provided filters, entities, queries, and returns the total number of subjects from API response.

#### Usage

```
GlobalSearch.subjects_count(env = .solveEnv, ...)
```

## **Arguments**

```
env (optional) Custom client environment.
... (optional) Additional query parameters (e.g. filters, entities, query, limit, offset).
```

#### References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
GlobalSearch.subjects_count(entities = '[["gene","BRCA2"]]')
## End(Not run)
```

login

login

# Description

Store and verify your SolveBio credentials.

#### Usage

```
login(api_key, api_host, env = .solveEnv)
```

## Arguments

api\_key Your SolveBio API key

env (optional) The R environment used to store API credentials.

52 Object.all

## References

```
https://docs.solvebio.com/
```

# Examples

```
## Not run:
login()
## End(Not run)
```

Object.all

Object.all

# Description

Retrieves the metadata about all objects on SolveBio accessible to the current user.

# Usage

```
Object.all(env = .solveEnv, ...)
```

# Arguments

```
env (optional) Custom client environment.
... (optional) Additional query parameters.
```

## References

```
https://docs.solvebio.com/
```

```
## Not run:
Object.all()
## End(Not run)
```

Object.create 53

Object.create

Object.create

# Description

Create a SolveBio object.

# Usage

```
Object.create(
  vault_id,
  parent_object_id,
  object_type,
  filename,
  env = .solveEnv,
  ...
)
```

## Arguments

## References

```
https://docs.solvebio.com/
```

54 Object.delete

Object.data

Object.data

## **Description**

Returns one page of documents from a SolveBio file (object) and processes the response.

## Usage

```
Object.data(id, filters, col.names = NULL, env = .solveEnv, ...)
```

# **Arguments**

id The ID of a SolveBio file (vault object).

filters (optional) Query filters.

col.names (optional) Force data frame column name ordering.

env (optional) Custom client environment.

... (optional) Additional query parameters (e.g. limit, offset).

#### References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
Object.data("1234567890")
## End(Not run)
```

Object.delete

Object.delete

## **Description**

Delete a specific object from SolveBio.

# Usage

```
Object.delete(id, env = .solveEnv)
```

## **Arguments**

id The ID of the object.

env (optional) Custom client environment.

## References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
Object.delete("1234567890")
## End(Not run)
```

```
Object. disable\_global\_beacon \\ Object. disable\_global\_beacon
```

# **Description**

Disables Global Beacon for the specified dataset.

# Usage

```
Object.disable_global_beacon(id, env = .solveEnv)
```

# Arguments

id The ID of a SolveBio dataset.

env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

```
## Not run:
Object.disable_global_beacon("1234567890")
## End(Not run)
```

56 Object.fields

```
Object.enable\_global\_beacon \\ Object.enable\_global\_beacon
```

## **Description**

Enables Global Beacon for the specified dataset.

## Usage

```
Object.enable_global_beacon(id, env = .solveEnv)
```

#### **Arguments**

id The ID of a SolveBio dataset.

env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
Object.enable_global_beacon("1234567890")
## End(Not run)
```

Object.fields

Object.fields

## **Description**

Retrieves the list of fields for a file (JSON, CSV, or TSV).

## Usage

```
Object.fields(id, env = .solveEnv, ...)
```

# Arguments

```
id The ID of a SolveBio file (vault object).
env (optional) Custom client environment.
```

... (optional) Additional query parameters (e.g. limit, offset).

## References

```
https://docs.solvebio.com/
```

# **Examples**

```
## Not run:
Object.fields("1234567890")
## End(Not run)
```

```
Object.get\_by\_full\_path \\ Object.get\_by\_full\_path
```

# Description

A helper function to get an object on SolveBio by its full path.

# Usage

```
Object.get_by_full_path(full_path, env = .solveEnv, ...)
```

## **Arguments**

```
full_path The full path to the object.
env (optional) Custom client environment.
... (optional) Additional query parameters.
```

#### References

```
https://docs.solvebio.com/
```

```
## Not run:
Object.get_by_full_path("solvebio:public:/ClinVar")
## End(Not run)
```

# Description

A helper function to get an object on SolveBio by its path. Used as a pass-through function from some Vault methods.

#### Usage

```
Object.get_by_path(path, env = .solveEnv, ...)
```

## **Arguments**

path The path to the object, relative to a vault.
env (optional) Custom client environment.
... (optional) Additional query parameters.

#### References

```
https://docs.solvebio.com/
```

#### **Examples**

```
## Not run:
Object.get_by_path("/ClinVar")
## End(Not run)
```

```
Object.get_download_url
```

Object.get\_download\_url

# Description

Helper method to get the download URL for a file object.

