

# Package ‘DataSpaceR’

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**Type** Package

**Title** Interface to 'the CAVD DataSpace'

**Version** 0.7.6

**Description** Provides a convenient API interface to access immunological data within 'the CAVD DataSpace'(<<https://dataspace.cavd.org>>), a data sharing and discovery tool that facilitates exploration of HIV immunological data from pre-clinical and clinical HIV vaccine studies.

**URL** <https://docs.ropensci.org/DataSpaceR/>,  
<https://github.com/ropensci/DataSpaceR>

**BugReports** <https://github.com/ropensci/DataSpaceR/issues>

**License** GPL-3

**Encoding** UTF-8

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jsonlite, data.table

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DataSpaceR-package	<i>DataSpaceR</i>
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### Description

DataSpaceR provides a convenient API for accessing datasets within the DataSpace database.

### Details

Uses the Rlabkey package to connect to DataSpace. Implements convenient methods for accessing datasets.

### Author(s)

Ju Yeong Kim

### See Also

[connectDS](#)

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checkNetrc	<i>Check netrc file</i>
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### Description

Check that there is a netrc file with a valid entry for the CAVD DataSpace.

### Usage

```
checkNetrc(netrcFile = getNetrcPath(), onStaging = FALSE, verbose = TRUE)
```

**Arguments**

netrcFile	A character. File path to netrc file to check.
onStaging	A logical. Whether to check the staging server instead of the production server.
verbose	A logical. Whether to print the extra details for troubleshooting.

**Value**

The name of the netrc file

**See Also**

[connectDS writeNetrc](#)

**Examples**

```
try(checkNetrc())
```

---

connectDS

*Create a connection to DataSpace*

---

**Description**

Constructor for [DataSpaceConnection](#)

**Usage**

```
connectDS(login = NULL, password = NULL, verbose = FALSE, onStaging = FALSE)
```

**Arguments**

login	A character. Optional argument. If there is no netrc file a temporary one can be written by passing login and password of an active DataSpace account.
password	A character. Optional. The password for the selected login.
verbose	A logical. Whether to print the extra details for troubleshooting.
onStaging	A logical. Whether to connect to the staging server instead of the production server.

**Details**

Instantiates an [DataSpaceConnection](#). The constructor will try to take the values of the various `labkey.*` parameters from the global environment. If they don't exist, it will use default values. These are assigned to 'options', which are then used by the [DataSpaceConnection](#) class.

**Value**

an instance of [DataSpaceConnection](#)

**See Also**[DataSpaceConnection](#)**Examples**

```
## Not run:
con <- connectDS()

## End(Not run)

con <- try(connectDS())
if (inherits(con, "try-error")) {
  warning("Read README for more information on how to set up a .netrc file.")
}
```

---

DataSpaceConnection    *The DataSpaceConnection class*

---

**Description**

The DataSpaceConnection class

The DataSpaceConnection class

**Constructor**[connectDS](#)**Active bindings**

`config` A list. Stores configuration of the connection object such as URL, path and username.

`availableStudies` A data.table. The table of available studies.

`availableGroups` A data.table. The table of available groups.

`availablePublications` A data.table. The table of available publications.

`mabGridSummary` A data.table. The filtered grid with updated `n_columns` and `geometric_mean_curve_ic50`.

`mabGrid` A data.table. The filtered mAb grid.

`virusMetadata` A data.table. Metadata about all viruses in the DataSpace.

`virusNameMappingTables` A list of data.table objects. This list contains ‘virusMetadataAll’, ‘virus-LabId’, and ‘virus\_synonym’ which are described in the vignette ‘Virus\_Name\_Mapping\_Tables’.

## Methods

### Public methods:

- [DataSpaceConnection\\$new\(\)](#)
- [DataSpaceConnection\\$print\(\)](#)
- [DataSpaceConnection\\$getStudy\(\)](#)
- [DataSpaceConnection\\$getGroup\(\)](#)
- [DataSpaceConnection\\$filterMabGrid\(\)](#)
- [DataSpaceConnection\\$resetMabGrid\(\)](#)
- [DataSpaceConnection\\$getMab\(\)](#)
- [DataSpaceConnection\\$downloadPublicationData\(\)](#)
- [DataSpaceConnection\\$refresh\(\)](#)
- [DataSpaceConnection\\$clone\(\)](#)

**Method** [new\(\)](#): Initialize a `DataSpaceConnection` object. See [connectDS](#).

*Usage:*

```
DataSpaceConnection$new(  
  login = NULL,  
  password = NULL,  
  verbose = FALSE,  
  onStaging = FALSE  
)
```

*Arguments:*

`login` A character. Optional argument. If there is no netrc file a temporary one can be written by passing login and password of an active DataSpace account.

