Package 'hipathia'

July 26, 2025

Title HiPathia: High-throughput Pathway Analysis

Version 3.9.0

Description Hipathia is a method for the computation of signal transduction along signaling pathways from transcriptomic data. The method is based on an iterative algorithm which is able to compute the signal intensity passing through the nodes of a network by taking into account the level of expression of each gene and the intensity of the signal arriving to it. It also provides a new approach to functional analysis allowing to compute the signal arriving to the

functions annotated to each pathway.

Depends R (>= 4.1), igraph (>= 1.0.1), AnnotationHub(>= 2.6.5), MultiAssayExperiment(>= 1.4.9), SummarizedExperiment(>= 1.8.1)

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Encoding UTF-8

Imports coin, stats, limma, grDevices, utils, graphics, preprocessCore, servr, DelayedArray, matrixStats, methods, S4Vectors, ggplot2, ggpubr, dplyr, tibble, visNetwork, reshape2, MetBrewer

RoxygenNote 7.2.2

Suggests BiocStyle, knitr, rmarkdown, testthat

VignetteBuilder knitr

biocViews Pathways, GraphAndNetwork, GeneExpression, GeneSignaling, GO

LazyData true

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

git_branch devel

git_last_commit 29effee

git_last_commit_date 2025-04-15

Repository Bioconductor 3.22

Date/Publication 2025-07-25

Author Marta R. Hidalgo [aut, cre], José Carbonell-Caballero [ctb], Francisco Salavert [ctb], Alicia Amadoz [ctb], Çankut Cubuk [ctb], Joaquin Dopazo [ctb]

Maintainer Marta R. Hidalgo <marta.hidalgo@outlook.es>

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annotate_paths

Annotates functions to pathways

Description

Annotates functions from a database to each pathway

Usage

annotate_paths(metaginfo, dbannot)

Arguments

| metaginfo | Pathways object |
|-----------|---|
| dbannot | Either a string indicating which precomputed annotation to use ("uniprot" for Uniprot Keywords or "GO" for Gene Ontology terms), or a dataframe with the annotation of the genes to the functions. First column are gene symbols, second column the functions. |

Value

Object of annotations from pathways to functions

#@examples #pathways <- load_pathways(species = "hsa", pathways_list = c("hsa03320", #"hsa04012")) #annotate_paths(pathways, "GO")

#@export

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brca

Description

A dataset containing a matrix with the Gene expression of 40 samples from the BRCA-US project from The Cancer Genome Atlas (TCGA), and their experimental design, containing 20 "Tumor" samples 20 "Normal" samples.

Usage

data(brca)

Format

SummarizedExperiment. The assay is a matrix with 40 columns and 18638 rows. Row names are Entrez IDs and column names are the TCGA identifyers of the samples. The colData() is a data.frame with 1 column and 40 rows, including the experimental design of the 40 samples from the BRCA-US project from TCGA. Field group is the type of sample, either "Tumor" or "Normal".

Details

The gene expression matrix includes 40 samples. The data has been log-transformed and normalized with TMM.

Value

SummarizedExperiment including a matrix with 40 columns and 18638 rows. Row names are Entrez IDs and column names are the TCGA identifyers of the samples.

Source

https://cancergenome.nih.gov/

brca_data

BRCA gene expression dataset

Description

Gene expression of 40 samples from the BRCA-US project from The Cancer Genome Atlas (TCGA).

Usage

```
data(brca_data)
```

Format

Matrix with 40 columns and 18638 rows. Row names are Entrez IDs and column names are the TCGA identifyers of the samples.

brca_design

Details

Gene expression matrix with 40 samples taken from the BRCA-US project from The Cancer Genome Atlas (TCGA). The data has been log-transformed and normalized with TMM.

Value

Matrix with 40 columns and 18638 rows. Row names are Entrez IDs and column names are the TCGA identifyers of the samples.

Source

https://cancergenome.nih.gov/

brca_design

BRCA experimental design

Description

Experimental design of the gene expression matrix brca_data with 40 samples taken from the BRCA-US project from The Cancer Genome Atlas (TCGA). 20 samples are "Tumor" samples and 20 samples are "Normal" samples.

Usage

data(brca_design)

Format

Dataframe with 1 column and 40 rows, including the experimental design of the 40 samples from the BRCA-US project from TCGA. Field group is the type of sample, either "Tumor" or "Normal".

Value

Dataframe with 1 column and 40 rows, including the experimental design of the 40 samples from the BRCA-US project from TCGA. Field group is the type of sample, either "Tumor" or "Normal".

Source

https://cancergenome.nih.gov/

comp

Description

Comparison object returned by hipathia::do_wilcoxon function, after calling comp <- do_wilcoxon(path_vals, sample_group, g1 = "Tumor", g2 = "Normal") path_names <- get_path_names(pathways, rownames(comp)) comp <- cbind(path_names, comp)

Usage

data(comp)

Format

Table with 1868 rows and 5 columns

Value

Pathway comparison result

create_report Create visualization HTML

Description

Saves the results of a Wilcoxon comparison for the Hipathia pathway values into a folder, and creates a HTML from which to visualize the results on top of the pathways. The results are stored into the specified folder. If this folder does not exist, it will be created. The parent folder must exist.

Usage

```
create_report(
   comp,
   metaginfo,
   output_folder = NULL,
   path = NULL,
   node_colors = NULL,
   group_by = "pathway",
   conf = 0.05,
   verbose = FALSE
)
```

DAcomp

Arguments

| comp | Comparison object as given by the do_wilcoxon function |
|---------------|---|
| metaginfo | Pathways object as returned by the load_pathways function |
| output_folder | Name of the folder in which the report will be stored. |
| path | Absolute path to the parent directory in which 'output_folder' will be saved. If it is not provided, it will be created in a temp folder. |
| node_colors | List of colors with which to paint the nodes of the pathways, as returned by the node_color_per_de function. Default is white. |
| group_by | How to group the subpathways to be visualized. By default they are grouped by the pathway to which they belong. Available groupings include "uniprot", to group subpathways by their annotated Uniprot functions, "GO", to group sub- pathways by their annotated GO terms, and "genes", to group subpathways by the genes they include. Default is set to "pathway". |
| conf | Level of significance. By default 0.05. |
| verbose | Boolean, whether to show details about the results of the execution |

Value

Saves the results and creates a report to visualize them through a server in the specified output_folder. Returns the folder where the report has been stored.

Examples

```
data(comp)
pathways <- load_pathways(species = "hsa", pathways_list = c("hsa03320",
    "hsa04012"))
report <- create_report(comp, pathways, "save_results")
## Not run:
data(results)
data(brca)
sample_group <- colData(brca)[,1]
colors_de <- node_color_per_de(results, pathways,
sample_group, "Tumor", "Normal")
report_colors <- create_report(comp, pathways, "save_results",
node_colors = colors_de)
## End(Not run)</pre>
```

| DAcomp | Compares the gene expression, pathway activation level and the func- |
|--------|--|
| | tion activation level of the |

Description

Compares the gene expression, pathway activation level and the function activation level of the

Usage

```
DAcomp(
  hidata,
  groups,
  expdes,
  g2 = NULL,
  path.method = "wilcoxon",
  node.method = "limma",
  fun.method = "wilcoxon",
  order = FALSE,
  paired = FALSE,
  adjust = TRUE,
  conf.level = 0.05,
  sel_assay = 1
)
```

