

Package ‘KEGGprofile’

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Type Package

Title An annotation and visualization package for multi-types and multi-groups expression data in KEGG pathway

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Description KEGGprofile is an annotation and visualization tool which integrated the expression profiles and the function annotation in KEGG pathway maps. The multi-types and multi-groups expression data can be visualized in one pathway map. KEGGprofile facilitated more detailed analysis about the specific function changes inner pathway or temporal correlations in different genes and samples.

License GPL (>= 2)

LazyLoad yes

Imports AnnotationDbi,png,TeachingDemos,XML,KEGG.db,KEGGREST,biomaRt

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biocViews Pathways, KEGG

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| | |
|--------------|---------------------|
| col_by_value | <i>col_by_value</i> |
|--------------|---------------------|

Description

The function will transfer a numeric matrix into a matrix of colors, in which the colors represent the values of numeric matrix

Usage

```
col_by_value(x, col, range = NA, breaks = NA, showColorBar = T)
```

Arguments

| | |
|--------------|---|
| x | a numeric matrix |
| col | colors used to represent the values. (See also 'Details') |
| range | values out of the range will be modified to in the range. |
| breaks | a numeric vector of three or more cut points giving the number of intervals into which x is to be cut. See also 'Details' |
| showColorBar | Logical. Indicates display the colorbar or not. The default value is TRUE. |

Details

A colorbar would also be plotted. The returned colors of the function can be used in function plot_profile. if breaks not equal to NA, col must have the same length with breaks-1.

Value

a matrix equal to x, but the values were instead by colors.

Examples

```
data(pho_sites_count)
col<-col_by_value(pho_sites_count,col=colorRampPalette(c('white','khaki2'))(4),breaks=c(0,1,4,10,Inf))
```

 convertId

convertId

Description

A function to convert ID based on the biomaRt package.

Usage

```
convertId(x, dataset = "hsapiens_gene_ensembl",
  filters = "uniprotswissprot", attributes = c(filters, "entrezgene"),
  genesKept = c("foldchange", "first", "random", "var", "abs"),
  keepNoId = T, keepMultipleId = F, verbose = F)
```

Arguments

| | |
|----------------|---|
| x | the expression data matrix. |
| dataset | Dataset you want to use. To see the different datasets available within a biomaRt you can e.g. do: mart = useMart('ensembl'), followed by listDatasets(mart). |
| filters | Filters (one or more) that should be used in the query. A possible list of filters can be retrieved using the function listFilters. |
| attributes | Attributes you want to retrieve. A possible list of attributes can be retrieved using the function listAttributes. |
| genesKept | The method to select target gene in more than one targets. "var"/"foldchange"/"abs" means selecting the gene with largest variation/fold change/absolute value. "first" means selecting the first target and "random" means randomly selection. |
| keepNoId | Logical. Indicate keep the source IDs without target IDs or not. |
| keepMultipleId | Logical. Indicate keep the multiple target IDs related to one source ID or not. |
| verbose | Logical. Indicate report extra information on progress or not. |

Details

A function to convert ID based on the biomaRt package..

Examples

```
temp<-cbind(rnorm(10),rnorm(10))
row.names(temp)<-c("Q04837", "P0C0L4", "P0C0L5", "075379", "Q13068", "A2MYD1", "P60709", "P30462", "P30475", "P30475")
colnames(temp)<-c("Exp1", "Exp2")
convertId(temp, filters="uniprotswissprot", keepMultipleId=TRUE)
## Not run:
temp<-cbind(rnorm(5000), rnorm(5000), rnorm(5000), rnorm(5000), rnorm(5000), rnorm(5000))
row.names(temp)<-1000:5999
colnames(temp)<-c("Control1", "Control2", "Control3", "Treatment1", "Treatment2", "Treatment3")
convertId(temp, filters="entrezgene", attributes =c("entrezgene", "uniprot_swissprot"), keepNoId=FALSE)

## End(Not run)
```

| | |
|-------------------|--------------------------|
| download_KEGGfile | <i>download_KEGGfile</i> |
|-------------------|--------------------------|

