

Package ‘gCMAPWeb’

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Type Package

Title A web interface for gene-set enrichment analyses

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Description The gCMAPWeb R package provides a graphical user interface for the gCMAP package. gCMAPWeb uses the Rook package and can be used either on a local machine, leveraging R's internal web server, or run on a dedicated rApache web server installation. gCMAPWeb allows users to search their own data sources and instructions to generate reference datasets from public repositories are included with the package. The package supports three common types of analyses, specifically queries with 1. one or two sets of query gene identifiers, whose members are expected to show changes in gene expression in a consistent direction. For example, an up-regulated gene set might contain genes activated by a transcription factor, a down-regulated geneset targets repressed by the same factor. 2. a single set of query gene identifiers, whose members are expected to show divergent differential expression (non-directional query). For example, members of a particular signaling pathway, some of which may be up- some down-regulated in response to a stimulus. 3. a query with the complete results of a differential expression profiling experiment. For example, gene identifiers and z-scores from a previous perturbation experiment. gCMAPWeb accepts three types of identifiers: EntrezIDs, gene Symbols and microarray probe ids and can be configured to work with any species supported by Bioconductor. For each query submission, significantly similar reference datasets will be identified and reported in graphical and tabular form.

Note gCMAPWeb is not compatible with the RStudio IDE, but can be used from the command line or via R's standard GUI.

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Depends Biobase, gCMAP (>= 1.3.0), methods, R (>= 3.3.0), Rook

Imports brew, BiocGenerics, annotate, AnnotationDbi, grDevices, GSEABase, hwriter, parallel, yaml

Suggests affy, ArrayExpress, hgfocus.db, hgu133a.db, mgug4104a.db,
org.Hs.eg.db, org.Mm.eg.db, RUnit

Enhances bigmemory, bigmemoryExtras

biocViews GUI, GeneSetEnrichment, Visualization

LazyLoad yes

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'cmap.R' 'cmapAnalysis.R' 'cmapHeatmap.R' 'createObject.R'
'example_objects.R' 'file_io.R' 'html_elements.R'
'plotting_functions.R' 'reporting.R'

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gCMAPWeb-packageA web-interface for gene-set enrichment analysis

Description

A web-interface for gene-set enrichment analysis

Details

This package provides a stand-alone web interface for gene-set enrichment analysis. Run either through R's internal Rhttpd server or deployed through an Apache2 webserver, gCMAPWeb allows users to search reference datasets stored in `NChannelSet` or `CMAPICollection` objects via three different methods:

- Directional queries performed by calculating the parametric JG score from the reference datasets
- Non-directional queries performed via Fisher's exact test
- Profile queries performed by calculating the parametric JG score from the query data

A web interface is populated based on a configuration file in YAML format and additional information from the reference dataset objects (if available). For each query, a report is generated in html format, including graphs and tabular output. Extensive customization of the graphical user interface can easily be performed through global options.

Author(s)

Thomas Sandmann

References

Jiang Z, Gentleman R., Extensions to gene set enrichment. Bioinformatics. 2007 Feb 1;23(3):306-13 <http://www.ncbi.nlm.nih.gov/pubmed/17127676>

Examples

```
example( gCMAPWeb )
```

addLinks

Adding links to columns of a data.frame

Description

This function identifies column pairs from a data.frame that differ only in a string suffix (default: `.url`). The column with the suffix is used to add http href tags to the other column and is then removed.

Usage

```
addLinks(df, pattern = ".url$", swap.colnames =getOption("swap.colnames",
  default = list(padj = "FDR", nFound = "Genes")))
```

Arguments

<code>df</code>	data.frame, must contain both target and url columns
<code>pattern</code>	character, the suffix linking target and url columns. Default=c('url')
<code>swap.colnames</code>	list, containing alternative names for CMAPResults columns.

Details

To annotate columns with links to other urls, the urls must be included in the data.frame in a separate column matching the target column name but carrying the additional suffix in the column name. For example, to add links to the 'id' column, an additional id.url column can be used. Multiple columns can be processed simultaneously, if each of them has a matching url column.

Value

Data frame with href html tags in the target column(s)

Author(s)

Thomas Sandmann

add_geneScores_to_df *CMAPResults data.frame with gene-level scores*

Description

This function converts a CMAPResults object into a data.frame and adds the gene.level scores as an additional column

Usage

```
add_geneScores_to_df(cmap.result)
```

Arguments

cmap.result CMAPResults object

Value

A data.frame with gene-level scores in column 'Scores'

Author(s)

Thomas Sandmann

body_first *Definition of html navigation bar elements*

Description

Shared html code, constituting the first part of the body section

Usage

```
body_first(url.base = NULL)
```

Arguments

url.base character, path to htdocs directory (optional)

Value

Html code as a character string

Author(s)

Thomas Sandmann

cmap-data

*Example reference datasets***Description**

Five example reference datasets containing simulated data to demonstrate the functionality of the gCMAPWeb package. These objects were generated with the [exampleCMAP](#) function using different seeds.

Format

cmap1 to 4 are [NChannelSet](#) objects each with 1000 simulated z-scores for 10 samples. cmap5 is a [CMAPCollection](#) with membership information about 1000 genes and 10 gene sets.