Arguments

| hidata | $Either \ a \ Summarized Experiment \ object \ or \ a \ matrix, returned \ by \ function \ hipathia.$ |
|-------------|--|
| groups | Either a character indicating the name of the column in colData including the classes to compare, or a character vector with the class to which each sample belongs. Samples must be ordered as in hidata. |
| expdes | String, either an equation expression to pas to limma, or the label of the first group to be compared |
| g2 | String, label of the second group to be compared, if not specified in expdes. |
| path.method | String, method to be used when comparing pathways. Options include wilcoxon (default, performs a Wilcoxon test comparing conditions expdes and g2 - in this case, mandatory parameter) and limma (performs a limma DE analysis using functions lmFit, contrasts.fit and eBayes using the formula in expdes or comparing conditions expdes and g2. |
| node.method | String, method to be used when comparing nodes. Options include wilcoxon (performs a Wilcoxon test comparing conditions expdes and g2 - in this case, mandatory parameter) and limma (default, performs a limma DE analysis using functions lmFit, contrasts.fit and eBayes using the formula in expdes or comparing conditions expdes and g2. |
| fun.method | String, method to be used when comparing functions. Options include wilcoxon (default, performs a Wilcoxon test comparing conditions expdes and g2 - in this case, mandatory parameter) and limma (performs a limma DE analysis using functions lmFit, contrasts.fit and eBayes using the formula in expdes or comparing conditions expdes and g2. |
| order | Boolean, whether to order the results table by the FDRp.value column. Default is FALSE. |
| paired | Boolean, whether the samples to be compared are paired. If TRUE, function wilcoxsign_test from package coin is used. If FALSE, function wilcox.test from package stats is used. |
| adjust | Boolean, whether to adjust the p.value with Benjamini-Hochberg FDR method. Default is TRUE. |
| conf.level | Numeric, cut off for significance. Default is 0.05. |
| sel_assay | Character or integer, indicating the assay to be normalized in the Summarized- Experiment. Default is 1. |

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DAdata

Value

List including comparison results for nodes, pathways and functions, if present.

Examples

```
data(hidata)
comp <- DAcomp(hidata, groups = "group", expdes = "Tumor", g2 = "Normal")</pre>
```

| DAdata |
|--------|
|--------|

Wilcoxon and limma comparison object for nodes, pathways and functional annotations

Description

Comparison object returned by hipathia::DAcomp function, after calling DAdata <- DAcomp(hidata, "group", g1 = "Tumor", g2 = "Normal")

Usage

data(DAdata)

Format

List object with 4 entries: Nodes includes a matrix with 6826 rows and 8 columns Paths includes a matrix with 1876 rows and 13 columns Uni.terms includes a matrix with 142 rows and 6 columns GO.terms includes a matrix with 1654 rows and 6 columns

Value

List of tibbles with the comparison results

| DAoverview | Table and plot of total number of altered and not altered nodes, paths |
|------------|--|
| | and functions (Uniprot keywords and/or GO terms, if present). |

Description

Table and plot of total number of altered and not altered nodes, paths and functions (Uniprot keywords and/or GO terms, if present).

Usage

```
DAoverview(DAdata, conf.level = 0.05, adjust = TRUE, colors = "hiro")
```

Arguments

| DAdata | List of comparison results, returned by function DAcomp. |
|------------|--|
| conf.level | Numeric, cut off for significance. Default is 0.05. |
| adjust | Boolean, whether to adjust the p.value with Benjamini-Hochberg FDR method. Default is TRUE. |
| colors | String with the color scheme or vector of colors to be used. See define_colors for available options. Default is "hiro". |

Value

Plot and tibble including the number of total, altered, UP- and DOWN-regulated features for nodes, paths and functions if present.

Examples

data(DAdata) DAoverview(DAdata)

DAreport

Create visualization HTML

Description

Saves the results of a DAdata comparison for the Hipathia pathway values into a folder, and creates a HTML from which to visualize the results on top of the pathways. The results are stored into the specified folder. If this folder does not exist, it will be created. The parent folder must exist.

Usage

```
DAreport(
   DAdata,
   pathways,
   conf.level = 0.05,
   adjust = TRUE,
   group_by = "pathway",
   colors = "classic",
   output_folder = NULL,
   path = NULL,
   verbose = TRUE
)
```

Arguments

| DAdata | List of comparison results, returned by function DAcomp. |
|------------|---|
| pathways | Pathways object as returned by the load_pathways function |
| conf.level | Level of significance. By default 0.05. |
| adjust | Boolean, whether to adjust the p.value with Benjamini-Hochberg FDR method. Default is TRUE. |

DAsummary

| group_by | How to group the subpathways to be visualized. By default they are grouped by the pathway to which they belong. Available groupings include "uniprot", to group subpathways by their annotated Uniprot functions, "GO", to group sub- pathways by their annotated GO terms, and "genes", to group subpathways by the genes they include. Default is set to "pathway". |
|---------------|---|
| colors | String with the color scheme or vector of colors to be used. See define_colors for available options. Default is "hiro". |
| output_folder | Name of the folder in which the report will be stored. |
| path | Absolute path to the parent directory in which 'output_folder' will be saved. If it is not provided, it will be created in a temp folder. |
| verbose | Boolean, whether to show details about the results of the execution |

Value

Saves the results and creates a report to visualize them through a server in the specified output_folder. Returns the folder where the report has been stored.

Examples

```
data(DAdata)
data(pathways)
DAreport(DAdata, pathways)
```

Lists and plots the top n altered pathways, taking into account the number of altered.

Description

Lists and plots the top n altered pathways, taking into account the number of altered .

Usage

```
DAsummary(
 DAdata,
 n = 10,
 conf.level = 0.05,
 adjust = TRUE,
 ratio = FALSE,
 colors = "hiro",
 order.by = "number"
```

```
)
```

Arguments

| DAdata | List of comparison results, returned by function DAcomp. |
|------------|--|
| n | Number of top features to show. |
| conf.level | Numeric, cut off for significance. Default is 0.05. |

| adjust | Boolean, whether to adjust the p.value with Benjamini-Hochberg FDR method. Default is TRUE. |
|----------|--|
| ratio | Boolean, whether to plot the ratio of significant paths with respect to the total paths in the pathway. Default is FALSE. |
| colors | String with the color scheme or vector of colors to be used. See define_colors for available options. Default is "hiro". |
| order.by | String, how to order table of results. Available options include ratio (default, uses the ratio of significant paths with respect to the total paths in the pathway) and number (uses the number of significant paths in the pathway). |

Value

Plot and tibble including top n altered pathways.

Examples

data(DAdata) DAsummary(DAdata)

| DAtop | Lists and plots the top n altered nodes, paths and functions (Uniprot |
|-------|---|
| | keywords and/or GO terms, if present). |

Description

Lists and plots the top n altered nodes, paths and functions (Uniprot keywords and/or GO terms, if present).

Usage

DAtop(DAdata, n = 10, conf.level = 0.05, adjust = TRUE, colors = "hiro")

Arguments

| DAdata | List of comparison results, returned by function DAcomp. |
|------------|--|
| n | Number of top features to show. |
| conf.level | Numeric, cut off for significance. Default is 0.05. |
| adjust | Boolean, whether to adjust the p.value with Benjamini-Hochberg FDR method. Default is TRUE. |
| colors | String with the color scheme or vector of colors to be used. See define_colors for available options. Default is "hiro". |

Value

Plot and list of tables including top n altered features for nodes, paths and functions if present.

Examples

data(DAdata) DAtop(DAdata) define_colors

Description

Color palettes to be used in plots.

Usage

```
define_colors(colors, no.col = NULL)
```

Arguments

| colors | String with the color scheme or vector of colors to be used. Available predefined options include: hipathia, classic, soft, okee, hiro, new, vg, orchid. |
|--------|--|
| no.col | String with the color given to non-significant nodes, if not given in parameter colors. |

Value

Plot and list of tables including top n altered features for nodes, paths and functions if present.

