Package 'ImmuneSpaceR'

October 16, 2019

Type Package

Title A Thin Wrapper around the ImmuneSpace Database

Version 1.12.2

Date 2014-06-12

VignetteBuilder knitr

Description Provides a convenient API for accessing data sets within ImmuneSpace (www.immunespace.org), the data repository and analysis platform of the Human Immunology Project Consortium (HIPC).

biocViews DataImport, DataRepresentation, ThirdPartyClient

URL https://github.com/RGLab/ImmuneSpaceR

BugReports https://github.com/RGLab/ImmuneSpaceR/issues

License GPL-2

Suggests knitr, testthat

Imports utils, R6, data.table, curl, httr, Rlabkey (>= 2.1.136), Biobase, pheatmap, ggplot2 (>= 3.2.0), scales, stats, gplots, plotly, heatmaply (>= 0.7.0), jsonlite, rmarkdown, preprocessCore, flowCore, flowWorkspace, digest

RoxygenNote 6.1.1

Collate 'CreateConnection.R' 'ISCon.R' 'ISCon-cytometry.R' 'ISCon-dataset.R' 'ISCon-geneExpression.R' 'ISCon-participantGroup.R' 'ISCon-plot.R' 'ISCon-utils.R' 'ImmuneSpaceR.R' 'netrc.R' 'template.R' 'theme.R' 'utils.R' 'zzz.R'

git_url https://git.bioconductor.org/packages/ImmuneSpaceR

git_branch RELEASE_3_9

git_last_commit fcb0494

git_last_commit_date 2019-07-09

Date/Publication 2019-10-15

Author Greg Finak [aut], Renan Sauteraud [aut], Mike Jiang [aut], Gil Guday [aut], Leo Dashevskiy [aut], Evan Henrich [aut], Ju Yeong Kim [aut], Lauren Wolfe [aut], Helen Miller [aut], Raphael Gottardo [aut], ImmuneSpace Package Maintainer [cre, cph]

Maintainer ImmuneSpace Package Maintainer <immunespace@gmail.com>

R topics documented:

ImmuneSpaceR-package	2
check_netrc	3
CreateConnection	3
ImmuneSpaceConnection	4
interactive_netrc	8
ISpalette	8
loadConnection	9
template_IS	9
theme_IS	10
write_netrc	11
	12

Index

ImmuneSpaceR-package A Thin Wrapper Around ImmuneSpace

Description

ImmuneSpaceR provides a convenient API for accessing data sets within the ImmuneSpace database.

Details

Uses the Rlabkey package to connect to ImmuneSpace. Implements caching, and convenient methods for accessing data sets.

See Also

CreateConnection

check_netrc

Check netrc file

Description

Check that there is a netrc file with a valid entry for ImmuneSpace.

Usage

check_netrc()

Details

In order to connect to ImmuneSpace, you will need a '.netrc' file in your contains a 'machine' name (hostname of ImmuneSpace), and 'login' and 'password'. See [here](https://www.labkey.org/wiki/home/Documentation/ for more information. By default RCurl will look for the file in your home directory.

If no netrc is available or it is not formatted properly, write_netrc can be used to write one. Otherwise, when specifying login and password in CreateConnection, a temporary file will be created for that connection.

Value

The name of the netrc file

See Also

CreateConnection write_netrc

Examples

try(check_netrc())

CreateConnection CreateConnection

Description

Constructor for ImmuneSpaceConnection class.

Usage

```
CreateConnection(study = NULL, login = NULL, password = NULL,
verbose = FALSE, onTest = FALSE)
```

Arguments

study	A "character" vector naming the study.
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 ImmuneSpace account.
password	A "character". Optional. The password for the selected login.
verbose	A "logical" whether to print the extra details for troubleshooting.
onTest	A "logical" whether to connect to the test server (https://test.immunespace.org/) instead of the production server (https://www.immunespace.org/).

Details

Instantiates an ImmuneSpaceConnection for study 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 ImmuneSpaceConnection class.