Details

cmap1, cmap2, cmap3 and cmap5 contain data for 1000 human Entrez identifiers. cmap1, cmap2 and cmap3 are [NChannelSet](#) objects with one channel, the 'z' AssayDataElement. cmap4 is an [NChannelSet](#) with data for 1000 mouse Entrez identifiers in the 'z' and 'log_fc' channels. cmap5 is a code[CMAPCollection](#).

cmapAnalysis

*Function controlling the gCMAP analysis workflow***Description**

This function controls the gCMAPWeb analysis workflow

Usage

```
cmapAnalysis(req, conf_data, reference.cmmaps, element =getOption("element",
  default = "z"), save.intermediates =getOption("save.intermediates", default
  = FALSE))
```

Arguments

req	a Rook Request object
conf_data	list, the configuration data as returned by the <code>read_config_file</code> function
reference.cmmaps	list, containing the reference database eSet objects
element	character, identifying the <code>elementName</code> of the channel extracted from <code>NChannelSet</code> objects
save.intermediates	logical, if TRUE rdata files of intermediate results will be stored in the temporary directory for debugging

Value

a list with three elements: conversion, reports and tmp_filename

Author(s)

Thomas Sandmann

cmapHeatmap

Function to create an annotated heatmap of gene scores

Description

This function takes a numerical matrix (with samples in columns and genes in rows), preprocesses the data (if desired), determines the optimal height for the heatmap and calls the .ImagePlot function to create the final heatmap figure.

Usage

```
cmapHeatmap(x, reference.name, col.anno = NULL, row.anno = NULL,
            file.name = "heatmap", url.base = NULL, main = "Query gene scores",
            ColorRamp = colorRampPalette(c("#044381FF", "grey95", "grey95",
            "firebrick"))(100), col.col = c(down = "black", up = "grey"),
            row.col = c(correlated = "#1B9E77", anticorrelated = "#044381FF", over =
            "#1B9E77", under = "#044381FF"), order.by.score = TRUE,
            cluster.rows = TRUE, score.cap = c(-5, 5),
            ylab = "Significant datasets")
```

Arguments

x	numerical matrix with samples in rows and genes in columns.
reference.name	character, names of the reference cmap, used to construct the html image reference
col.anno	character vector with column annotations to be displayed as (horizontal) annotation bar above the heatmap. If not NULL, must contain one element for each column of 'x'.
row.anno	character vector with row annotations to be displayed as (vertical) annotation to the right of the heatmap. If not NULL, must contain one element for each row of 'x'.
file.name	character, the path and filename (without suffix) to save the png file to
url.base	character, prefix for the html image reference
main	Character, main title of the plot
ColorRamp	vector of colors used for the heatmap, e.g. generated by a call to colorRampPalette
col.col	named vector with a color for each level of col.anno (e.g. c(up="firebrick", down="blue"))
row.col	named vector with a color for each level of row.anno (e.g. c(correlated="firebrick", anticorrelated="#044381FF", over = "#1B9E77", under = "#044381FF"))
order.by.score	logical, should gene scores be reordered independently for each sample ?
cluster.rows	logical, perform hierarchical clustering on significant gene sets ?
score.cap	numerical vector of length two, specifying the limits of the color scale. Scores > max(score.cap) or < min(score.cap) will be set to score.cap. Default: c(-5,5)
ylab	character, y-axis label

Value

list with two elements, 1. image.html: a character string with html code with the image tag referring to the output png 2. row.order: integer vector with thw row.order obtained after hierarchical clustering or NULL if no clustering was performed

Author(s)

Thomas Sandmann

cmapRun

cmapRun

Description

This S4 method dispatches a gCMAP analysis and selects a suitable gene-set-enrichment analysis method based on the class of its arguments.

Usage

```
cmapRun(user.input, cmap, ...)

## S4 method for signature 'CMAPCollection,CMAPCollection'
cmapRun(user.input, cmap)

## S4 method for signature 'GeneSet,CMAPCollection'
cmapRun(user.input, cmap, ...)

## S4 method for signature 'GeneSet,eSet'
cmapRun(user.input, cmap, ...)

## S4 method for signature 'CMAPCollection,eSet'
cmapRun(user.input, cmap,
        lower = getOption("lower.threshold", default = -3),
        higher = getOption("higher.threshold", default = 3),
        element = getOption("element", default = "z"),
        min.set.size = getOption("min.set.size", default = 5), keep.scores = TRUE)

## S4 method for signature 'SignedGeneSet,eSet'
cmapRun(user.input, cmap,
        element = getOption("element", default = "z"))

## S4 method for signature 'eSet,CMAPCollection'
cmapRun(user.input, cmap,
        element = getOption("element", default = "z"))

## S4 method for signature 'eSet,eSet'
cmapRun(user.input, cmap,
        lower = getOption("lower.threshold", default = -3),
        higher = getOption("higher.threshold", default = 3),
        element = getOption("element", default = "z"))
```

Arguments

<code>user.input</code>	An object of class GeneSet, SignedGeneSet, eSet or CMAPCollection with the user-provided query.
<code>cmap</code>	The reference database, a CMAPCollection or eSet object.
<code>...</code>	Arguments specific to individual methods
<code>lower</code>	Numeric, when reference databases are thresholded, genes with scores less than 'lower' are considered down-regulated. Default: <code>getOption("lower.threshold", default=-3")</code>
<code>higher</code>	Numeric, when reference databases are thresholded, genes with scores larger than 'higher' are considered up-regulated. Default: <code>getOption("higher.threshold", default=3")</code>
<code>element</code>	Character, specifying which channel / assayDataElement of the reference database to query. Default: <code>getOption("element", default="z")</code>
<code>min.set.size</code>	integer, gene sets with less than <code>min.set.size</code> members will be dropped from CMAPCollections. Default: <code>getOption("min.set.size", default=5)</code>
<code>keep.scores</code>	Scalar boolean, include scores in the result object ?