## Usage

```
Object.get_download_url(id, env = .solveEnv)
```

# Arguments

id The ID of the object.

env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
Object.get_download_url("1234567890")
## End(Not run)
```

```
Object.get\_global\_beacon\_status \\ Object.get\_global\_beacon\_status
```

## **Description**

Retrieves the global beacon status for the specified dataset.

## Usage

```
Object.get_global_beacon_status(id, raise_on_disabled = FALSE, env = .solveEnv)
```

# Arguments

```
id The ID of a SolveBio dataset.

raise_on_disabled

Whether to raise an exception if Global Beacon is disabled or to return NULL.

env (optional) Custom client environment.
```

#### References

```
https://docs.solvebio.com/
```

```
## Not run:
Object.get_global_beacon_status("1234567890")
Object.get_global_beacon_status("1234567890", raise_on_disabled=TRUE)
## End(Not run)
```

```
Object.get_or_upload_file 
 Object.get_or_upload_file
```

# Description

Upload a local file to a vault on SolveBio only if it does not yet exist (by name, at the provided path). The vault path provided is the parent directory for uploaded file. Accepts the same arguments as 'Object.upload\_file'.

## Usage

```
Object.get_or_upload_file(
  local_path,
  vault_id,
  vault_path,
  filename,
  env = .solveEnv
)
```

# Arguments

```
local_path The path to the local file

vault_id The SolveBio vault ID

vault_path The remote path in the vault

filename (optional) The filename for the uploaded file in the vault (default: the basename of the local_path)

env (optional) Custom client environment.
```

#### References

```
https://docs.solvebio.com/
```

```
## Not run:
Object.get_or_upload_file("my_file.json.gz", vault$id, "/parent/directory/")
## End(Not run)
```

Object.query 61

Object.query

Object.query

## **Description**

Queries a SolveBio file (vault object) and returns an R data frame containing all records. Returns a single page of results otherwise (default).

#### Usage

```
Object.query(id, paginate = FALSE, env = .solveEnv, ...)
```

#### **Arguments**

id The ID of a SolveBio file (vault object).

paginate When set to TRUE, retrieves all records (memory permitting).

env (optional) Custom client environment.

... (optional) Additional query parameters (e.g. filters, limit, offset).

#### References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
Object.query("12345678790", paginate=TRUE)
## End(Not run)
```

Object.retrieve

Object.retrieve

## Description

Retrieves the metadata about a specific object on SolveBio.

## Usage

```
Object.retrieve(id, env = .solveEnv)
```

# **Arguments**

id The ID of the object.

env (optional) Custom client environment.

62 Object.update

## References

```
https://docs.solvebio.com/
```

# **Examples**

```
## Not run:
Object.retrieve("1234567890")
## End(Not run)
```

Object.update

Object.update

# Description

Updates the attributes of an existing vault object.

# Usage

```
Object.update(id, env = .solveEnv, ...)
```

# Arguments

id The ID of the vault to update.env (optional) Custom client environment.... Object attributes to change.

## References

```
https://docs.solvebio.com/
```

Object.upload\_file 63

## **Description**

Upload a local file to a vault on SolveBio. The vault path provided is the parent directory for uploaded file.

## Usage

```
Object.upload_file(local_path, vault_id, vault_path, filename, env = .solveEnv)
```

## **Arguments**

local\_path The path to the local file

vault\_id The SolveBio vault ID

vault\_path The remote path in the vault

filename (optional) The filename for the uploaded file in the vault (default: the basename of the local\_path)

env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
Object.upload_file("my_file.json.gz", vault$id, "/parent/directory/")
## End(Not run)
```

protectedServer protectedServer

## **Description**

Wraps an existing Shiny server in an OAuth2 flow.

64 protectedServerJS

#### Usage

```
protectedServer(
   server,
   client_id,
   client_secret = NULL,
   base_url = "https://my.solvebio.com"
)
```

# Arguments

server Your original Shiny server function.

client\_id Your application's client ID.

client\_secret (optional) Your application's client secret.

base\_url (optional) Override the default login host (default: https://my.solvebio.com).

## References

```
https://docs.solvebio.com/
```

# Examples

protectedServerJS

protectedServerUI

## **Description**

Returns ShinyJS-compatible JS code to support cookie-based token storage.

## Usage

```
protectedServerJS()
```

#### References

```
https://docs.solvebio.com/
```

SavedQuery.all 65

## **Examples**

SavedQuery.all

SavedQuery.all

# Description

Retrieves the all saved queries on SolveBio.

# Usage