`password` A character. Optional. The password for the selected login.

`verbose` A logical. Whether to print the extra details for troubleshooting.

`onStaging` A logical. Whether to connect to the staging server instead of the production server.

*Returns:* A new 'DataSpaceConnection' object.

**Method** [print\(\)](#): Print the `DataSpaceConnection` object.

*Usage:*

```
DataSpaceConnection$print()
```

**Method** [getStudy\(\)](#): Create a [DataSpaceStudy](#) object.

*Usage:*

```
DataSpaceConnection$getStudy(studyName)
```

*Arguments:*

`studyName` A character. Name of the study to retrieve.

**Method** [getGroup\(\)](#): Create a [DataSpaceStudy](#) object.

*Usage:*

```
DataSpaceConnection$getGroup(groupId)
```

*Arguments:*

groupId An integer. ID of the group to retrieve.

**Method filterMabGrid():** Filter rows in the mAb grid by specifying the values to keep in the columns found in the mabGrid field. It takes the column and the values and filters the underlying tables.

*Usage:*

```
DataSpaceConnection$filterMabGrid(using, value)
```

*Arguments:*

using A character. Name of the column to filter.

value A character vector. Values to keep in the mAb grid.

**Method resetMabGrid():** Reset the mAb grid to the unfiltered state.

*Usage:*

```
DataSpaceConnection$resetMabGrid()
```

**Method getMab():** Create a [DataSpaceMab](#) object.

*Usage:*

```
DataSpaceConnection$getMab()
```

**Method downloadPublicationData():** Download publication data for a chosen publication.

*Usage:*

```
DataSpaceConnection$downloadPublicationData(
  publicationId,
  outputDir = getwd(),
  unzip = TRUE,
  verbose = TRUE
)
```

*Arguments:*

publicationId A character/integer. ID for the publication to download data for.

outputDir A character. Path to directory to download publication data.

unzip A logical. If TRUE, unzip publication data to outputDir.

verbose A logical. Default TRUE.

**Method refresh():** Refresh the connection object to update available studies and groups.

*Usage:*

```
DataSpaceConnection$refresh()
```

**Method clone():** The objects of this class are cloneable with this method.

*Usage:*

```
DataSpaceConnection$clone(deep = FALSE)
```

*Arguments:*

deep Whether to make a deep clone.

## See Also

[connectDS DataSpaceR-package](#)

**Examples**

```

## Not run:
# Create a connection (Initiate a DataSpaceConnection object)
con <- connectDS()
con

# Connect to cvd408
# https://dataspace.cavd.org/cds/CAVD/app.view#learn/learn/Study/cvd408?q=408
cvd408 <- con$getStudy("cvd408")

# Connect to all studies
cvd <- con$getStudy("cvd408")

# Connect to the NYVAC durability comparison group
# https://dataspace.cavd.org/cds/CAVD/app.view#group/groupssummary/220
nyvac <- con$getGroup(220)

# Refresh the connection object to update available studies and groups
con$refresh()

## End(Not run)

```

---

DataSpaceMab

*The DataSpaceMab class*


---

**Description**

The DataSpaceMab class

The DataSpaceMab class

**Constructor**

DataSpaceConnection\$getMab()

**Active bindings**

`config` A list. Stores configuration of the connection object such as URL, path and username.

`studyAndMabs` A data.table. The table of available mAbs by study.

`mabs` A data.table. The table of available mAbs and their attributes.

`nabMab` A data.table. The table of mAbs and their neutralizing measurements against viruses.

`studies` A data.table. The table of available studies.

`assays` A data.table. The table of assay status by study.

`variableDefinitions` A data.table. The table of variable definitions.