Details

The `cmapRun` methods choose one of three approaches for gene-set-enrichment analyses, depending on the nature of the user-provided query and the data available in the reference databases.

1. for non-directional gene set queries (e.g. a list of gene identifiers) a Fisher exact test is performed (with a call to the `fisher_score` method from the gCMAP package) to evaluate the overlap with significantly up- or down-regulated gene sets from the reference database. If the reference object is an NChannelSet, it is thresholded on the fly to obtain a set of up- or down-regulated genes. If the reference database is a CMAPCollection, the gene sets are used directly. Please note that the Fisher exact test does not take into account information about the directionality of gene regulation (potentially) available in the reference database.
2. for directional gene set queries (e.g. two separate lists of up- and down-regulated genes of interest) the gene expression scores available in the reference database are summarized as the JG score by running the `gsealm_jg_score` method from the gCMAP package.
3. for queries with complete differential gene expression profiles, directional gene sets are induced from the reference database. As above, the JG score is used to summarize the expression changes for each gene set, but this time the sets are derived from the database, while the scores are provided by the user.

This method is called by the `cmapAnalysis` function of the gCMAPWeb package.

Please note: Most of the parameters of the `cmapRun` methods can be set as global options.

Value

A CMAPResults object

See Also

[gsealm_jg_score](#) and [fisher_score](#)

`conversion_html` *Identifier conversion html output*

Description

This function formats a data.frame with the results of the gene identifier conversion to html code, including an interactive dataTable. The html page is saved to disk and the path to the output file is returned as a character string.

Usage

```
conversion_html(df, result.dir, file.name, tmp_filename, url.base = NULL,
  message = NULL)
```

Arguments

<code>df</code>	data.frame with old / new gene identifiers
<code>result.dir</code>	character, path to result directory
<code>file.name</code>	character, output filename
<code>tmp_filename</code>	character, name of the session-specific subdirectory
<code>url.base</code>	character, path to htdocs directory (optional)
<code>message</code>	character, optional message to display above the result table

Value

character, name of the output html file

Author(s)

Thomas Sandmann

`convert_gene_identifiers`
Entrez mapper

Description

This function maps the submitted gene identifiers to Entrez identifiers and returns a translation table with the original and mapped ids

Usage

```
convert_gene_identifiers(gene.ids, query.IdType, species)
```

Arguments

<code>gene.ids</code>	Character vector with gene ids
<code>query.IdType</code>	A GeneIdentifierType object
<code>species</code>	String identifying the species, will be inserted into a new EntrezIdentifier object and returned with the translated results

Value

A list with the following elements

- gene.ids, a character vector of unique EntrezIds
- query.IdType, an EntrezIdentifier object with the species
- conversion, a list containing the full translation table and a summary message
- original.ids, a character vector of the submitted ids corresponding to the returned EntrezIds

Author(s)

Thomas Sandmann

create_figure_legend *This function generates the figure legend, with a separate dom identifier for each output panel*

Description

This function generates the figure legend, with a separate dom identifier for each output panel

Usage

```
create_figure_legend(reference.name)
```

Arguments

reference.name character, name of the reference database used

Value

character string with the html code element required to insert the figure legend into the html report

Author(s)

Thomas Sandmann

create_GeneIdentifierType
GeneIdentifierType creator

Description

This function extracts the species or platform information from the POST request and constructs GeneIdentifierType objects

Usage

```
create_GeneIdentifierType(post, conf_data)
```

Arguments

post	list, POST component of the Rook request
conf_data	list, the configuration data as returned by the read_config_file function

Value

GeneIdentifierType object

Author(s)

Thomas Sandmann

create_GeneSet	<i>GeneSet creator</i>
----------------	------------------------

Description

This function creates a GeneSet object from user-specified identifiers

Usage

```
create_GeneSet(post, conf_data)
```

Arguments

post	list, POST component of the Rook request
conf_data	list, the configuration data as returned by the read_config_file function

Value

GeneSet object

Author(s)

Thomas Sandmann

create_gene_report	<i>This function compiles a gene-level report from a CMAPResults object.</i>
--------------------	--

Description

Usually called by the generate_report function

Usage

```
create_gene_report(cmap.result, query, result.dir, url.base, reference.cmap,
  figure.dir, element =getOption("element", default = "z"),
  annotation.db = "org.Hs.eg.db",
  gene.level.plot =getOption("gene.level.plot", default = TRUE))
```

Arguments

cmap.result	a CMAPResults object
query	a GeneSet, SignedGeneSet or ExpressionSet object used as query for the gene-set-enrichment analysis giving rise to 'cmap.result'
result.dir	character, path to gene_results folder in the per-session output directory
url.base	character, the name of the reference database
reference.cmap	eSet, the reference database used for the gCMAP analysis
figure.dir	character, path to figure folder in the per-session output directory
element	character, the assayDataElementName to extract from NChannelSet objects
annotation.db	character, the name of the annotation package used to lookup gene identifiers
gene.level.plot,	logical: should gene-level plots be included in the report ?