Examples

define_colors("hiro")

do_pca

Performs a Principal Components Analysis

Description

Performs a Principal Components Analysis

Usage

```
do_pca(data, sel_assay = 1, cor = FALSE)
```

Arguments

| data | SummarizedExperiment or matrix of values to be analyzed. Samples must be represented in the columns. |
|-----------|--|
| sel_assay | Character or integer, indicating the assay to be normalized in the Summarized-Experiment. Default is 1. |
| cor | A logical value indicating whether the calculation should use the correlation matrix or the covariance matrix. (The correlation matrix can only be used if there are no constant variables.) |

Value

do_pca returns a list with class princomp.

Examples

```
data(path_vals)
pca_model <- do_pca(path_vals[seq_len(ncol(path_vals)),])</pre>
```

do_wilcoxon Apply Wilcoxon test

Description

Performs a Wilcoxon test for the values in sel_vals comparing conditions g1 and g2

Usage

```
do_wilcoxon(
   data,
   group,
   g1,
   g2,
   paired = FALSE,
   adjust = TRUE,
   sel_assay = 1,
   order = FALSE
)
```

Arguments

| data | Either a SummarizedExperiment object or a matrix, containing the values. Columns represent samples. |
|-----------|---|
| group | Either a character indicating the name of the column in colData including the classes to compare, or a character vector with the class to which each sample belongs. Samples must be ordered as in data |
| g1 | String, label of the first group to be compared |
| g2 | String, label of the second group to be compared |
| paired | Boolean, whether the samples to be compared are paired. If TRUE, function wilcoxsign_test from package coin is used. If FALSE, function wilcox.test from package stats is used. |
| adjust | Boolean, whether to adjust the p.value with Benjamini-Hochberg FDR method |
| sel_assay | Character or integer, indicating the assay to be normalized in the Summarized- Experiment. Default is 1. |
| order | Boolean, whether to order the results table by the FDRp.value column. Default is FALSE. |

Value

Dataframe with the result of the comparison

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exp_data

Examples

```
data(path_vals)
data(brca_design)
sample_group <- brca_design[colnames(path_vals),"group"]
comp <- do_wilcoxon(path_vals, sample_group, g1 = "Tumor", g2 = "Normal")</pre>
```

exp_data

Normalized BRCA gene expression dataset

Description

Experimental design matrix once expression matrix brca_data has been translated to Entrez geens with translate_matrix and normalized using normalize_data.

Usage

data(exp_data)

Format

Matrix with 40 columns and 3184 rows. Row names are Entrez IDs and column names are the TCGA identifyers of the samples.

Details

To create the data, the following functions have been called: trans_data <- translate_matrix(brca_data, "hsa") exp_data <- normalize_data(trans_data)

Value

Matrix with 40 columns and 3184 rows. Row names are Entrez IDs and column names are the TCGA identifyers of the samples.

get_go_names Tranlates GO IDs to GO names

Description

Translates the GO IDs to readable and comprensible names.

Usage

```
get_go_names(names, species, maxchar = NULL, disambiguate = FALSE)
```

Arguments

| names | Character vector with the GO IDs to be translated. |
|--------------|---|
| species | Species of the samples. |
| maxchar | Integer, describes the number of maximum characters to be shown. By default no filter is applied. |
| disambiguate | Boolean, whether to return unique strings by disambiguating with numbers. |

Value

A character vector including the readable names of the GO IDs, in the same order as provided.

Examples

```
data(go_vals)
get_go_names(rownames(go_vals), "hsa")
```

get_highest_sig_ancestor

Get highest common GO ancestor of GO annotations

Description

Get highest common GO ancestor of GO annotations

Usage

```
get_highest_sig_ancestor(
  go_terms,
  go_comp,
  metaginfo,
  unique = TRUE,
  pval = 0.05
)
```

Arguments

| go_terms | GO terms for which the highest common ancestors are to be looked for. |
|-----------|---|
| go_comp | Wilcoxon comparison of the matrix of GO values as returned by do_wilcoxon. |
| metaginfo | Pathways object |
| unique | Boolean, whether to return only one highest significant GO ancestor or all of them. By default, TRUE. |
| pval | P-value cut-off. Default values is set to 0.05. |

Value

highest common ancestors

#@export

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get_nodes_data

Description

This function returns the object with the levels of activation of each node for each sample. Rows represent the nodes and columns represent the samples. Each cell is the value of activation of a node in a sample.

Rownames are the IDs of the nodes In order to transform IDs into readable names, use get_node_names.

Effector subpathways are subgraphs of a pathway including all the paths leading to an effector protein. Effector proteins are defined as final nodes in the graph. Each effector protein (final node) in a pathway defines its own effector subpathway as the nodes and edges in a path leading to it.

Decomposed subpathways are subgraphs of a pathway including all the paths starting in a receptor protein and ending in an effector protein. Receptor proteins are defined as initial nodes and effector proteins are defined as final nodes in the graph. Each effector subpathway can be decomposed in as many decomposed subpathways as initial nodes it includes.

Usage

get_nodes_data(results, matrix = FALSE)

Arguments

| results | Results object as returned by hipathia. |
|---------|--|
| matrix | Boolean, if TRUE the function returns a matrix object, if FALSE (as default) |
| | returns a SummarizedExperiment object. |

Value

Object, either a SummarizedExperiment or a matrix, with the levels of activation of each decomposed subpathway for each sample.

Examples

```
data(results)
path_vals <- get_paths_data(results)</pre>
```

get_node_names Tranlates node IDs to node names

Description

Translates the node IDs to readable and comprensible names.

The names of the nodes are encoded as "pathway: name", where "pathway" is the pathway to which the node belongs and "node" is the name of the node. Nodes may include more genes than the one depicted in the name.

Usage

```
get_node_names(metaginfo, names, maxchar = NULL)
```

Arguments

| metaginfo | Pathways object |
|-----------|---|
| names | Character vector with the subpathway IDs to be translated |
| maxchar | Integer, describes the number of maximum characters to be shown. By default no filter is applied. |

Value

A character vector including the readable names of the subpathways IDs, in the same order as provided.

Examples

```
data(results)
pathways_list <- c("hsa03320", "hsa04012")
pathways <- load_pathways(species = "hsa", pathways_list)
node_vals <- get_nodes_data(results)
translated_names <- get_node_names(pathways, rownames(node_vals))</pre>
```

get_paths_data Gets the object of subpathway activation values

Description

This function returns the object with the levels of activation of each subpathway for each sample. Rows represent the subpathways and columns represent the samples. Each cell is the value of activation of a subpathway in a sample.

Rownames are the IDs of the subpathways. In order to transform IDs into readable names, use get_path_names.

Effector subpathways are subgraphs of a pathway including all the paths leading to an effector protein. Effector proteins are defined as final nodes in the graph. Each effector protein (final node) in a pathway defines its own effector subpathway as the nodes and edges in a path leading to it.

Decomposed subpathways are subgraphs of a pathway including all the paths starting in a receptor protein and ending in an effector protein. Receptor proteins are defined as initial nodes and effector proteins are defined as final nodes in the graph. Each effector subpathway can be decomposed in as many decomposed subpathways as initial nodes it includes.

Usage

```
get_paths_data(results, matrix = FALSE)
```

Arguments

| results | Results object as returned by hipathia. |
|---------|--|
| matrix | Boolean, if TRUE the function returns a matrix object, if FALSE (as default) |
| | returns a SummarizedExperiment object. |

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get_pathways_annotations

Value

Object, either a SummarizedExperiment or a matrix, with the levels of activation of each decomposed subpathway for each sample.