```
SavedQuery.all(env = .solveEnv, ...)
```

## **Arguments**

```
env (optional) Custom client environment.
... (optional) Additional query parameters (e.g. page).
```

## References

```
https://docs.solvebio.com/
```

```
## Not run:
SavedQuery.all()
## End(Not run)
```

SavedQuery.delete

SavedQuery.create

SavedQuery.create

# Description

Create a SolveBio saved query.

# Usage

```
SavedQuery.create(env = .solveEnv, ...)
```

# **Arguments**

```
env (optional) Custom client environment.
... (optional) Saved query attributes.
```

#### References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
SavedQuery.create(name="My Dataset Template")
## End(Not run)
```

SavedQuery.delete

SavedQuery.delete

# Description

Deletes a specific saved query on SolveBio.

## Usage

```
SavedQuery.delete(id, env = .solveEnv)
```

# **Arguments**

id String The ID of the SolveBio saved query. env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

SavedQuery.retrieve 67

## **Examples**

```
## Not run:
SavedQuery.delete(<ID>)
## End(Not run)
```

SavedQuery.retrieve

SavedQuery.retrieve

# Description

Retrieves a specific saved query on SolveBio by ID.

# Usage

```
SavedQuery.retrieve(id, env = .solveEnv)
```

## **Arguments**

id String The ID of a SolveBio saved query. env (optional) Custom client environment.

## References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
SavedQuery.retrieve(<ID>)
## End(Not run)
```

SavedQuery.update

SavedQuery.update

# Description

Updates the attributes of an existing saved query.

# Usage

```
SavedQuery.update(id, env = .solveEnv, ...)
```

68 Task.all

#### **Arguments**

id The ID of the saved query to update.env (optional) Custom client environment.... Saved query attributes to change.

## References

```
https://docs.solvebio.com/
```

# **Examples**

Task.all

Task.all

# Description

Retrieves the metadata about all tasks on SolveBio accessible to the current user.

## Usage

```
Task.all(env = .solveEnv, ...)
```

# Arguments

env (optional) Custom client environment.
... (optional) Additional query parameters.

#### References

```
https://docs.solvebio.com/
```

```
## Not run:
Task.all()
## End(Not run)
```

Task.follow 69

Task.follow

Task.follow

#### **Description**

A helper function to follow a specific task until it gets completed.

# Usage

```
Task.follow(id, env = .solveEnv, interval = 2)
```

# **Arguments**

id String The ID of a task.

env (optional) Custom client environment.

interval 2. Delay in seconds between each completion status query

#### Value

the task object

#### References

```
https://docs.solvebio.com/
```

# Examples

```
## Not run:
Task.follow("1234567890")
## End(Not run)
```

Task.retrieve

Task.retrieve

## **Description**

Retrieves the metadata about a specific task on SolveBio.

## Usage

```
Task.retrieve(id, env = .solveEnv)
```

## **Arguments**

id The ID of a task.

env (optional) Custom client environment.

70 User.retrieve

## References

```
https://docs.solvebio.com/
```

# **Examples**

```
## Not run:
Task.retrieve("1234567890")
## End(Not run)
```

User.retrieve

User.retrieve

# Description

Retrieves information about the current user.

# Usage

```
User.retrieve(env = .solveEnv)
```

# Arguments

env

(optional) Custom client environment.

## References

```
https://docs.solvebio.com/
```

```
## Not run:
User.retrieve()
## End(Not run)
```

Vault.all 71

Vault.all

Vault.all

# Description

Retrieves the metadata about all accessible vaults.

# Usage

```
Vault.all(..., env = .solveEnv)
```

## **Arguments**

```
(optional) Additional query parameters (e.g. limit, offset).env (optional) Custom client environment.
```

#### References

```
https://docs.solvebio.com/
```

# **Examples**

```
## Not run:
Vault.all()
## End(Not run)
```

Vault.create

Vault.create

# Description

Create a new SolveBio vault.

# Usage

```
Vault.create(name, env = .solveEnv, ...)
```

# Arguments

name	The unique name of the vault.
env	(optional) Custom client environment.
	(optional) Additional vault attributes.

72 Vault.create\_dataset

## References

```
https://docs.solvebio.com/
```

# **Examples**

```
## Not run:
Vault.create(name="my-domain:MyVault")
## End(Not run)
```

# Description

Create a new dataset in a vault.

# Usage

```
Vault.create_dataset(id, path, name, env = .solveEnv, ...)
```

# Arguments

id	The ID of the vault.
path	The path to the dataset, within the vault.
name	The name (filename) for the dataset.
env	(optional) Custom client environment.
	(optional) Additional dataset creation parameters.

## References