Value

character string with the relative url to the report html, which is directly written to disk.

Author(s)

Thomas Sandmann

create_gene_table_legend

This function reads an eSet object and returns the html code to display a legend for all columns that have more descriptive annotations in the varMetadata slot.

Description

This function reads an eSet object and returns the html code to display a legend for all columns that have more descriptive annotations in the varMetadata slot.

Usage

```
create_gene_table_legend(col.names, eset)
```

Arguments

col.names	character, a vector with column names that are always included in the legend
eset	eSet object

Value

character string with the html code element required to insert the legend into the html report

Author(s)

Thomas Sandmann

`create_heatmap_legend` *Legend for overview heatmap*

Description

Content for the legend of the overview heatmap plot displayed on the main results page for directional and non-directional queries

Usage

```
create_heatmap_legend(text)
```

Arguments

<code>text</code>	Character, text for the legend of the density plot the main report page. Can be set as the global variabel "gene.set.legend".
-------------------	---

Value

Character

Author(s)

Thomas Sandmann

`create_legend`

This function reads a CMAPResults object and returns the html code to display a legend for all columns that have more descriptive annotations in the varMetadata slot. A standard set of columns is selected for display in the legend through the 'keep' parameter. In addition, all columns with annotations in the labelDescription field of the varMetadata slot are included if the field content differs from the column name.

Description

This function reads a CMAPResults object and returns the html code to display a legend for all columns that have more descriptive annotations in the varMetadata slot. A standard set of columns is selected for display in the legend through the 'keep' parameter. In addition, all columns with annotations in the labelDescription field of the varMetadata slot are included if the field content differs from the column name.

Usage

```
create_legend(res, reference.name, keep = c("set", "trend", "FDR", "effect",
  "nSet", "Genes"), swap.colnames = getOption("swap.colnames", default =
  list(padj = "FDR", nFound = "Genes")))
```

Arguments

res	CMAPResults object
reference.name	character, name of the reference database used
keep	character, a vector with column names that are always included in the legend
swap.colnames	list, containing alternative names for CMAPResults columns.

Value

character string with the html code element required to insert the legend into the html report

Author(s)

Thomas Sandmann

create_overview_legend

Legend for overview density plot

Description

Content for the legend of the overview density plot displayed on every main results page

Usage

create_overview_legend(text)

Arguments

text	Character, text for the legend of the density plot the main report page. Can be set as the global variabel "gene.set.legend".
------	---

Value

Character

Author(s)

Thomas Sandmann

`create_overview_plot` *Overview plot*

Description

This function creates an overview density plot summarizing the similarity scores obtained for all instances in the reference database.

Usage

```
create_overview_plot(effect.sample, effect.population, file.name,
                     reference.name, main = "Distribution of similarity scores",
                     xlab = "Effect size", col.set = "black", col.up = "#1B9E77",
                     col.down = "blue", strip.cutoffs = c(-3, 3), strip.bounds = c(-6, 6),
                     strip.col = c("#1B9E77", "white", "blue"), url.base = NULL,
                     up.label = "Correlated", down.label = "Anti-correlated")
```

Arguments

<code>effect.sample</code>	numeric, the scores for all gene sets in the reference database deemed significantly similar
<code>effect.population</code>	numeric, the scores for all gene sites in the reference database
<code>file.name</code>	character, path and name of the output file
<code>reference.name</code>	name of the reference database
<code>main</code>	main title of the plot
<code>xlab</code>	x-axis label
<code>col.set</code>	color used for the significant samples in the density plot
<code>col.up</code>	color used to indicate significantly correlated / enriched gene sets in the rug plot
<code>col.down</code>	color used to indicate significantly anti-correlated / depleted gene sets in the rug plot
<code>strip.cutoffs</code>	numeric vector, indicating the area of the heat-strip set to the intermediate strip.col. Default=c(-3,3)
<code>strip.bounds</code>	numeric vector, indicating the cutoffs above / below which the color of the heat-strip is set to maximum. Default=c(-6,6)
<code>strip.col</code>	character vector with three elements, indicating high, neutral and low scores, respectively. Default=c("#1B9E77", "white", "blue")
<code>url.base</code>	path the location of the output files relative to the gCMAPWeb home directory
<code>up.label</code>	character, legend label for positive scores
<code>down.label</code>	character, legend label for negative scores

Value

None. Pdf and png versions of the overview file are written to file.

Author(s)

Thomas Sandmann

create_profile_ExpressionSet
ExpressionSet creator

Description

This function creates an ExpressionSet object from user-specified identifiers and scores.

Usage

```
create_profile_ExpressionSet(post, conf_data)
```

Arguments

post	list, POST component of the Rook request
conf_data	list, the configuration data as returned by the read_config_file function

Value

ExpressionSet object

Author(s)

Thomas Sandmann

create_query_objects *Query object creator*

Description

This function generates an appropriate R data object from the user query

Usage

```
create_query_objects(post, conf_data)
```

Arguments

post	list, POST component of the Rook request
conf_data	list, the configuration data as returned by the read_config_file function

Value

one of GeneSet, SignedGeneSet or ExpressionSet

Author(s)

Thomas Sandmann

create_SignedGeneSet *SignedGeneSet creator*

Description

This function creates a SignedGeneSet object from user-specified identifiers.