Examples

```
data(results)
path_vals <- get_paths_data(results)</pre>
```

get_pathways_annotations

Get Pathways functional annotations

Description

Get functional annotation of the pathways, either for a particular annotation or a stored one.

Usage

get_pathways_annotations(pathway_names, metaginfo, dbannot, collapse = FALSE)

Arguments

| pathway_names | Character vector of the names of the pathways |
|---------------|---|
| metaginfo | Pathways object |
| dbannot | Either a string indicating which precomputed annotation to use ("uniprot" for Uniprot Keywords or "GO" for Gene Ontology terms), or a dataframe with the annotation of the genes to the functions. First column are gene symbols, second column the functions. |
| collapse | Boolean, whether to collapse all functions of the same path in a single character string. |

Value

2-columns matrix with the annotations of each pathway ID in the annotation dbannot.

Examples

```
pathways <- load_pathways(species = "hsa", pathways_list = c("hsa03320",
    "hsa04012"))
pathway_names <- c("P-hsa03320-37", "P-hsa03320-61", "P-hsa03320-46",
    "P-hsa03320-57", "P-hsa03320-64", "P-hsa03320-47", "P-hsa03320-65")
## Not run: get_pathways_annotations(pathway_names, pathways, "GO")
get_pathways_annotations(pathway_names, pathways, "uniprot")
```

get_pathways_list Lists the IDs of the pathways in a pathways object

Description

Lists the IDs of the pathways included in the pathways object metaginfo

Usage

```
get_pathways_list(metaginfo)
```

Arguments

metaginfo Pathways object

Value

List of the pathway IDs included in the pathways object

Examples

```
pathways <- load_pathways(species = "hsa", pathways_list = c("hsa03320",
"hsa04012"))
pathways_list <- get_pathways_list(pathways)</pre>
```

get_pathways_summary Compute pathway summary

Description

Computes a summary of the results, summarizing the number and proportion of up- and down-regulated subpathways in each pathway.

Usage

```
get_pathways_summary(comp, metaginfo, conf = 0.05)
```

Arguments

| comp | Comparison data frame as returned by the do_wilcoxon function. |
|-----------|--|
| metaginfo | Pathways object |
| conf | Level of significance of the comparison for the adjusted p-value. Default is 0.05. |

Value

Table with the summarized information for each of the pathways. Rows are the analized pathways. Columns are: * num_total_paths Number of total subpathways in which each pathway is decomposed. * num_significant_paths Number of significant subpathways in the provided comparison. * percent_significant_paths Percentage of significant subpathways from the total number of subpathways in a pathway. * num_up_paths Number of significant up-regulated subpathways in the provided comparison. * percent_up_paths Percentage of significant up-regulated subpathways from the total number of subpathways in a pathway. * num_down_paths Number of significant down-regulated subpathways in the provided comparison. * percent_down_paths Percentage of significant down-regulated subpathways from the total number of subpathways in a pathway.

Examples

```
data(comp)
pathways <- load_pathways(species = "hsa", pathways_list = c("hsa03320",
    "hsa04012"))
get_pathways_summary(comp, pathways)</pre>
```

get_pathway_functions Returns functions related to a pathway

Description

Returns functions related to a pathway

Usage

```
get_pathway_functions(
   pathigraph,
   dbannot,
   entrez2hgnc,
   use_last_nodes = TRUE,
   unique = TRUE
)
```

Arguments

| pathigraph | Pathway object |
|----------------|--|
| dbannot | Dataframe with the annotation of the genes to the functions. First column are gene symbols, second column the functions. |
| entrez2hgnc | Relation between Entrez and HGNC genes. |
| use_last_nodes | Boolean, whether to annotate functions to the last nodes of the pathways or not. If FALSE, functions will refer to all the nodes of the pathway. |
| unique | Boolean, whether to return the first function for each path. |

Value

List of annotations from pathways to functions

```
get_path_names
```

Description

Translates the subpathway IDs to readable and comprensible names.

For effector subpathways, the names of the subpathways are encoded as "pathway: effector_protein", where "pathway" is the pathway to which the subpathway belongs and "effector_protein" is the name of the last node in the subpathway.

For decomposed subpathways, the names of the subpathways are encoded as "pathway: receptor_protein - effector_protein", where "pathway" is the pathway to which the subpathway belongs, "receptor_protein" is the name of the initial node of the subpathway and "effector_protein" is the name of the last node in the subpathway.

Usage

```
get_path_names(metaginfo, names, maxchar = NULL)
```

Arguments

| metaginfo | Pathways object |
|-----------|---|
| names | Character vector with the subpathway IDs to be translated |
| maxchar | Integer, describes the number of maximum characters to be shown. By default no filter is applied. |

Value

A character vector including the readable names of the subpathways IDs, in the same order as provided.

Examples

```
data(path_vals)
pathways <- load_pathways(species = "hsa", pathways_list = c("hsa03320",
    "hsa04012"))
translated_names <- get_path_names(pathways, rownames(path_vals))</pre>
```

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Gene Ontology matrix of the BRCA gene expression dataset

Description

Matrix of Gene Ontology terms activation values for the BRCA dataset. This matrix is computed from the Results object returned by the hipathia function by means of the quantify_terms function.

Usage

data(go_vals)

heatmap_plot

Format

Matrix with 40 columns and 1654 rows. Row names are Gene Ontology terms and column names are the TCGA identifyers of the samples.

Details

```
go_vals <- quantify_terms(results, pathways, "GO")</pre>
```

Value

Matrix with 40 columns and 1654 rows. Row names are Gene Ontology terms and column names are the TCGA identifyers of the samples.

heatmap_plot Plots subpathways heatmap

Description

Plots a heatmap with the values of the subpathways.

Usage

```
heatmap_plot(
  data,
  group = NULL,
  sel_assay = 1,
  colors = "classic",
  sample_clust = TRUE,
  variable_clust = FALSE,
  labRow = NULL,
  labCol = NULL,
  sample_colors = NULL,
  scale = TRUE,
  save_png = NULL,
  legend = TRUE,
  legend_xy = "topright",
  pch = 15,
  main = NULL
)
```

Arguments

| | Either a SummarizedExperiment or a matrix with the values to be plotted. Rows are features and columns are samples. |
|-----|---|
| | Either a character indicating the name of the column in colData including the classes to plot, or a character vector with the class to which each sample belongs. Samples must be ordered as in data. By default, all samples will be assigned to the same class. |
| - 5 | Character or integer, indicating the assay to be normalized in the Summarized-Experiment. Default is 1. |

| colors | Either a character vector with colors or a key name indicating the color scheme to be used in the heatmap. If a character vector is provided, it is recommended to provide at least 3 colors. Three different predefined color schemes may be selected by providing a key name. Options are: * classic Blue for lower values, white for medium values, red for higher values. * hipathia Hipathia predefined color scheme: Green for lower values, white for medium values, orange for higher values. * redgreen Green for lower values, black for medium values, red for higher values. By default classic color scheme is applied. |
|-------------------------|--|
| <pre>sample_clust</pre> | Boolean, whether to cluster samples (columns). By default TRUE. |
| variable_clust | Boolean, whether to cluster variables (rows). By default FALSE. If TRUE, rows with 0 variance are removed. |
| labRow, labCol | Character vectors with row and column labels to be used. By default row-names(data) or colnames(data) are used, respectively. |
| sample_colors | Named character vector of colors. The names of the colors must be the classes in group. Each sample will be assigned the color corresponding to its class, taken from the group vector. By default a color will be assigned automatically to each class. |
| scale | Boolean, whether to scale each row to the interval [0,1]. Default is TRUE. |
| save_png | Path to the file where the image as PNG will be saved. By default, the image is not saved. |
| legend | Boolean, whether to display a legend. |
| legend_xy | Position for the legend, in case legend is TRUE. |
| pch | Graphical parameter from par() function. |
| main | Main title of the image |
| | |

Value

Heatmap of the values of the subpathways

Examples

```
data(brca_design)
data(path_vals)
sample_group <- brca_design[colnames(path_vals),"group"]
heatmap_plot(path_vals, group = sample_group)
heatmap_plot(path_vals, group = "group", colors = "hipathia",
variable_clust = TRUE)</pre>
```

| Head function for SummarizedExperiment, data.frames and matrix |
|--|
| objects |

Description

hhead

Shows the first n rows and the first n columns of a matrix, in case the matrix has more than n+5 rows or columns. Otherwise, it shows all the rows or columns, respectively.

hidata

Usage

hhead(mat, n = 5, sel_assay = 1)

Arguments

| mat | Object to be shown |
|-----------|--|
| n | Number of rows and columns |
| sel_assay | Character or integer, indicating the assay to be translated in the SummarizedExperiment. Default is 1. |

Value

Matrix with as much as n rows and n columns.