## Methods

### Public methods:

- [DataSpaceMab\\$new\(\)](#)
- [DataSpaceMab\\$print\(\)](#)
- [DataSpaceMab\\$refresh\(\)](#)
- [DataSpaceMab\\$getLanlMetadata\(\)](#)
- [DataSpaceMab\\$clone\(\)](#)

**Method** `new()`: Initialize DataSpaceMab object. See [DataSpaceConnection](#).

*Usage:*

```
DataSpaceMab$new(mabMixture, filters, config)
```

*Arguments:*

`mabMixture` A character vector.

`filters` A list.

`config` A list.

**Method** `print()`: Print the DataSpaceMab object summary.

*Usage:*

```
DataSpaceMab$print()
```

**Method** `refresh()`: Refresh the DataSpaceMab object to update datasets.

*Usage:*

```
DataSpaceMab$refresh()
```

**Method** `getLanlMetadata()`: Applies LANL metadata to mabs table.

*Usage:*

```
DataSpaceMab$getLanlMetadata()
```

**Method** `clone()`: The objects of this class are cloneable with this method.

*Usage:*

```
DataSpaceMab$clone(deep = FALSE)
```

*Arguments:*

`deep` Whether to make a deep clone.

## See Also

[connectDS DataSpaceConnection](#)



**Examples**

```
## Not run:
# Create a connection (Initiate a DataSpaceConnection object)
con <- connectDS()

# Browse the mAb Grid
con$mabGridSummary

# Filter the grid by viruses
con$filterMabGrid(using = "virus", value = c("242-14", "Q23.17", "6535.3", "BaL.26", "DJ263.8"))

# Filter the grid by donor species (llama)
con$filterMabGrid(using = "donor_species", value = "llama")

# Check the updated grid
con$mabGridSummary

# Retrieve available viruses in the filtered grid
con$mabGrid[, unique(virus)]

# Retrieve available clades for 1H9 mAb mixture in the filtered grid
con$mabGrid[mab_mixture == "1H9", unique(clade)]

# Create a DataSpaceMab object that contains the filtered mAb data
mab <- con$getMab()
mab

# Inspect the `nabMab` field
mab$nabMab

## End(Not run)
```

---

DataSpaceStudy

*The DataSpaceStudy class*

---

**Description**

The DataSpaceStudy class

The DataSpaceStudy class

**Constructor**

DataSpaceConnection\$getStudy() DataSpaceConnection\$getGroup()

**Active bindings**

study A character. The study name.

config A list. Stores configuration of the connection object such as URL, path and username.

availableDatasets A data.table. The table of datasets available in the DataSpaceStudy object.

cache A list. Stores the data to avoid downloading the same tables multiple times.

dataDir A character. Default directory for storing nonstandard datasets. Set with setDataDir(dataDir).

treatmentArm A data.table. The table of treatment arm information for the connected study. Not available for all study connection.

group A character. The group name.

studyInfo A list. Stores the information about the study.

## Methods

### Public methods:

- [DataSpaceStudy\\$new\(\)](#)
- [DataSpaceStudy\\$print\(\)](#)
- [DataSpaceStudy\\$getDataset\(\)](#)
- [DataSpaceStudy\\$clearCache\(\)](#)
- [DataSpaceStudy\\$getDatasetDescription\(\)](#)
- [DataSpaceStudy\\$setDataDir\(\)](#)
- [DataSpaceStudy\\$refresh\(\)](#)
- [DataSpaceStudy\\$clone\(\)](#)

**Method new():** Initialize DataSpaceStudy class. See [DataSpaceConnection](#).

*Usage:*

```
DataSpaceStudy$new(study = NULL, config = NULL, group = NULL, studyInfo = NULL)
```

*Arguments:*

study A character. Name of the study to retrieve.

config A list. Stores configuration of the connection object such as URL, path and username.

group An integer. ID of the group to retrieve.

studyInfo A list. Stores the information about the study.

**Method print():** Print DataSpaceStudy class.

*Usage:*

```
DataSpaceStudy$print()
```

**Method getDataset():** Get a dataset from the connection.

*Usage:*

```
DataSpaceStudy$getDataset(
  datasetName,
  mergeExtra = FALSE,
  colFilter = NULL,
  reload = FALSE,
  outputDir = NULL,
  ...
)
```

*Arguments:*

`datasetName` A character. Name of the dataset to retrieve. Accepts the value in either the "name" or "label" field from `availableDatasets`.