Usage

```
create_SignedGeneSet(post, conf_data)
```

Arguments

post	list, POST component of the Rook request
conf_data	list, the configuration data as returned by the read_config_file function

Value

SignedGeneSet object

Author(s)

Thomas Sandmann

create_tab *Create tab-delimited output*

Description

This function exports results in tab-delimited format

Usage

```
create_tab(df, result.dir, url.base = NULL, file.name)
```

Arguments

df	data.frame
result.dir	character, path to the output directory
url.base	character, the name of the reference database
file.name	character, name of the report file

Value

character string with the html code pointing to the download URL

Author(s)

Thomas Sandmann

create_zip	<i>Export gCMAPWeb report as zip archive</i>
------------	--

Description

This function exports the complete report in a zip file

Usage

```
create_zip(tmp_filename, out.dir)
```

Arguments

tmp_filename	character, name of the per-session result directory to be archived
out.dir	character, output directory to save zip archive into

Value

Nothing, the zip archive is written to disk.

Author(s)

Thomas Sandmann

entrez_GeneSets	<i>GeneSet Entrez mapper</i>
-----------------	------------------------------

Description

This function maps the gene identifiers of a GeneSet object to Entrez identifiers and returns an error if none of them could be found.

Usage

```
entrez_GeneSets(gs)
```

Arguments

gs	GeneSet or SignedGeneSet
----	--------------------------

Details

gCMAPWeb uses this function to ensure that all submitted / retrieved Entrez Ids are valid.

Value

GeneSet or SignedGeneSet

Author(s)

Thomas Sandmann

`exampleCMAP`*Generate example NChannelSets*

Description

This function generates NChannelSets from randomly selected gene identifiers and scores. Scores of the first (simulated) sample will be shifted up by the amount specified by the 'add' parameter for the first 50 genes and down by the same amount for genes 51-100.

Usage

```
exampleCMAP(universe = "org.Hs.eg.db", idType = "ENTREZID", rows = 1000,
            cols = 20, seed = 123, add = 3)
```

Arguments

<code>universe</code>	character, the name of an available annotation package. Default=org.Hs.eg.db
<code>idType</code>	character, a valid identifier type supported by the selected annotation package. Default=ENTREZID
<code>rows</code>	integer, the number of genes to sample. Default=1000
<code>cols</code>	integer, the number of samples to generate. Default=20
<code>seed</code>	numeric, seed for the random number generator. Default=123
<code>add</code>	numeric, score shift for scores in the first sample column. Scores for genes 1-50 will be shifted up, those for genes 51-100 down.

Details

This function is mainly intended for testing purposes.

Value

NChannelSet with simulated scores.

Author(s)

Thomas Sandmann

Examples

```
## generate example NChannelSet with 100 features and 10 columns
## for human EntrezIds
if( require( "org.Hs.eg.db", character.only=TRUE)){
  e <- exampleCMAP( universe="org.Hs.eg.db", rows=100, cols=10)
  dim( e )
  head( featureNames( e ))
  assayDataElementNames( e )
  ## or gene symbols
  s <- exampleCMAP( universe="org.Hs.eg.db", idType="SYMBOL")
  head( featureNames( s ))
}
```

examplePost	<i>Simulate POST request</i>
-------------	------------------------------

Description

This function returns a simulated POST request, similar to those generated from user-input into the web interface.

Usage

```
examplePost(cmap, inputType = "non-directional", species = "human",
array.platform = NULL, idType = "entrez", rows = 1000, add = 3,
seed = 123)
```

Arguments

cmap	NChannelSet object, e.g. generated by the exampleCMAP function
inputType	character, one of directional, non-directional, profile
species	character, should match a species supported by the loaded gCMAPWeb configuration file
array.platform	character, should match an array platform supported by the loaded gCMAPWeb configuration file
idType	character, one of entrez, symbol or probe
rows	integer, number of genes to include in a 'profile' query.
add	numeric, score shift for scores in the first sample column. Scores for genes 1-50 will be shifted up, those for genes 51-100 down. Only used to simulate 'profile' queries.
seed	numeric, seed for the random number generator. Default=123

Details

This function is mainly intended for testing purposes.

Value

list, containing the same elements as a gCMAPWeb POST request

Author(s)

Thomas Sandmann

Examples

```
if( require( "org.Hs.eg.db", character.only=TRUE)){
  ## generate an example NChannelSet for human EntrezIds
  e <- exampleCMAP( universe="org.Hs.eg", rows=1000, cols=10)
  ## generate a matching example Post, simulating a non-directional query
  examplePost( cmap=e )
  ## or a directional query
  examplePost( cmap=e, inputType="directional")
}
```

gCMAPWeb

*Start local gCMAPWeb instance***Description**

This function starts a gCMAPWeb instance on the local machine using R's internal web server.

Usage

```
gCMAPWeb(config.file.path = system.file("config", "config.yml", package =
  "gCMAPWeb"), url.root = system.file("htdocs", package = "gCMAPWeb"))
```

Arguments

config.file.path	character, path to a gCMAPWeb configuration file in yaml format
url.root	character, path to the htdocs folder

Value

an Rhttpd class object

Author(s)

Thomas Sandmann

Examples

```
if (interactive()) {
  ## start a gCMAPWeb instance with the example data and
  ## configuration provided in the package
  gCMAPWeb()
  ## same as above, explicitely specifying the location
  ## of the configuration file
  gCMAPWeb(
    config.file.path = system.file("config", "config.yml",
                                   package = "gCMAPWeb" )
  )
}
```

generate_report

*gCMAPWeb reporting function***Description**

This function receives the results of a gene-set enrichment analysis and the original reference databases to generate plots and html code elements for the final report.