Examples

```
mat <- matrix(rnorm(100), ncol = 10)
hhead(mat)
hhead(mat, 3)
hhead(mat, 7)</pre>
```

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Results object

Description

Results object returned by hipathia::hipathia function, after calling hidata <- hipathia(brca, pathways, verbose=TRUE, uni.terms = TRUE,GO.terms = TRUE)

Usage

data(hidata)

Format

MultiAssayExperiment object of 4 listed experiments, with the activity values of nodes, paths and functional annotations for each sample: Nodes includes a matrix with 6826 rows Paths includes a matrix with 1876 rows Uni.terms includes a matrix with 142 rows GO.terms includes a matrix with 1654 rows

Value

Object of results, including nodes, pathways and functional information.

hipathia

Description

#@importFrom igraph

Usage

```
hipathia(
  genes_vals,
  metaginfo,
  uni.terms = FALSE,
  GO.terms = FALSE,
  sel_assay = 1,
  decompose = FALSE,
  scale = TRUE,
  maxnum = 100,
  verbose = TRUE,
  tol = 1e-06,
  test = TRUE
)
```

Arguments

| A SummarizedExperiment or matrix with the normalized expression values of the genes. Rows represent genes and columns represent samples. Rownames() must be accepted gene IDs. |
|--|
| Pathways object |
| Boolean, whether to compute functional analysis with Uniprot keywords. |
| Boolean, whether to compute functional analysis with Gene Ontology terms. |
| Character or integer, indicating the assay to be processed in the SummarizedExperiment. Only applied if genes_vals is a SummarizedExperiment.Default is 1. |
| Boolean, whether to compute the values for the decomposed subpathways. By default, effector subpathways are computed. |
| Boolean, whether to scale the values matrix to $[0,1]$. Default is TRUE. |
| Number of maximum iterations when iterating the signal through the loops into the pathways |
| Boolean, whether to show details about the results of the execution of hipathia |
| Tolerance for the difference between two iterations when iterating the signal through the loops into the pathways |
| Boolean, whether to test the input objects. Default is TRUE. |
| |

Value

A MultiAssayExperiment object with the level of activation of the subpathways from the pathways in pathigraphs for the experiment with expression values in genes_vals.

igraphs_upgrade

Examples

```
data(exp_data)
pathways <- load_pathways(species = "hsa", pathways_list = c("hsa03320",
    "hsa04012"))
results <- hipathia(exp_data, pathways, verbose = TRUE)
## Not run: results <- hipathia(exp_data, pathways, decompose = TRUE,
verbose = FALSE)
## End(Not run)</pre>
```

igraphs_upgrade Upgrade igraphs to current version

Description

Upgrades the igraph objects in metaginfo object to the corresponding version of the igraph package.

Usage

```
igraphs_upgrade(metaginfo)
```

Arguments

metaginfo Pathways object

Value

The pathways object with the upgraded igraph objects

is_accepted_species Checks whether a species is accepted

Description

Checks whether a species is accepted

Usage

```
is_accepted_species(species)
```

Arguments

| species | Species of the samples. |
|---------|--|
| | #@examples #is_accepted_species("hsa") #is_accepted_species("fca") |

Value

Boolean, whether species is accepted or not.

load_annofuns

Description

Loads annotations object

Usage

load_annofuns(db, species)

Arguments

| db | Database to be used. Either "GO" or "uniprot". |
|---------|--|
| species | Species of the samples. |
| | <pre>#@examples #load_annofuns("GO", "hsa") #load_annofuns("uniprot", "hsa")</pre> |

Value

Annotations object

| load_annots | Loads functional annotations to genes |
|-------------|---------------------------------------|
|-------------|---------------------------------------|

Description

Loads functional annotations from HGNC to the selected database.

Usage

load_annots(db, species)

Arguments

| db | Database to be used. Either "GO" or "uniprot". |
|---------|--|
| species | Species of the samples. |
| | #@examples #load_annots("GO", "hsa") |

Value

Functional annotations from HGNC to the selected database.

load_entrez_hgnc Loads table of translation from HGNC to Entrez

Description

Loads table of translation from HGNC to Entrez

Usage

load_entrez_hgnc(species)

Arguments

| species | Species of the samples. | |
|---------|--|--|
| | <pre>#@examples #load_entrez_hgnc("hsa")</pre> | |

Value

Table of translation from HGNC to Entrez

load_gobp_frame Loads GO graph information

Description

#@examples #load_gobp_frame()

Usage

load_gobp_frame()

Value

GO graph information

load_gobp_net Loads GO graph

Description

#@examples #load_gobp_net()

Usage

load_gobp_net()

Value

GO graph

load_mgi

Description

Loads object with graph information

Usage

load_mgi(species)

Arguments

| species | Species of the samples. |
|---------|-----------------------------|
| | #@examples #load_mgi("hsa") |

Value

Graph information object

| load_pathways | Loads the pathways object. | |
|---------------|----------------------------|--|
|---------------|----------------------------|--|

Description

Loads the pathways object, which includes information about the pathways to be analyzed.

Usage

```
load_pathways(species, pathways_list = NULL)
```

Arguments

| species | Species of the samples. |
|---------------|--|
| pathways_list | Vector of the IDs of the pathways to load. By default all available pathways are |
| | load. |

Details

The object of pathways includes information about the pathways and the subpathways which will be analyzed. This object must be provided to some of the functions (like hipathia or quantify_terms) in the package. These functions will analyze all the pathways included in this object. By default, all available pathways are load. In order to restrict the analysis to a predefined set of pathways, specify the set of pathways to load with the parameter pathways_list.

Value

An pathways object including * species Species to which the pathways are related. * pathigraphs List of Pathigraph objects. Each Pathigraph contains the necessary information of a pathway for it to be analyzed with Hipathia. * all_genes List of all the genes included in the selection of pathways stored in pathigraphs. * eff_norm Vector of normalization values for effector subpathways. * path_norm Vector of normalization values for decomposed subpathways.

load_pseudo_mgi

Examples

```
## Not run: pathways <- load_pathways("hsa") # Loads all pathways for human
pathways <- load_pathways("mmu", c("mmu03320", "mmu04024", "mmu05200"))
    # Loads pathways 03320, 04024 and 05200 for mouse</pre>
```

load_pseudo_mgi Loads object with pseudo graph information

Description

Loads object with pseudo graph information

Usage

load_pseudo_mgi(species, group_by)

Arguments

| species | Species of the samples. |
|----------|---|
| group_by | How to group the subpathways to be visualized. By default they are grouped by the pathway to which they belong. Available groupings include "uniprot", to group subpathways by their annotated Uniprot functions, "GO", to group sub- pathways by their annotated GO terms, and "genes", to group subpathways by the genes they include. #@examples #load_pseudo_mgi("hsa", "uniprot") |

Value

Pseudo graph information object

```
load_xref
```

Loads table of references

Description

Loads table of references

Usage

```
load_xref(species)
```

Arguments

| species | Species of the samples. | |
|---------|---|--|
| | <pre>#@examples #load_xref("hsa")</pre> | |

Value

Table of references

mgi_from_sif

Description

Creates a Pathways object from the information of a pathway stored in a SIF file with some attributes. This pathways object can be used by function hipathia to analyze data.