Description

The function download XML files and png files from KEGG website to local disk

Usage

```
download_KEGGfile(pathway_id = "00010", species = "hsa",
  target_dir = getwd())
```

Arguments

| | |
|------------|---|
| pathway_id | the KEGG pathway id, such as '00010' |
| species | the species id in KEGG database, 'hsa' means human, 'mmu' means mouse, 'rno' means rat, etc |
| target_dir | the local directory where the downloaded files are saved |

Details

If pathway_id is set as 'all', all KEGG pathway ids in KEGG.db package will be used and downloaded from KEGG website

Examples

```
download_KEGGfile(pathway_id="00010",species='hsa')
```

| | |
|-------------------------|--------------------------------|
| download_latest_pathway | <i>download_latest_pathway</i> |
|-------------------------|--------------------------------|

Description

The function will download the latest pathway gene link from KEGG website.

Usage

```
download_latest_pathway(species)
```

Arguments

| | |
|---------|---|
| species | the species id in KEGG database, 'hsa' means human, 'mmu' means mouse, 'rno' means rat, etc |
|---------|---|

Details

The function will download the latest pathway gene link from KEGG website.

Value

a list with two parts

```

name keggpathway2gene
      description a list with the genes for each pathway
name pathway2name
      description a list with the names for each pathway

```

Examples

```
## Not run: download_latest_pathway(species="hsa")
```

```
find_enriched_pathway find_enriched_pathway
```

Description

The function will map the genes in KEGG pathway database, and then hypergeometric tests would be used to estimate the significance of enrichment for each pathway

Usage

```
find_enriched_pathway(gene, species = "hsa", returned_pvalue = 0.01,
  returned_adjpvalue = 0.05, returned_genenumber = 5,
  download_latest = FALSE, refGene = NULL)
```

Arguments

```

gene          a numeric matrix
species       the species id in KEGG database, 'hsa' means human, 'mmu' means mouse,
              'rno' means rat, etc
returned_pvalue
              the minimum p value for enriched pathways
returned_adjpvalue
              the minimum adjusted p value for enriched pathways
returned_genenumber
              the minimum number of annotated genes for enriched pathways
download_latest
              logical. Indicate if the function will download the latest pathway/gene link from
              KEGG website. As the KEGG.db package was not updated for a long time due
              to the KEGG policy change, we provided this parameter so that the users could
              get the latest KEGG database.
refGene       the names of genes used as reference. If not provided, all genes in KEGG
              database will be used.

```

Details

Only the pathways with p value \leq returned_pvalue in hypergeometric tests and number of annotated genes \geq returned_genenumber would be taken as enriched and returned.

Value

a list with two parts

| | | | |
|------|---------|-------------|--|
| name | stastic | description | a matrix containing the pathway IDs of enriched pathways, and their names, p values, number of annotated genes |
| name | detail | description | a list with the genes annotated for each pathway |

Examples

```
data(pho_sites_count)
#the 300 genes with most phosphorylation sites quantified
genes<-names(rev(sort(pho_sites_count[,1]))[1:300])
pho_KEGGresult<-find_enriched_pathway(genes,species='hsa')
```

newIdMatrix

newIdMatrix

Description

A function to convert ID.

Usage

```
newIdMatrix(x, convertIdTable, genesKept = c("var", "foldchange", "abs",
      "first", "random"))
```

Arguments

| | |
|----------------|---|
| x | the expression data matrix. |
| convertIdTable | A vector. The names should be the source IDs, and the values should be the target IDs. |
| genesKept | The method to select target gene in more than one targets. "var"/"foldchange"/"abs" means selecting the gene with largest variation/fold change/absolute value. "first" means selecting the first target and "random" means randomly selection. |

Details

A function to convert ID.