Usage

```
generate_report(cmap.result, reference, reference.name, annotation.db, element,
  query, tmp_filename, title = "", max.results = getOption("max.results",
  default = 50), min.found = getOption("min.found", default = 1),
  max.padj = getOption("max.padj", default = 0.1),
  gene.level.report = getOption("gene.level.report", default = TRUE),
  gene.level.plot = getOption("gene.level.plot", default = TRUE),
  reportDirectory = tempdir(), excluded.cols = getOption("excluded.cols",
  default = c("geneScores", "signed", "pval", "UID", "z.shift", "log_fc.shift",
  "mod_fc.shift")), swap.colnames = getOption("swap.colnames", default =
  list(padj = "FDR", nFound = "Genes")))
```

Arguments

cmap.result	CMAPResults object
reference	eSet object, the full reference database, typically an NChannelSet or CMAPCollection object
reference.name	character, the name of the reference database
annotation.db	character, the name of the annotation package used to lookup gene identifiers
element	character, the assayDataElementName to extract from NChannelSet objects
query	a GeneSet, SignedGeneSet or ExpressionSet object used as query for the gene-set-enrichment analysis giving rise to 'cmap.result'
tmp_filename	character, the name of the temporary output folder
title	character, title of the report
max.results	integer, maximum number of results to return
min.found	integer, minimum number of gene set members that need to overlap with the user query. Gene sets with matching members less than min.found will not be returned as results.
max.padj	numeric, maximum adjusted p-value for a gene set to be considered significantly similar. Results with adjusted p-values larger than max.padj will not be returned.
gene.level.report	logical, create gene-level reports ?
gene.level.plot	logical, create gene-level plots ?
reportDirectory	character, path to temporary output directory for this session. Default=tempdir()
excluded.cols	character vector listing columns in the CMAPResults objects that should NOT be included in the report
swap.colnames	list, containing alternative names for CMAPResults columns.

Details

This function is called by the cmapAnalysis function

Value

a list of character strings, either containing html code snippets or paths to result files. These elements are used to brew the final result html page.

Author(s)

Thomas Sandmann

gene_density_chart_legend

Html code for the legend of pie the pie chart of the gene-level report

Description

Html code for the legend of pie the pie chart of the gene-level report

Usage

`gene_density_chart_legend(text)`

Arguments

text Character, text for the legend of the density plot on gene-level reports for non-directional and directional queries. Can be set as the global variabel "gene.density.legend".

Value

Character string with html code

Author(s)

Thomas Sandmann

gene_density_profile_legend

Html code for the legend of the density charts of the gene-level reports from Profile queries

Description

Html code for the legend of the density charts of the gene-level reports from Profile queries

Usage

`gene_density_profile_legend(text)`

Arguments

text Character, text for the legend of the density plot on gene-level reports for profile queries. Can be set as the global variabel "gene.profile.legend".

Value

Character string with html code

Author(s)

Thomas Sandmann

gene_pie_chart_legend *Html code for the legend of the pie chart of the gene-level report*

Description

Html code for the legend of the pie chart of the gene-level report

Usage

```
gene_pie_chart_legend(text)
```

Arguments

text Character, text for the legend of the pie chart on gene-level reports for non-directional queries. Can be set as the global variabel "gene.pie.legend".

Value

Character string with html code

Author(s)

Thomas Sandmann

hide_species_js *Javascript code to hide html elements*

Description

Javascript code to hide html elements for all but the first species listed in the configuration file

Usage

```
hide_species_js(conf_data)
```

Arguments

conf_data list, the configuration data as returned by the read_config_file function

Value

javascript call as character string

Author(s)

Thomas Sandmann

`html_body_last` *Html footer*

Description

Closing html code, including references to jquery, bootstrap and dataTables javascript libraries

Usage

```
html_body_last(url.base = NULL)
```

Arguments

`url.base` character, path to htdocs directory (optional)

Value

Html code as character string

Author(s)

Thomas Sandmann

`html_header` *Html header and bootstrap css stylesheet*

Description

Html header including link to the bootstrap cerulean css stylesheet

Usage

```
html_header(url.base = NULL)
```

Arguments

`url.base` character, path to htdocs directory (optional)

Value

Html code as a character string

Author(s)

Thomas Sandmann

html_table	<i>Format data.frame as html table</i>
------------	--

Description

Function to format a data.frame as an html table

Usage

```
html_table(df, table_id)
```

Arguments

df	A data frame
table_id	Character, specifying the id tag for the html table

Details

This function uses the thead and tbody tags required for the dataTables javascript to create interactive html tables.

Value

Html code as a character string.

Author(s)

Thomas Sandmann

identifier_radio_html	<i>Html code for identifier type radio button selectors</i>
-----------------------	---

Description

This function generates the html code required to display radio buttons for the supported identifier types specified in the gCMAPWeb configuration file

Usage

```
identifier_radio_html(conf_data, single = FALSE)
```

Arguments

conf_data	list, the configuration data as returned by the read_config_file function
single,	logical, use singular nouns for radiobutton labels ?

Value

Html code as character string.