Usage

```
mgi_from_sif(sif.folder, spe, entrez_symbol = NULL, dbannot = NULL)
```

Arguments

| sif.folder | Path to the folder in which SIF and ATT files are stored. |
|---------------|---|
| spe | Species |
| entrez_symbol | Relation between Entrez (NCBI) genes and gene symbols. Data.frame with 2 columns: First column is the EntrezGene ID, second column is the gene Symbol. The genes in the nodes of the pathways should be defined by Entrez IDs in the SIF and ATT files of the pathways. In order to be more readable, gene names are used when plotting the pathways. |
| dbannot | Functional annotation of the genes in the pathways to create function nodes. |

Value

A pathways object with the same structure of that returned by function load_pathways.

multiple_pca_plot Plots multiple components of a PCA

Description

Plots multiple components of a PCA analysis computed with do_pca

Usage

```
multiple_pca_plot(
    fit,
    group = NULL,
    sample_colors = NULL,
    comps = seq_len(3),
    plot_variance = FALSE,
    legend = TRUE,
    cex = 2,
    pch = 20,
    main = "Multiple PCA plot",
    save_png = NULL
)
```

node_color

Arguments

| fit | princomp object as returned by do_pca |
|---------------|---|
| group | Vector with the group to which each sample belongs. The samples must be ordered as in path_vals. By default, all samples will be assigned to the same class. |
| sample_colors | Named character vector of colors. The names of the colors must be the classes in group. Each sample will be assigned the color corresponding to its class, taken from the group vector. By default a color will be assigned automatically to each class. |
| comps | Vector with the components to be plot |
| plot_variance | Logical, whether to plot the cumulative variance. |
| legend | Boolean, whether to plot a legend in the plot. Default is TRUE. |
| cex | Graphical parameter from par() function. |
| pch | Graphical parameter from par() function. |
| main | Main title of the image |
| save_png | Path to the file where the image as PNG will be saved. By default, the image is not saved. |

Value

Plots multiple components of a PCA

Examples

```
data(path_vals)
sample_group <- brca_design[colnames(path_vals),"group"]
pca_model <- do_pca(path_vals[seq_len(ncol(path_vals)),])
multiple_pca_plot(pca_model, sample_group, cex = 3, plot_variance = TRUE)</pre>
```

node_color

Get colors of the nodes from a comparison file

Description

Computes the colors of the nodes depending on the sign and p.value from the provided file. Significant up- and down-regulated nodes are depicted with the selected color, with a gradient towards the non-significant color depending on the value of the p-value. Smaller p-values give rise to purer colors than higher p-values.

Usage

```
node_color(
   comp,
   metaginfo,
   group_by = "pathway",
   colors = "classic",
   conf = 0.05,
   adjust = TRUE
)
```

Arguments

| Comparison file as returned by do_wilcoxon. Must include a column named "UP/DOWN" with the sign of the comparison coded as UP or DOWN, a column named "p.value" of raw p.values and a column named "FDRp.value" of adjusted p.values. |
|---|
| Object of pathways. |
| How to group the subpathways to be visualized. By default they are grouped by the pathway to which they belong. Available groupings include "uniprot", to group subpathways by their annotated Uniprot functions, "GO", to group sub- pathways by their annotated GO terms, and "genes", to group subpathways by the genes they include. Default is set to "pathway". |
| Either a character vector with 3 colors (indicating, in this order, down-regulation, non-significance and up-regulation colors) or a key name indicating the color scheme to be used. Options are: |
| Level of significance of the comparison for the adjusted p-value. |
| Boolean, whether to adjust the p.value from the comparison. Default is TRUE. |
| |

Value

List of color vectors, named by the pathways to which they belong. The color vectors represent the differential expression of the nodes in each pathway.

Slots

classic ColorBrewer blue, white and colorBrewer red.

hipathia Hipathia predefined color scheme: Green, white and orange. By default classic color scheme is applied.

Examples

```
data(results)
data(brca)
pathways_list <- c("hsa03320", "hsa04012")
pathways <- load_pathways(species = "hsa", pathways_list)
comp <- do_wilcoxon(results[["nodes"]], "group", "Tumor", "Normal")
colors_de <- node_color(comp, pathways)</pre>
```

node_color_per_de Colors of the nodes by its differential expression

Description

Performs a Limma differential expression on the nodes and computes the colors of the nodes depending on it_ Significant up- and down-regulated nodes are depicted with the selected color, with a gradient towards the non-significant color depending on the value of the p-value. Smaller p-values give rise to purer colors than higher p-values. node_color_per_de

Usage

```
node_color_per_de(
  results,
  metaginfo,
  group,
  expdes,
  g2 = NULL,
  group_by = "pathway",
  colors = "classic",
  conf = 0.05,
  adjust = TRUE
)
```

Arguments

| results | Object of results as provided by the hipathia function_ |
|-----------|---|
| metaginfo | Object of pathways_ |
| group | Character indicating the column in which the group variable is stored, in case the object provided to hipathia was a SummarizedExperiment, or a vector with the class to which each sample belongs. Samples must be ordered as in results. |
| expdes | String, either the comparison to be performed or the label of the first group to be compared. |
| g2 | String, label of the second group to be compared. Only necessary in case expdes is the name of the first group, not the comparison. |
| group_by | How to group the subpathways to be visualized. By default they are grouped by the pathway to which they belong. Available groupings include "uniprot", to group subpathways by their annotated Uniprot functions, "GO", to group sub- pathways by their annotated GO terms, and "genes", to group subpathways by the genes they include. Default is set to "pathway". |
| colors | Either a character vector with 3 colors (indicating, in this order, down-regulation, non-significance and up-regulation colors) or a key name indicating the color scheme to be used. Options are: |
| conf | Level of significance of the comparison for the adjusted p-value. |
| adjust | Boolean, whether to adjust the p.value from the comparison. Default is TRUE. |

Value

List of color vectors, named by the pathways to which they belong. The color vectors represent the differential expression of the nodes in each pathway.

Slots

classic ColorBrewer blue, white and colorBrewer red.

hipathia Hipathia predefined color scheme: Green, white and orange. By default classic color scheme is applied.

Examples

```
data(results)
data(brca)
pathways_list <- c("hsa03320", "hsa04012")
pathways <- load_pathways(species = "hsa", pathways_list)
colors_de <- node_color_per_de(results, pathways, "group", "Tumor - Normal")
colors_de <- node_color_per_de(results, pathways, "group", "Tumor", "Normal")</pre>
```

normalize_data

Normalize expression data from a SummarizedExperiment or matrix to be used in hipathia

Description

Transforms the rank of the SummarizedExperiment or matrix of gene expression to [0,1] in order to be processed by hipathia. The transformation may be performed in two different ways. If percentil = FALSE, the transformation is a re-scaling of the rank of the matrix. If percentil = TRUE, the transformation is performed assigning to each cell its percentil in the corresponding distribution. This option is recommended for distributions with very long tails.