Examples

```
convertIdTable<-paste("New",c(1,2,2,2,1,3,4,4,5,5))
names(convertIdTable)<-paste("Old",1:length(convertIdTable))
temp<-matrix(rnorm(20),ncol=2)
row.names(temp)<-names(convertIdTable)
colnames(temp)<-c("Exp1","Exp2")
newIdMatrix(temp,genesKept="foldchange",convertIdTable)
```

| | |
|---------------|----------------------|
| parse_XMLfile | <i>parse_XMLfile</i> |
|---------------|----------------------|

Description

The function parses KEGG XML (KGML) files

Usage

```
parse_XMLfile(pathway_id, species, database_dir = getwd())
```

Arguments

| | |
|--------------|---|
| pathway_id | the KEGG pathway id, such as '00010' |
| species | the species id in KEGG database, 'hsa' means human, 'mmu' means mouse, 'rno' means rat, etc |
| database_dir | the directory where the XML files and png files are located |

Details

This function will parse the KEGG XML (KGML) file. Then a matrix with genes in this pathway and related informations will be returned. This matrix can be used for plot the expression profiles on the pathway figure.

Value

a matrix containing genes in this pathway, and their names, locations etc, which could be used in the function plot_profile as param KEGG_database

Examples

```
XML2database<-parse_XMLfile(pathway_id="04110",species="hsa",database_dir=system.file("extdata",package="K
```

| | |
|-----------------|---|
| pho_sites_count | <i>number of phosphorylation sites quantified for each gene</i> |
|-----------------|---|

Description

This data set is a data.frame with number of phosphorylation sites quantified for each gene in the analysis.

Usage

```
pho_sites_count
```

Source

Olsen, J.V., et al. (2010) Quantitative phosphoproteomics reveals widespread full phosphorylation site occupancy during mitosis, Sci Signal, 3, ra3.

| | |
|--------------|---------------------|
| plot_pathway | <i>plot_pathway</i> |
|--------------|---------------------|

Description

A wrapper for function `download_KEGGfile`, `parse_XMLfile` and `plot_profile`

Usage

```
plot_pathway(gene_expr, line_col, groups, pathway_id = "00010",
             species = "hsa", pathway_min = 5, database_dir = getwd(),
             speciesRefMap = TRUE, ...)
```

Arguments

| | |
|----------------------------|---|
| <code>gene_expr</code> | the matrix for gene expression, row.names should be NCBI gene ID, such as 67040, 93683 |
| <code>line_col</code> | line color for expression in different samples in the pathway map, valid when <code>type='lines'</code> |
| <code>groups</code> | a character used to indicate expression values from different types of samples |
| <code>pathway_id</code> | the KEGG pathway id, such as '00010' |
| <code>species</code> | the species id in KEGG database, 'hsa' means human, 'mmu' means mouse, 'rno' means rat, etc |
| <code>pathway_min</code> | The pathways with number of annotated genes less than <code>pathway_min</code> would be ignored |
| <code>database_dir</code> | the directory where the XML files and png files are located |
| <code>speciesRefMap</code> | Logical, use the species specific figure as reference map. if set as FALSE, the reference pathway figure without species information will be used |
| <code>...</code> | any other Arguments for function <code>plot_profile</code> |

Details

This wrapper function is developed to make the visualization process more easier. Firstly the existence of XML file and png file would be checked, if not, the `download_KEGGfile` function would be used to download the files. Then the `parse_XMLfile` function would be used to parse the XML file. At last the `plot_profile` function would be used to generate the pathway map.