```
https://docs.solvebio.com/
```

```
## Not run:
vault = Vault.get_personal_vault()
Vault.create_dataset(vault$id, path="/", name="My Dataset")
## End(Not run)
```

Vault.create\_folder 73

Vault.create\_folder Vault.create\_folder

## **Description**

Create a new folder in a vault.

## Usage

```
Vault.create_folder(id, path, recursive = FALSE, env = .solveEnv, ...)
```

## Arguments

id The ID of the vault.

path The path to the folder, within the vault.

recursive Create all parent directories that do not yet exist (default: FALSE).

env (optional) Custom client environment.

... (optional) Additional folder creation parameters.

#### References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
vault = Vault.get_personal_vault()
Vault.create_folder(vault$id, "/My Folder")
## End(Not run)
```

Vault.datasets

Vault.datasets

## **Description**

Retrieves all datasets in a specific vault.

## Usage

```
Vault.datasets(id, env = .solveEnv, ...)
```

74 Vault.delete

## **Arguments**

id The ID of the vault.

env (optional) Custom client environment.

... (optional) Additional query parameters (e.g. limit, offset).

#### References

```
https://docs.solvebio.com/
```

# **Examples**

```
## Not run:
vault = Vault.get_personal_vault()
Vault.datasets(vault$id)
## End(Not run)
```

Vault.delete

Vault.delete

# Description

Delete a specific vault from SolveBio. This operation cannot be undone.

## Usage

```
Vault.delete(id, env = .solveEnv)
```

# **Arguments**

id String The ID of a SolveBio vault.
env (optional) Custom client environment.

#### References

```
https://docs.solvebio.com/
```

```
## Not run:
Vault.delete("1")
## End(Not run)
```

Vault.files 75

Vault.files

Vault.files

# Description

Retrieves all files in a specific vault.

## Usage

```
Vault.files(id, env = .solveEnv, ...)
```

# **Arguments**

id The ID of the vault.

env (optional) Custom client environment.

... (optional) Additional query parameters (e.g. limit, offset).

#### References

```
https://docs.solvebio.com/
```

# **Examples**

```
## Not run:
vault = Vault.get_personal_vault()
Vault.files(vault$id)
## End(Not run)
```

Vault.folders

Vault.folders

# **Description**

Retrieves all folders in a specific vault.

## Usage

```
Vault.folders(id, env = .solveEnv, ...)
```

## **Arguments**

```
id The ID of the vault.
```

env (optional) Custom client environment.

... (optional) Additional query parameters (e.g. limit, offset).

#### References

```
https://docs.solvebio.com/
```

# **Examples**

```
## Not run:
vault = Vault.get_personal_vault()
Vault.folders(vault$id)
## End(Not run)
```

## **Description**

Retrieves a specific vault by its full path (domain:vault).

# Usage

```
Vault.get_by_full_path(full_path, verbose = TRUE, env = .solveEnv)
```

# Arguments

full\_path The full path of a SolveBio vault.

verbose Print warning/error messages (default: TRUE).

env (optional) Custom client environment.

## References

```
https://docs.solvebio.com/
```

```
## Not run:
Vault.get_by_full_path("SolveBio:Public")
## End(Not run)
```

```
Vault.get\_or\_create\_by\_full\_path \\ \textit{Vault.get\_or\_create\_by\_full\_path}
```

## **Description**

Retrieves or creates a specific vault by its full path (domain:vault).

## Usage

```
Vault.get_or_create_by_full_path(full_path, env = .solveEnv, ...)
```

## **Arguments**

```
full_path The full path of a SolveBio vault.
env (optional) Custom client environment.
... (optional) Additional parameters.
```

#### References

```
https://docs.solvebio.com/
```

# **Examples**

```
## Not run:
Vault.get_or_create_by_full_path("My New Vault")
## End(Not run)
```

```
Vault.get_personal_vault
```

Vault.get\_personal\_vault

## **Description**

Retrieves the current users"s personal, private vault.

#### Usage

```
Vault.get_personal_vault(env = .solveEnv)
```

#### **Arguments**

env (optional) Custom client environment.