Usage

```
normalize_data(
    data,
    sel_assay = 1,
    by_quantiles = FALSE,
    by_gene = FALSE,
    percentil = FALSE,
    truncation_percentil = NULL
)
```

Arguments

| data | Either a SummarizedExperiment or a matrix of gene expression. | |
|----------------------|--|--|
| sel_assay | Character or integer, indicating the assay to be normalized in the Summarized-Experiment. Default is 1. | |
| by_quantiles | Boolean, whether to normalize the data by quantiles. Default is FALSE. | |
| by_gene | Boolean, whether to transform the rank of each row of the matrix to [0,1]. Default is FALSE. | |
| percentil | Boolean, whether to take as value the percentil of each sample in the correspond- ing distribution. | |
| truncation_percentil | | |
| | Real number p in [0,1]. When provided, values beyond percentil p are truncated to the value of percentil p, and values beyond 1-p are truncated to percentil 1-p. By default no truncation is performed. | |

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normalize_paths

Details

This transformation may be applied either to the whole matrix (by setting by_gene = FALSE), which we strongly recommend, or to each of the rows (by setting by_gene = TRUE), allowing each gene to have its own scale.

A previous quantiles normalization may be applied by setting by_quantiles = TRUE. This is recommended for noisy data.

For distributions with extreme outlayer values, a percentil p may be given to the parameter truncation_percentil. When provided, values beyond percentil p are truncated to the value of percentil p, and values beyond 1-p are truncated to percentil 1-p. This step is performed before any other transformation. By default no truncation is performed.

Value

Matrix of gene expression whose values are in [0,1].

Examples

```
data("brca_data")
trans_data <- translate_data(brca_data, "hsa")
exp_data <- normalize_data(trans_data)
exp_data <- normalize_data(trans_data, by_quantiles = TRUE,
truncation_percentil=0.95)</pre>
```

normalize_paths Normalize the pathway matrix by rows

Description

Due to the nature of the Hipathia method, the length of a pathway may influence its signal rank. In order to compare signal values among subpathways, we strongly recommend to normalize the matrix with this normalization.

Usage

normalize_paths(path_vals, metaginfo)

Arguments

| path_vals | SummarizedExperiment or matrix of the pathway values |
|-----------|--|
| metaginfo | Pathways object |

Details

This function removes the bias caused by the length of the subpathways by dividing by the value obtained from running the method with a basal value of 0.5 at each node.

Value

SummarizedExperiment or matrix of normalized pathway values, depending on the class of path_vals.

Examples

```
data(path_vals)
pathways <- load_pathways(species = "hsa", pathways_list = c("hsa03320",
    "hsa04012"))
path_normalized <- normalize_paths(path_vals, pathways)</pre>
```

paths_to_go_ancestor Create path results table with highest significant GO ancestors

Description

Create table of results with the comparison of the paths together with the GO functional annotation and the highest significant GO ancestor (HSGOA).

Usage

```
paths_to_go_ancestor(pathways, comp_paths, comp_go, pval = 0.05)
```

Arguments

| pathways | Pathways object |
|------------|--|
| comp_paths | $Wilcoxon\ comparison\ of\ the\ matrix\ of\ pathways\ values\ as\ returned\ by\ do_wilcoxon.$ |
| comp_go | Wilcoxon comparison of the matrix of GO values as returned by do_wilcoxon. |
| pval | P-value cut-off. Default values is set to 0.05. |

Details

The table returns in each row: the name of a pathway and its Wilcoxon comparison information (direction, adjusted p-value), the GO term to which the path is related (not necessarily unique), the Wilcoxon comparison information for this GO (direction, adjusted p-value), the HSGOA of this GO and its Wilcoxon comparison information (direction, adjusted p-value).

The HSGOA is computed as the GO term with minimum level from all the significant (with respect to value pval) ancestors of a GO. The level of a GO term is computed as the number of nodes in the shortest path from this GO term to the term "GO:0008150". The ancestors of a node are defined as all the nodes from which a path can be defined from the ancestor to the node.

Value

Table of comparisons with Highest common ancestors

Examples

```
data(comp)
data(go_vals)
data(brca_design)
data(path_vals)
sample_group <- brca_design[colnames(path_vals), "group"]
comp_go <- do_wilcoxon(go_vals, sample_group, g1 = "Tumor", g2 = "Normal")
## Not run: pathways <- load_pathways(species = "hsa", pathways_list =
c("hsa03320", "hsa04012"))</pre>
```

pathways

```
table <- paths_to_go_ancestor(pathways, comp, comp_go)
## End(Not run)</pre>
```

pathways

Pathways object including pathways has03320 and hsa04012.

Description

Pathways object returned by hipathia::load_pathways function, after calling pathways <- load_pathways(species = "hsa",pathways_list = c("hsa03320", "hsa04012"))

Usage

data(pathways)

Format

Pathways object

Value

Pathways object including pathways has03320 and hsa04012.

pathway_comparison_plot

Plots pathway with colored significant paths

Description

Plots the layout of a pathway, coloring the significant subpathways in different colors depending on whether they are significantly up- or down-regulated. Nodes may be also colored providing a suitable list of colors for each node. Function node_color_per_de assigns colors to the nodes depending on their differential expression.

Usage

```
pathway_comparison_plot(
   comp,
   metaginfo,
   pathway,
   conf = 0.05,
   node_colors = NULL,
   colors = "classic"
)
```

Arguments

| comp | Comparison data frame as returned by the do_wilcox function. |
|-------------|--|
| metaginfo | Pathways object. |
| pathway | Name of the pathway to be plotted. |
| conf | Level of significance of the comparison for the adjusted p-value. Default is 0.05. |
| node_colors | List, named by the pathway name, including the color of each node for each pathway. |
| colors | Either a character vector with 3 colors (indicating, in this order, down-regulation, non-significance and up-regulation colors) or a key name indicating the color scheme to be used. Options are: |

Value

Image in which a pathway is ploted. Edges are colored so that the UP- and DOWN-activated subpathways are identified.

Slots

classic ColorBrewer blue, white and colorBrewer red.

hipathia Hipathia predefined color scheme: Green, white and orange. By default classic color scheme is applied.

Examples

```
data(comp)
pathways_list <- c("hsa03320", "hsa04012")
pathways <- load_pathways(species = "hsa", pathways_list)
pathway_comparison_plot(comp, metaginfo = pathways, pathway = "hsa03320")
## Not run:
data(results)
data(brca)
colors_de <- node_color_per_de(results, pathways, group, "Tumor", "Normal")
pathway_comparison_plot(comp, metaginfo = pathways, pathway = "hsa04012",
node_colors = colors_de)</pre>
```

End(Not run)

path_vals

Pathways matrix of the BRCA gene expression dataset

Description

Matrix of pathway activation values for the BRCA dataset. This matrix is extracted from the Results object returned by the hipathia function by means of the get_paths_matrix function.

Usage

data(path_vals)

pca_plot

Format

Matrix with 40 columns and 1868 rows. Row names are Pathway IDs and column names are the TCGA identifyers of the samples.