Value

an instance of an ImmuneSpaceConnection

See Also

ImmuneSpaceConnection

Examples

```
## Not run:
# Single study
con <- CreateConnection("SDY269")
# Cross study
con <- CreateConnection("")
## End(Not run)
sdy <- try(CreateConnection("SDY269"))
if (inherits(sdy, "try-error")) {
  warning("Read the Introduction vignette for more information on how to set
up a .netrc file.")
}
```

ImmuneSpaceConnection The ImmuneSpaceConnection class

Description

A connection respresents a study or a set of studies available on ImmuneSpace. It provides function to download and display the data within these studies.

Usage

ISCon

ImmuneSpaceConnection

Format

An object of class R6ClassGenerator of length 24.

Details

The ImmuneSpaceConnection will initialize itself, and look for a .netrc file in "~/" the user's home directory. The .netrc file should contain a machine, login, and password entry to allow access to ImmuneSpace, where machine is the host name like "www.immunespace.org".

It can also use global variables labkey.url.base, and labkey.url.path, to access a study. labkey.url.base should be https://www.immunespace.org/.labkey.url.path should be /Studies/studyname, where 'studyname' is the accession number of the study.

Value

An instance of an ImmuneSpaceConnection for a study in labkey.url.path.

Constructor

CreateConnection

Fields

study A character. The study accession number. Use an empty string ("") to create a connection at the project level.

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

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

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

Methods

initialize() Initialize ImmuneSpaceConnection class. See CreateConnection.

print() Print ImmuneSpaceConnection class.

- listDatasets(output = c("datasets", "expression")) Lists the datasets available in the study
 or studies of the connection.
- listGEMatrices(verbose = FALSE, reload = FALSE) Lists available gene expression matrices
 for the connection.

verbose: A logical. If TRUE, whether to print the extra details for troubleshooting.

reload: A logical. If TRUE, retrieve the table of available gene expression matrices whether a cached version exist or not.

- listGEAnalysis() Lists available gene expression analysis for the connection.
- listParticipantGroups() Lists available participant groups on the ImmuneSpace portal.
- listWorkspaces(reload = FALSE) Lists available workspaces for the connection.

reload: A logical. If TRUE, download the table whether a cached version exist or not.

listGatingSets(reload = FALSE) Lists available gating sets for the connection.

reload: A logical. If TRUE, download the table whether a cached version exist or not.

- summarizeCyto() Prints a summary of cytometry data for the connection.
- summarizeGatingSet(gatingSet) Prints a summary of a gating set. Note that this method currently works only in the ImmuneSpace RStudio session.

gatingSet: A character. The name of the gating set to summarize.

loadGatingSet(gatingSet) Loads a gating set via flowWorkspace::load_gs to the current environment. Note that this method currently works only in the ImmuneSpace RStudio Docker session.

gatingSet: A character. The name of the gating set to load.

- getDataset(x, original_view = FALSE, reload = FALSE, colFilter = NULL, ...) Get a dataset
 form the connection.
 - x: A character. The name of the dataset to download.
 - original_view: A logical. If TRUE, download the original ImmPort view; else, download the default grid view.
 - reload: A logical. If TRUE, download the dataset whether a cached version exist or not.
 - colFilter: A character. A filter as returned by Rlabkey's makeFilter function.
 - ...: Extra arguments to be passed to labkey.selectRows.
- getGEMatrix(matrixName = NULL, cohortType = NULL, outputType = "summary", annotation = "latest", reloa Downloads a probe-level or gene-symbol summarized expression matrix from ImmuneSpace and constructs an ExpressionSet. Use experimentData() on the resulting ExpressionSet object to see version info for annotation.

matrixName: A character. The name of the gene expression matrix to download.

cohortType: A character. The name of a cohortType that has an associated gene expression matrix. Note that if this argument is not NULL, then matrixName is ignored. CohortType is a concatenation of "cohort" and "cell type" that allows the user to specify a matrix for the cell type subset of a cohort.

outputType: A character. one of 'raw', 'normalized' or 'summary'. If 'raw', returns an expression matrix of non-normalized values by probe. 'normalized' returns normalized values by probe. 'summary' returns normalized values averaged by gene symbol.

annotation: A character. one of 'default', 'latest', or 'ImmSig'. Determines which feature annotation set (FAS) is used. 'default' uses the FAS from when the matrix was generated. latest' uses a recently updated FAS based on the original. 'ImmSig' is specific to studies involved in the ImmuneSignatures project and uses the annotation from when the meta-study's manuscript was created.

reload: A logical. If set to TRUE, the matrix will be downloaded again, even if a cached cop exist in the ImmuneSpaceConnection object.

verbose: A logical. If set to TRUE, notes on how the expressionSet object was created will be printed, including normalization, summarization, feature_annotation_set, and alias2symbol mapping version of org.Hs.eg.db.

getGEAnalysis(...) Downloads data from the gene expression analysis results table.

