Package 'PathQuant'

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Type Package
Title Annotation of gene-metabolite pairs using topology of KEGG metabolic pathway maps
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Description Modeling of KEGG metabolism pathway maps into compound graph format to compute shortest reactional distance (SRD) for gene-metabolite pairs to assess their biological relevance and output results in data.frame and graphical outputs.
License GNU-3
Imports igraph, XML, ggplot2, data.table, knitr, rmarkdown, scales, stringr, gdata, RCurl, reshape2, stringr, dplyr, plyr, doParallel, parallel, foreach
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annotateAssociationData

Gene annotation classification of associations

Description

Gene annotation involve EC number, kegg brite annotation High level classification made by involved brite trnasporter, enzyme or protein

Usage

annotateAssociationData(associations)

Arguments

association

Dataframe with 2 columns, where each line reprensents an association. First column are the genes and the sencond column are the metabolites. Only use KEGG Ids.

Examples

annotateAssociationData(associations)

exportDFtoTxt

Export results data of a data frame into a text files, for which each lines separates the data from every data frame colum by tabulation. This text file can be easily opened with excel.

Description

Export results data of a data frame into a text files, for which each lines separates the data from every data frame colum by tabulation. This text file can be easily opened with excel.

Usage

```
exportDFtoTxt(export_data, exportNameFile)
```

Arguments

а

dataframe.

```
exportDFtoTxt(dataFrame, exportFileName)
```

3 gene.distribution

gene.distribution Distance distribution plots for single gene

Description

Function ploting the distribution of distances between a gene and all measured metabolite, highlithing the distance of its associated metabolites.

Usage

```
gene.distribution(pathway, gene, pairs, metabolites)
```

Arguments

pathway Selected pathway. Only use KEGG Ids. gene Gene selected. Only use KEGG Ids. pairs Dataframe with 2 columns, where each line reprensents an association. First

column are the genes and the sencond column are the metabolites. Only use

KEGG Ids.

metabolites Dataframe of 1 column containing a lsit of selected metabolites. Default = "All".

All metabolite of the selected pathway map. Only use KEGG Ids.

Details

The plot is depicted as frequency bars, which represent the number of metabolites at a given distance for the selected gene. Frequency bars are shown in grey for metabolites that are not associated with the selected gene and in red if there is at least one metabolite associated with this gene.

If a gene or a metabolite is present on multiple edges or nodes, then the shortest distance is selected.

Examples

```
gene.distribution(hsa01100, "hsa:1373" , pairs.df, metabolite.df)
```

 $is GeneIn Map, \quad metabolite Common Name, \quad metabolite KEGGId, \quad is - metabolite Common Name, \quad metabolite KEGGId, \quad is - metabolite Common Name, \quad metabolite Common Name, \quad metabolite Common Name, \quad metabolite Name, \quad me$ get.srd MetaboliteInMap, distance

Description

isGeneInMap, metaboliteCommonName, metaboliteKEGGId, isMetaboliteInMap, distance

Usage

```
get.srd(pairs, organism_code, pathway = "All", ordered = FALSE,
  commonNames = FALSE, path = FALSE,
  computer_path_to_map_folder = "")
```

4 heatmap

Arguments

pairs dataframe with 2 columns, where each line reprensents a pair. The first column

is genes and the sencond column is metabolites. Only use KEGG Ids.

pathway list of selected metabolic pathway in KEGG. Default = "All". Only use KEGG

Ids.

ordered [option] ascendent ordering of distance

commonNames [option] get KEGG's Common Names of the KEGG Id in the results.

path [option] get reactional path used for srd computation.

Examples

get.srd(pairs.df,"All")

heatmap Function that output a heatmap to visualize distance computed be-

tween every gene-metabolite associations in input using the overview

map only

Description

Function computing shortest distance between every genes and metabolites in a gene-metabolite pairs of your association dataset on a graph model of KEGG map selected, where nodes are metabolites and reactions/genes are edges.

Usage

```
heatmap(association, pathway, computer_path_to_map_folder,
  commonNames = TRUE)
```

Arguments

association dataframe with 2 columns, where each line reprensents a uniq association. First

column are genes and sencond column are metabolites. Only use KEGG Ids.

pathway KEGG Id.

display common names of KEGG ids input (TRUE) or display KEGG Ids for axis de-

scription.

Details

If a gene or a metabolite is present on multiple edges or nodes, then the shortest distance is selected within the group

Output: Heatmap of distances calculated between associated genes-metabolites. Columns represent genes and rows represent metabolites. The calculated distance is shown in each cell with the corresponding color code (from red - closest; to yellow - farthest).

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Examples

heatmap(shinAndAlDF, TRUE)

importTXTtoDF

Importing data from an txt file (with table separators), in the context of this package, one of the data to import is the association file.

Description

Importing data from an txt file (with table separators), in the context of this package, one of the data to import is the association file.

Usage

```
importTXTtoDF(file)
```

Arguments

path

to txt file.

Examples

```
inputXLSXtoDF("file_path/fileName.xlsx")
```

importXLSXtoDF

Importing data from an XSLX file, in the context of this package, one of the data to import is the association file.

Description

Importing data from an XSLX file, in the context of this package, one of the data to import is the association file.

Usage

```
importXLSXtoDF(file, sheet.num = 1)
```

Arguments

path

to XSLX file.

```
inputXLSXtoDF("file_path/fileName.xlsx")
```

6 permutationTest

permutationTest	Statistical permutation test to asses the relevance of associated gene- metabolite pairs vs. randomly selected pairs

Description

Permutation test to evauluate if gene-metabolite associations are significantly closer than randomly selected gene-metabolite pairs.

Usage

```
permutationTest(pathwayId, association, gene, metabolite, permutation,
  output = c("medians", "pvalue", "histogram"))
```

Arguments

pathwayId	KEGG Id of selected pathway.
association	Dataframe with 2 columns, where each line reprensents an association. First column are genes and the sencond column are metabolites. Only use KEGG Ids.
gene	Dataframe of 1 column, representing all genes reported. Only use KEGG Ids.
metabolite	Dataframe of 1 column, representing all the measured metabolites. Only use KEGG Ids.
permutation	Number desired permutations
output	'medians' of all permutations, 'pvalue' of the permutation test, 'histogram' representing the distribution of all permutations' median

Details

If a gene or a metabolite is present on multiple edges or nodes, then the shortest distance is selected.

srd.distribution 7

srd.distribution

srd.distribution

Description

Function ploting the distribution of srds computed by get.srd function.

Usage

```
srd.distribution(distance)
```

Arguments

distance

list of computed srd values of the data.frame obtained by get.srd function

Details

The plot is depicted as frequency bars, which represent the number of pairs at a given srd value. Frequency bars are shown in heatmap colors to depict the small from larger srd values.

```
srd.distribution(res[,7])
```

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