Author(s)

Thomas Sandmann

index_message	<i>Html code for text of the additional message on the index page (warning box)</i>
---------------	---

Description

Html code for text of the additional message on the index page (warning box)

Usage

```
index_message(text)
```

Arguments

text Character, text for the additional message on the index page (warning box)

Value

text Character string with html code

Author(s)

Thomas Sandmann

index_quote	<i>Html code for quote on the index page</i>
-------------	--

Description

Html code for quote on the index page

Usage

```
index_quote(text)
```

Arguments

text, Character, the content of the blockquote field on the index.rhtml page. Can be set as the global variabel "index.quote".

Value

Character string with html code

Author(s)

Thomas Sandmann

<code>index_subtitle</code>	<i>Html code for subtitle of the hero unit of the index.rhtml page</i>
-----------------------------	--

Description

Html code for subtitle of the hero unit of the index.rhtml page

Usage

```
index_subtitle(text)
```

Arguments

<code>text</code>	Character, subtitle of the hero unit on the index.rhtml page. Can be set as the global variabel "index.sub".
-------------------	--

Value

Character string with html code

Author(s)

Thomas Sandmann

<code>index_text</code>	<i>Html code for text of the hero unit of the index.rhtml page</i>
-------------------------	--

Description

Html code for text of the hero unit of the index.rhtml page

Usage

```
index_text(text)
```

Arguments

<code>text</code>	Character, text of the hero unit on the index.rhtml page. Can be set as the global variabel "index.text".
-------------------	---

Value

Character string with html code

Author(s)

Thomas Sandmann

<code>index_title</code>	<i>Html code for title of the hero unit of the index.rhtml page</i>
--------------------------	---

Description

Html code for title of the hero unit of the index.rhtml page

Usage

```
index_title(text = getOption("index.main", default = "gConnectivity Map"))
```

Arguments

<code>text</code>	Character, main title of the hero unit on the index.rhtml page. Can be set as the global variabel "index.main".
-------------------	---

Value

Character string with html code

Author(s)

Thomas Sandmann

<code>inputType_buttons</code>	<i>Html code for the query type selection buttons on the gCMAPWeb index page</i>
--------------------------------	--

Description

This function generates the html code required to generate the query type selection buttons on the main index.rhtml page

Usage

```
inputType_buttons(conf_data)
```

Arguments

<code>conf_data</code>	list, the configuration data as returned by the <code>read_config_file</code> function
------------------------	--

Value

Html code as character string.

Author(s)

Thomas Sandmann

load_cmmaps

*Loading reference datasets***Description**

This function connects to / loads all reference datasets and returns them in a list

Usage

```
load_cmmaps(conf_data, min.set.size = getOption("min.set.size", default = 5))
```

Arguments

conf_data	list, the configuration data as returned by the read_config_file function
min.set.size	integer, gene sets with less than min.set.size members will be dropped from CMAPCollections

Value

list of eSet objects

Author(s)

Thomas Sandmann

Examples

```
library(yaml)
## read the example configuration file
conf_data <- yaml.load_file( system.file("config", "config.yml",
                                         package="gCMAPWeb") )
## load the example reference NChannelSet objects specified in
## the configuration file
ref.cmmaps <- load_cmmaps( conf_data )
class( ref.cmmaps )
names( ref.cmmaps )
ref.cmmaps[[1]]
```

parse_file_input

*Uploaded file parser***Description**

This functions parses the user-uploaded files

Usage

```
parse_file_input(temp.file, regexp = "[+,;\t ]+", n.score.col = 1)
```

Arguments

<code>temp.file</code>	character, path to temporary file (usually part of the POST request)
<code>regexp</code>	character, a regular expression matching all separators used to separate gene identifiers
<code>n.score.col</code>	integer, for profile uploads <code>n.score.col</code> specifies the number of data columns (usually 1)

Value

list of character vectors, one element for each row of `temp.file`

Author(s)

Thomas Sandmann

<code>parse_request</code>	<i>POST request parser</i>
----------------------------	----------------------------

Description

This function parses the a POST request and decodes the user-provided information

Usage

```
parse_request(request, conf_data,
  cmaps.concatenate_by = getOption("cmaps.concatenate_by", default = ","))
```

Arguments

<code>request</code>	Request object as defined by the Rook package
<code>conf_data</code>	list, the configuration data as returned by the <code>read_config_file</code> function
<code>cmaps.concatenate_by</code>	character, specifying the separator (e.g. semicolon, comma) used to concatenate multiple requested reference databases

Value

list, all elements from POST request, decoded if necessary

Author(s)

Thomas Sandmann

parse_textarea_input *Textarea parser*

Description

This function parses user-specified data pasted into the textarea fields of the input form

Usage

```
parse_textarea_input(textarea.input, regexp = "[+,;\\t ]+")
```

Arguments

textarea.input character, the user-input extracted from the POST request

regexp character, a regular expression matching all separators used to separate gene identifiers

Value

list of character vectors, one element for each original row of textarea.input

Author(s)

Thomas Sandmann

platform_radio_html *Html code for platform radio button selectors*

Description

This function generates the html code required to display radio buttons for all array platforms specified in the gCMAPWeb configuration file

Usage

```
platform_radio_html(conf_data)
```

Arguments

conf_data list, the configuration data as returned by the read_config_file function

Value

Html code as character string.