Details

```
path_vals <- get_paths_matrix(results)</pre>
```

Value

Matrix with 40 columns and 1868 rows. Row names are Pathway IDs and column names are the TCGA identifyers of the samples.

pca_plot

Plots two components of a PCA

Description

Plots two components of a PCA computed with do_pca

Usage

```
pca_plot(
    fit,
    group = NULL,
    sample_colors = NULL,
    cp1 = 1,
    cp2 = 2,
    legend = TRUE,
    legend_xy = "bottomleft",
    cex = 2,
    pch = 20,
    mgp = c(3, 1, 0),
    main = "PCA plot",
    save_png = NULL
)
```

Arguments

| fit | princomp object as returned by do_pca |
|---------------|---|
| group | Vector with the group to which each sample belongs. The samples must be ordered as in rownames(fit\$scores). By default, all samples will be assigned to the same class. |
| sample_colors | Named character vector of colors. The names of the colors must be the classes in group. Each sample will be assigned the color corresponding to its class, taken from the group vector. By default a color will be assigned automatically to each class. |
| cp1 | Integer, number of the component in the X-axis. Default is 1, the first component. |

| cp2 | Integer, number of the component in the Y-axis. Default is 2, the second component. |
|-----------|--|
| legend | Boolean, whether to plot a legend in the plot. Default is TRUE. |
| legend_xy | Situation of the legend in the plot. Available options are: "bottomright", "bottom", "bottomleft", "left", "topleft", "top", "topright", "right" and "center". |
| cex | Graphical parameter from par() function. |
| pch | Graphical parameter from par() function. |
| mgp | Graphical parameter from par() function. |
| main | Title of the graphics |
| save_png | Path to the file where the image as PNG will be saved. By default, the image is not saved. |

Value

Plots two components of a PCA

Examples

```
data(path_vals)
sample_group <- brca_design[colnames(path_vals),"group"]
pca_model <- do_pca(path_vals[seq_len(ncol(path_vals)),])
pca_plot(pca_model, sample_group)</pre>
```

| _ | |
|--------|---|
| plotVG | Plots a pathway with or without the comparison information, using |
| | the visNetwork library. |

Description

Plots a pathway with or without the comparison information, using the visNetwork library.

Usage

```
plotVG(
   name,
   pathways,
   DAdata = NULL,
   colors = "hiro",
   conf = 0.05,
   adjust = TRUE,
   main = "Pathway",
   submain = "",
   no.col = "BlanchedAlmond",
   height = "800px"
)
```

quantify_terms

Arguments

| KEGG ID of the pathway to plot. |
|--|
| Pathways object. |
| List of comparison results, returned by function DAcomp. |
| String with the color scheme or vector of colors to be used. See define_colors for available options. Default is "hiro". |
| Numeric, cut off for significance. Default is 0.05. |
| Boolean, whether to adjust the p.value with Benjamini-Hochberg FDR method. Default is TRUE. |
| Title of the plot. |
| Subtitle of the plot. |
| String with the color given to non-significant nodes. |
| Height of the plot. Default is "800px". |
| |

Value

Plot of the pathway.

Examples

```
data(pathways)
plotVG("hsa03320", pathways)
data(DAdata)
plotVG("hsa04012", pathways, DAdata)
```

| quantify_terms | Computes the level of activation of the functions related to the previ- |
|----------------|---|
| | ously computed subpathways |

Description

Computes the level of activation of the functions related to the previously computed subpathways

Usage

```
quantify_terms(
  results,
  metaginfo,
  dbannot,
  out_matrix = FALSE,
  normalize = TRUE
)
```

results

Arguments

| results | List of results as returned by the hipathia function |
|------------|---|
| metaginfo | Pathways object |
| dbannot | Either a string indicating which precomputed annotation to use ("uniprot" for Uniprot Keywords or "GO" for Gene Ontology terms), or a dataframe with the annotation of the genes to the functions. First column are gene symbols, second column the functions. |
| out_matrix | Boolean, whther the output object should be a matrix object. Default is FALSE, returning a SummarizedExperiment object. |
| normalize | Boolean, whether to normalize the matrix of pathway values with normalize_paths before quantifying the signal. Due to the nature of the Hipathia method, in which the length of each pathway may alter its signal rank, we strongly recommend to perform this normalization. This normalization removes the bias. Default is set to TRUE. |

Value

Matrix with the level of activation of the functions in dbannot

Examples

```
data(results)
pathways <- load_pathways(species = "hsa", pathways_list = c("hsa03320",
    "hsa04012"))
go_values <- quantify_terms(results, pathways, "GO")
uniprot_values <- quantify_terms(results, pathways, "uniprot")</pre>
```

results

Results object

Description

Results object returned by hipathia::hipathia function, after calling results <- hipathia(exp_data, pathways, verbose=TRUE)

Usage

```
data(results)
```

Format

Object of results, including pathways information.

Value

Object of results, including pathways information.

save_results Save results to folder

Description

Saves results to a folder. In particular, it saves the matrix of subpathway values, a table with the results of the provided comparison, the accuracy of the results and the .SIF and attributes of the pathways.

Usage

```
save_results(results, comp, metaginfo, output_folder = NULL, path = NULL)
```

Arguments

| results | Results object as returned by the hipathia function. |
|---------------|---|
| comp | Comparison as returned by the do_wilcoxon function. |
| metaginfo | Pathways object |
| output_folder | Name of the folder in which the results will be stored. |
| path | Absolute path to the parent directory in which 'output_folder' will be saved. If it is not provided, it will be created in a temp folder. |

Value

Creates a folder in disk in which all the information to browse the pathway results is stored.

Examples

```
data(results)
data(comp)
pathways <- load_pathways(species = "hsa", pathways_list = c("hsa03320",
    "hsa04012"))
save_results(results, comp, pathways, "output_results")</pre>
```

top_pathways Computes

Computes pathway significance

Description

Performs a test for each pathway checking if the number of significant paths is significant, compared to not having any of the paths as significant.

Usage

```
top_pathways(comp)
```

Arguments

comp

Comparison data frame as returned by the do_wilcoxon function.

Value

Table with the names of the pathways and their p-value for the Fisher test comparing the proportion of significant subpaths vs. 0.

Examples

```
data(comp)
top_pathways(comp)
```

| translate_data | Translation of the rownames IDs of a SummarizedExperiment to En- |
|----------------|--|
| | trez IDs. |

Description

Translates the IDs in the rownames of a SummarizedExperiment to Entrez IDs. For accepted IDs to be transformed see the DOCUMENTATION.

Usage

```
translate_data(data, species, sel_assay = 1, verbose = TRUE)
```

Arguments

| data | Either a SummarizedExperiment object or a matrix of gene expression. |
|-----------|--|
| species | Species of the samples. |
| sel_assay | Character or integer, indicating the assay to be translated in the SummarizedExperiment. Default is 1. |
| verbose | Boolean, whether to show details about the results of the execution. |

Value

Either a SummarizedExperiment or a matrix (depending on the input type) of gene expression with Entrez IDs as rownames.

Examples

```
data("brca_data")
trans_data <- translate_data(brca_data, "hsa")</pre>
```

translate_matrix Translation of the rownames IDs of a matrix to Entrez IDs.

Description

Translates the IDs in the rownames of a matrix to Entrez IDs. For accepted IDs to be transformed see the DOCUMENTATION.

Usage

translate_matrix(exp, species, verbose = TRUE)

Arguments

| exp | Matrix of gene expression. |
|---------|--|
| species | Species of the samples. |
| verbose | Boolean, whether to show details about the results of the execution. |

Value

Matrix of gene expression with Entrez IDs as rownames.

visualize_report Visualize a HiPathia report

Description

Visualize a HiPathia report

Usage

```
visualize_report(output_folder, port = 4000)
```

Arguments

```
output_folderFolder in which results to visualize are storedportPort to use
```

Value

The instructions to visualize a HiPathia report in a web browser

Examples

```
data(comp)
pathways <- load_pathways(species = "hsa", pathways_list = c("hsa03320",
    "hsa04012"))
report <- create_report(comp, pathways, "save_results")
visualize_report(report)
## Not run:
data(results)
data(brca)
sample_group <- colData(brca)[,1]
colors_de <- node_color_per_de(results, pathways,
    sample_group, "Tumor", "Normal")
report <- create_report(comp, pathways, "save_results",
    node_colors = colors_de)
visualize_report(report)
visualize_report(report, port = 5000)</pre>
```

End(Not run)

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