...: A list of arguments to be passed to labkey.selectRows.

- getGEInputs() Downloads data from the gene expression input samples table.

group: A character or integer. Call con\$listParticipantGroups() to see available participants groups. Use group_id or group_name as input.

dataType: A character. Use con\$availableDatasets to see available dataset names.

downloadGEFiles(files, destdir = ".") Downloads gene expression raw data files.

files: A character. Filenames as shown on the gene_expression_files dataset.

destdir: A character. The local path to store the downloaded files.

addTreatment(expressionSet) Adds treatment information to the phenoData of an Expression-Set.

expressionSet: An ExpressionSet. The ExpressionSet object that has been downloaded from the connection.

mapSampleNames(EM = NULL, colType = "participant_id") Changes the sampleNames of an ExpressionSet fetched by getGEMatrix using the information in the phenodData slot. EM: An ExpressionSet, as returned by getGEMatrix.

colType: A character. The type of column names. Valid options are 'expsample_accession' and 'participant_id'.

plot(...) Visualizes a selected dataset. This method is used by the DataExplorer module on the ImmuneSpace portal.

dataset: A character. The name of the dataset to plot, as displayed by the listDataset method. normalize_to_baseline: A logical. If TRUE, the values are plotted as log2 fold-change from baseline.

type: A character. The type of plot. Valid choices are 'auto', 'heatmap', 'boxplot', 'lineplot', 'violinplot'. If set to 'auto', the function will select an appropriate plot type for the selected data.

filter: A filter as created by the makeFilter function from Rlabkey.

facet: The facetting for ggplot2 based plots. Valid choices are 'grid' and 'wrap'.

text_size: The size of all text elements in the plot.

legend: A character. Columns of the dataset or demographics to be added as legend on the heatmap. This argument is ignored if the plot type isn't heatmap.

show_virus_strain: A logical. Should all the virus strains be shown or should the values be averaged. Only used when dataset = 'hai'.

interactive: A logical. If TRUE, an interactive plot will be created. The default is FALSE.

...: Extra argument to be passed to ggplot. e.g: shape = 'Age', color = 'Race'.

clearCache() Clears the cache. Removes downloaded datasets and expression matrices.

See Also

CreateConnection ImmuneSpaceR-package

Examples

```
## Not run:
# Create a connection (Initiate a ImmuneSpaceConnection object)
sdy269 <- CreateConnection("SDY269")</pre>
# Print the connection object
sdy269
# Retrieve the HAI dataset
HAI <- sdy269$getDataset("hai")</pre>
# Fetch a summarized gene expresssion matrix with latest annotation
LAIV <- sdy269$getGEMatrix("LAIV_2008")</pre>
# Visualize the ELISA dataset
sdy269$plot("elisa")
## End(Not run)
sdy <- try(CreateConnection("SDY269"))</pre>
if (inherits(sdy, "try-error")) {
 warning("Read the Introduction vignette for more information on how to set
up a .netrc file.")
}
```

interactive_netrc Interactively write a netrc file

Description

Write a netrc file that is valid for accessing ImmuneSpace

Usage

```
interactive_netrc()
```

Value

A netrc file that is verified to connect to ImmuneSpace

Examples

```
## Not run:
interactive_netrc()
```

End(Not run)

ISpalette ImmuneSpace palette

Description

Create a color gradient of the selected length that matches the ImmuneSpace theme.

Usage

ISpalette(n)

Arguments

n

A numeric. The length of the desired palette.

Value

A character vector colors in hexadecimal code of length n.

Examples

plot(1:10, col = ISpalette(10), cex = 10, pch = 16)

loadConnection Save/Load an ImmuneSpaceConnection object from disk

Description

Connection can hold a lot of data in cache. If a lot of work has been done (e.g: lots of downloaded datasets and gene-expression matrices), it can be useful to save the connection for later work or even offline use.