Author(s)

Thomas Sandmann

`process_score_input` *Score parser*

Description

This function reformats score input uploaded or pasted by the user

Usage

```
process_score_input(query_data, n.score.col = 1)
```

Arguments

<code>query_data</code>	list of row-vectors, as generated by <code>parse_file_input</code> function
<code>n.score.col</code>	integer, specifies the number of data columns (usually 1)

Value

matrix of expression scores

Author(s)

Thomas Sandmann

`profile_input_example` *Html code for generating an example query button on the profile submission page*

Description

Content can be provided / modified through the global parameters `profile.input.example.popover` (text for popover help) `profile.input.example` (gene identifiers and scores)

Usage

```
profile_input_example()
```

Value

Character string with html code

Author(s)

Thomas Sandmann

read_config_file *Parser for gCMAPWeb configuration file*

Description

This function parses the gCMAPWeb configuration file (in yaml format) and returns the information as a list

Usage

```
read_config_file(config.file.path = system.file("config", "config.yml",
  package = "gCMAPWeb"))
```

Arguments

config.file.path
character, full path to gCMAPWeb configuration file

Value

a nested list

Author(s)

Thomas Sandmann

reference_radio_html *Html code for reference cmap radio button selectors*

Description

This function generates the html code required to display radio buttons for the reference datasets specified in the gCMAPWeb configuration file

Usage

```
reference_radio_html(conf_data, reference.cmmaps)
```

Arguments

conf_data list, the configuration data as returned by the read_config_file function
reference.cmmaps
list containing all reference cmmaps as eSet-like objects (e.g. NChannelSet, CMAP-Collection)

Value

Html code as character string.

Author(s)

Thomas Sandmann

`retrieve_annotation` *Gene symbol and name lookup*

Description

This function retrieves the official gene symbol and name for Entrez identifiers

Usage

```
retrieve_annotation(entrez, annotation.db)
```

Arguments

<code>entrez</code>	character, Entrez identifier to look up
<code>annotation.db</code>	character, name of the annotation package to use for lookup

Value

`data.frame` with `entrez`, `symbol` and `name` columns

Author(s)

Thomas Sandmann

`signed_input_example` *Html code for generating and example query button on the signed gene set submission page*

Description

Content can be provided / modified through the global parameters `signed.input.example.popover` (text for popover help) `signed.input.example.up` (up-regulated gene identifiers) `signed.input.example.down` (down-regulated gene identifiers)

Usage

```
signed_input_example()
```

Value

Character string with html code

Author(s)

Thomas Sandmann

single_input_example *Html code for generating and example query button on the single gene query submission page*

Description

Content can be provided / modified through the global parameters single.gene.example.popover (text for popover help) single.gene.example (gene identifiers)

Usage

```
single_input_example()
```

Value

Character string with html code

Author(s)

Thomas Sandmann

species_radio_html *Html code for species radio button selectors*

Description

This function generates the html code required to display radio buttons for all species specified in the gCMAPWeb configuration file

Usage

```
species_radio_html(conf_data)
```

Arguments

conf_data list, the configuration data as returned by the read_config_file function

Value

Html code as character string.

Author(s)

Thomas Sandmann

stripPrefix	<i>Strip prefix from gene identifier names</i>
-------------	--

Description

This function strips a prefix from gene identifier strings

Usage

```
stripPrefix(ids, prefix = c("^GeneID:", "^GeneID"))
```

Arguments

ids	character, vector of gene identifiers
prefix	character, one or more strings to be removed

Value

character, gene identifiers without prefix

Author(s)

Thomas Sandmann

unsigned_input_example	<i>Html code for generating and example query button on the unsigned gene set submission page</i>
------------------------	---

Description

Content can be provided / modified through the global parameters `unsigned.input.example.popover` (text for popover help) `unsigned.input.example` (gene identifiers)

Usage

```
unsigned_input_example()
```

Value

Character string with html code

Author(s)

Thomas Sandmann

validate_config_file *Validation of gCMAPWeb configuration file*

Description

This function validates the content of the gCMAPWeb configuration file.

Usage

```
validate_config_file(config.file.path)
```

Arguments

`config.file.path`
full path to gCMAPWeb configuration file (in yaml format)

Details

Verifies that 1. all required fields are present 2. at least one species has been defined 3. all supported annotation packages are available 4. all cmaps have unique labels 5. each supported species has at least one associated reference dataset

Value

Information from a valid configuration file is returned as a nested list. Invalid entries in the config file will cause an error.

Author(s)

Thomas Sandmann

Examples

```
## read the example configuration yaml file without validation
library(yaml)
conf1 <- yaml.load_file( system.file("config", "config.yml",
                                         package="gCMAPWeb") )
## read the example configuration file and validate that
## all required information is provided and valid
conf2 <- validate_config_file( system.file("config", "config.yml",
                                         package="gCMAPWeb") )
identical( conf1, conf2)
```

`validate_request` *Html request validation*

Description

This function extracts the content from a POST request and validates its content

Usage

```
validate_request(req, conf_data,  
                 cmaps.concatenate_by = getOption("cmaps.concatenate_by", default = ",", ))
```

Arguments

<code>req</code>	Request object as defined by the Rook package
<code>conf_data</code>	list, the configuration data as returned by the <code>read_config_file</code> function
<code>cmaps.concatenate_by</code>	character, specifying the separator (e.g. semicolon, comma) used to concatenate multiple requested reference databases

Value

Logical, TRUE if all required fields could be validated, FALSE otherwise

Author(s)

Thomas Sandmann

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