`mergeExtra` A logical. If set to TRUE, merge extra information. Ignored for non-integrated datasets.

`colFilter` A matrix. A filter as returned by Rlabkey's `makeFilter`.

`reload` A logical. If set to TRUE, download the dataset, whether a cached version exist or not.

`outputDir` A character. Optional, specifies directory to download nonstandard datasets. If NULL, data will be downloaded to `dataDir`, set with `setDataDir(dataDir)`. If `dataDir` is not set, and `outputDir` is NULL, a tmp directory will be used.

... Extra arguments to be passed to `labkey.selectRows`

**Method** `clearCache()`: Clear cache. Remove downloaded datasets.

*Usage:*

```
DataSpaceStudy$clearCache()
```

**Method** `getDatasetDescription()`: Get variable information.

*Usage:*

```
DataSpaceStudy$getDatasetDescription(datasetName, outputDir = NULL)
```

*Arguments:*

`datasetName` A character. Name of the dataset to retrieve. Accepts the value in either the "name" or "label" field from `availableDatasets`.

`outputDir` A character. Directory path.

**Method** `setDataDir()`: Set default directory to download non-integrated datasets. If no `dataDir` is set, a tmp directory will be used.

*Usage:*

```
DataSpaceStudy$setDataDir(dataDir)
```

*Arguments:*

`dataDir` A character. Directory path.

**Method** `refresh()`: Refresh the study object to update available datasets and treatment info.

*Usage:*

```
DataSpaceStudy$refresh()
```

**Method** `clone()`: The objects of this class are cloneable with this method.

*Usage:*

```
DataSpaceStudy$clone(deep = FALSE)
```

*Arguments:*

`deep` Whether to make a deep clone.

## See Also

[connectDS DataSpaceConnection](#)

**Examples**

```
## Not run:
# Create a connection (Initiate a DataSpaceConnection object)
con <- connectDS()

# Connect to cvd408 (Initiate a DataSpaceStudy object)
# https://dataspace.cavd.org/cds/CAVD/app.view#learn/learn/Study/cvd408?q=408
cvd408 <- con$getStudy("cvd408")
cvd408

# Retrieve Neutralizing antibody dataset (NAb) for cvd408 from DataSpace
NAb <- cvd408$getDataset("NAb")

# Get variable information of the NAb dataset
cvd408$getDatasetDescription("NAb")

# Take a look at cvd408's treatment arm information
cvd408$treatmentArm

# Clear cache of a study object
cvd408$clearCache()

# Connect to the NYVAC durability comparison group
# https://dataspace.cavd.org/cds/CAVD/app.view#group/groupsummary/220
nyvac <- con$getGroup(220)

# Connect to all studies
cvd <- con$getStudy("")

# Refresh the study object to update available datasets and treatment info
cvd$refresh()

## End(Not run)
```

---

getNetrcPath

*Get a default netrc file path*

---

**Description**

Get a default netrc file path

**Usage**

```
getNetrcPath()
```

**Value**

A character vector containing the default netrc file path

## Examples

```
getNetrcPath()
```

---

writeNetrc	<i>Write a netrc file</i>
------------	---------------------------

---

## Description

Write a netrc file that is valid for accessing DataSpace.

## Usage

```
writeNetrc(  
    login,  
    password,  
    netrcFile = NULL,  
    onStaging = FALSE,  
    overwrite = FALSE  
)
```

## Arguments

login	A character. Email address used for logging in on DataSpace.
password	A character. Password associated with the login.
netrcFile	A character. Credentials will be written into that file. If left NULL, netrc will be written into a temporary file.
onStaging	A logical. Whether to connect to the staging server instead of the production server.
overwrite	A logical. Whether to overwrite the existing netrc file.

## Details

The database is accessed with the user's credentials. A netrc file storing login and password information is required. See [here](#) for instruction on how to register and set DataSpace credential. By default curl will look for the file in your home directory.

## Value

A character vector containing the netrc file path

## See Also

[connectDS](#) [checkNetrc](#)

**Examples**

```
# First, create an account in the DataSpace App and read the terms of use
# Next, create a netrc file using writeNetrc()
writeNetrc(
  login = "dataspaceuser@email.com",
  password = "yourSecretPassword"
)
# Specify `netrcFile = getNetrcPath()` to write netrc in the default path
```

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