78 Vault.objects

#### References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
Vault.get_personal_vault()
## End(Not run)
```

Vault.objects

Vault.objects

# Description

Retrieves all objects in a specific vault.

# Usage

```
Vault.objects(id, env = .solveEnv, ...)
```

# Arguments

id The ID of the vault.

env (optional) Custom client environment.

... (optional) Additional query parameters (e.g. limit, offset).

## References

```
https://docs.solvebio.com/
```

```
## Not run:
vault = Vault.get_personal_vault()
Vault.objects(vault$id)
## End(Not run)
```

Vault.retrieve 79

Vault.retrieve

Vault.retrieve

## **Description**

Retrieves the metadata about a specific SolveBio vault.

## Usage

```
Vault.retrieve(id, env = .solveEnv)
```

## Arguments

id String The ID of a SolveBio vault env (optional) Custom client environment.

## References

```
https://docs.solvebio.com/
```

## **Examples**

```
## Not run:
Vault.retrieve("1234567890")
## End(Not run)
```

Vault.search

Vault.search

## Description

Search for objects in a specific vault.

## Usage

```
Vault.search(id, query, env = .solveEnv, ...)
```

# Arguments

id The ID of the vault. query The search query.

env (optional) Custom client environment.

... (optional) Additional query parameters (e.g. limit, offset).

80 Vault.update

#### References

```
https://docs.solvebio.com/
```

# **Examples**

```
## Not run:
vault = Vault.get_personal_vault()
Vault.search('test')
## End(Not run)
```

Vault.update

Vault.update

# Description

Updates the attributes of an existing vault.

# Usage

```
Vault.update(id, env = .solveEnv, ...)
```

# Arguments

id The ID of the vault to update.

env (optional) Custom client environment.

... Vault attributes to change.

# References

```
https://docs.solvebio.com/
```

# **Index**

Annotator.annotate, 4	DatasetCommit.delete, 29
Application.all, 5	DatasetCommit.retrieve, 30
Application.create, 5	DatasetExport.all, 31
Application.delete, 6	DatasetExport.create, 31
Application.retrieve, 7	DatasetExport.delete, 32
Application.update, 7	<pre>DatasetExport.get_download_url, 33</pre>
	DatasetExport.retrieve, 33
Beacon.all, 8	DatasetField.all, 34
Beacon.create, 9	DatasetField.create, 35
Beacon.delete, 10	DatasetField.facets, 35
Beacon.query, 10	DatasetField.retrieve, 36
Beacon.retrieve, 11	DatasetField.update, 37
Beacon.update, 12	DatasetImport.all, 37
BeaconSet.all, 12	DatasetImport.create, 38
BeaconSet.create, 13	DatasetImport.delete, 39
BeaconSet.delete, 14	DatasetImport.retrieve, 39
BeaconSet.query, 14	DatasetMigration.all,40
BeaconSet.retrieve, 15	DatasetMigration.create, 41
BeaconSet.update, 16	DatasetMigration.delete,42
	DatasetMigration.retrieve, 42
createEnv, 16	DatasetTemplate.all, 43
	DatasetTemplate.create, 44
Dataset.activity, 17	DatasetTemplate.delete,44
Dataset.all, 18	DatasetTemplate.retrieve, 45
Dataset.count, 18	DatasetTemplate.update, 46
Dataset.create, 19	
Dataset.data, 20	Expression.evaluate, 46
Dataset.delete, 20	
Dataset.disable_global_beacon, 21	GlobalSearch.facets,47
Dataset.enable_global_beacon, 22	GlobalSearch.request,48
Dataset.facets, 22	GlobalSearch.search,49
Dataset.fields, 23	GlobalSearch.subjects, 50
Dataset.get_by_full_path, 24	GlobalSearch.subjects_count, 51
Dataset.get_global_beacon_status, 24	
Dataset.get_or_create_by_full_path, 25	login, 51
Dataset.query, 26	
Dataset.retrieve, 27	Object.all, 52
Dataset.template, 27	Object.create, 53
Dataset.update, 28	Object.data, 54
DatasetCommit.all, 29	Object.delete, 54

82 INDEX

```
Object.disable_global_beacon, 55
Object.enable_global_beacon, 56
Object.fields, 56
Object.get_by_full_path, 57
Object.get_by_path, 58
Object.get_download_url,58
Object.get_global_beacon_status, 59
Object.get_or_upload_file, 60
Object.query, 61
Object.retrieve, 61
Object.update, 62
Object.upload_file, 63
protectedServer, 63
protectedServerJS, 64
SavedQuery.all, 65
SavedQuery.create, 66
SavedQuery.delete, 66
SavedQuery.retrieve, 67
SavedQuery.update, 67
Task.all.68
Task.follow, 69
Task.retrieve, 69
User.retrieve, 70
Vault.all, 71
Vault.create, 71
Vault.create_dataset, 72
Vault.create_folder, 73
Vault.datasets, 73
Vault.delete, 74
Vault.files, 75
Vault.folders, 75
Vault.get_by_full_path, 76
Vault.get_or_create_by_full_path, 77
Vault.get_personal_vault,77
Vault.objects, 78
Vault.retrieve, 79
Vault.search, 79
Vault.update, 80
```