Usage

```
loadConnection(file)
```

saveConnection(con, file)

Arguments

file	The file name to be saved to or loaded from
con	An ${\tt ImmuneSpaceConnection}.$ The connection to save to file. To be loaded later using loadConnection.

Value

An ImmuneSpaceConnection object

Examples

```
# Sample saved connection with pre-downloaded expression matrices and datasets
saved <- system.file("extdata/saved_con.rds", package = "ImmuneSpaceR")
new_con <- loadConnection(saved)
new_con
names(new_con$cache)
## Not run:
saveConnection(new_con, tempfile())
```

End(Not run)

template_IS template_IS

Description

A HTML template for knitted reports that matches ImmuneSpace's graphic style. It is based on html_document from the **rmarkdown** package with css, theme, and template parameters disabled.

Usage

template_IS(...)

Arguments

See html_document

Details

See the documentation for html_document or the oneline documentation for additional details on using the html_document format. Compared to html_document, it:

- uses a custom css stylesheet
- · does not use bootstrap themes

Value

R Markdown output format to pass to render

Examples

```
## Not run:
library(ImmuneSpaceR)
rmarkdown::render("input.Rmd", template_IS())
rmarkdown::render("input.Rmd", template_IS(toc = TRUE))
```

```
## End(Not run)
template_IS()
```

theme_IS theme_IS

Description

Theme that matches ImmuneSpace's graphic style. The theme modifies the background, the grid lines, the axis, and the colors used by continuous and gradient scales.

Usage

theme_IS(base_size = 12)

Arguments

base_size A numeric. Base font size.

Details

List of modified ggplot2 elements: panel.background, panel.grid.major, panel.grid.minor, axis.ticks, axis.line.x, axis.line.y, plot.title, and strip.background.

The default scale_fill_gradient, scale_fill_continuous, scale_colour_gradient and scale_colour_contine are also replaced by a custom scale.

Value

A theme object

write_netrc

Examples

```
library(ggplot2)
p <- ggplot(data = mtcars) + geom_point(aes(x = mpg, y = cyl, color = hp)) + facet_grid(vs ~ am)
p + theme_IS()</pre>
```

write_netrc Write a netrc file

Description

Write a netrc file that is valid for accessing ImmuneSpace

Usage

```
write_netrc(login, password, machine = "www.immunespace.org",
    file = NULL)
```

Arguments

login	A character. The email address used for loging in on ImmuneSpace.
password	A character. The password associated with the login.
machine	A character. The server to connect.
file	A character. The credentials will be written into that file. If left NULL, the netrc will be written into a temporary file.

Value

A character vector containing the file paths for netrc

Examples

```
write_netrc("immunespaceuser@gmail.com", "mypassword")
```

Index

```
*Topic datasets
    ImmuneSpaceConnection, 4
addTreatmentt (ImmuneSpaceConnection), 4
check_netrc, 3
CreateConnection, 2, 3, 5, 7
getDataset (ImmuneSpaceConnection), 4
getGEAnalysis (ImmuneSpaceConnection), 4
getGEFiles (ImmuneSpaceConnection), 4
getGEInputs (ImmuneSpaceConnection), 4
getGEMatrix (ImmuneSpaceConnection), 4
getParticipantData
        (ImmuneSpaceConnection), 4
html_document, 9, 10
ImmuneSpaceConnection, 4, 4
ImmuneSpaceR (ImmuneSpaceR-package), 2
ImmuneSpaceR-package, 2
interactive_netrc, 8
ISCon (ImmuneSpaceConnection), 4
ISpalette, 8
listDatasets (ImmuneSpaceConnection), 4
listGatingSets (ImmuneSpaceConnection),
        4
listGEAnalysis(ImmuneSpaceConnection),
        4
listGEMatrices (ImmuneSpaceConnection),
        4
listParticipantGroups
        (ImmuneSpaceConnection), 4
listWorkspaces (ImmuneSpaceConnection),
        4
loadConnection, 9
loadGatingSet(ImmuneSpaceConnection), 4
mapSampleNames (ImmuneSpaceConnection),
        4
render, 10
saveConnection (loadConnection), 9
```

template_IS, 9
theme_IS, 10

write_netrc, 11