See Also

[download_KEGGfile](#), [parse_XMLfile](#), [plot_profile](#)

Examples

```
data(pro_pho_expr)
data(pho_sites_count)
#type='lines'
col<-col_by_value(pho_sites_count,col=colorRampPalette(c('white','khaki2'))(4),breaks=c(0,1,4,10,Inf))
temp<-plot_pathway(pro_pho_expr,bg_col=col,line_col=c("brown1","seagreen3"),groups=c(rep("Proteome ",6),rep(" ",6)))
#type='bg'
pho_expr<-pro_pho_expr[,7:12]
```



```

temp<-apply(pho_expr,1,function(x) length(which(is.na(x))))
pho_expr<-pho_expr[which(temp==0),]
col<-col_by_value(pho_expr,col=colorRampPalette(c('green','black','red'))(1024),range=c(-6,6))
temp<-plot_pathway(pho_expr,type="bg",bg_col=col,text_col="white",magnify=1.2,species='hsa',database_dir=sy
#Compound and gene data
set.seed(124)
testData1<-rbind(rnorm(6),rnorm(6),rnorm(6),rnorm(6),rnorm(6),rnorm(6),rnorm(6),rnorm(6))
row.names(testData1)<-c("4967","55753","1743","8802","47","50","cpd:C15972","cpd:C16255")
colnames(testData1)<-c("Control0","Control2","Control5","Sample0","Sample2","Sample5")
temp<-plot_pathway(testData1,type="lines",line_col=c("brown1","seagreen3"),groups=c(rep("Control",3),rep("S
testData2<-testData1[,4:6]-testData1[,1:3]
col<-col_by_value(testData2,col=colorRampPalette(c('green','black','red'))(1024),range=c(-2,2))
temp<-plot_pathway(testData2,type="bg",bg_col=col,text_col="white",magnify=1.2,species='hsa',database_dir=s

```

plot_pathway_cor

plot_pathway_cor

Description

The function will plot the correlation distributions for each enriched pathway (result from `find_enriched_pathway` function), and then Wilcoxon tests would be used to estimate the significance of correlations distribution between genes in each pathway and all genes.

Usage

```
plot_pathway_cor(gene_expr, kegg_enriched_pathway, groups = NULL,
  side = c("both", "pos", "neg"), alternative = NULL)
```

Arguments

| | |
|------------------------------------|---|
| <code>gene_expr</code> | the matrix for gene expression, row.names should be NCBI gene ID, such as 67040, 93683 |
| <code>kegg_enriched_pathway</code> | The returned value from <code>find_enriched_pathway</code> function, the enriched pathways. |
| <code>groups</code> | a character used to indicate expression values from different types of samples |
| <code>side</code> | a character string specifying the correlation directions interested, must be one of "both" (default), "pos" or "neg". |
| <code>alternative</code> | a character string specifying the alternative hypothesis, must be one of "two.sided" (default), "greater" or "less". You can specify just the initial letter. |

Value

p values for Wilcoxon tests in each pathway

Examples

```

data(pro_pho_expr)
data(pho_sites_count)
genes<-row.names(pho_sites_count)[which(pho_sites_count>=10)]
pho_KEGGresult<-find_enriched_pathway(genes,species='hsa')
result<-plot_pathway_cor(gene_expr=pro_pho_expr,kegg_enriched_pathway=pho_KEGGresult)

```

| | |
|--------------|---------------------|
| plot_profile | <i>plot_profile</i> |
|--------------|---------------------|

Description

The function plot gene expression profiles on KEGG pathway maps

Usage

```
plot_profile(gene_expr, pathway_name, result_name = paste(pathway_name,
  "_profile_", type, ".png", sep = ""), KEGG_database, groups,
  bg_col = "white", text_col = "black", line_col, border_col = "grey",
  text_cex = 0.25, magnify = 1, type = c("lines", "bg"),
  pathway_min = 5, genes_kept = c("foldchange", "first", "random", "var",
  "abs"), species = "hsa", database_dir = getwd(), max_dist, lwd = 1.2,
  speciesRefMap = TRUE)
```

Arguments

| | |
|---------------|--|
| gene_expr | the matrix for gene expression, row.names should be NCBI gene ID, such as 67040, 93683 |
| pathway_name | the species id and KEGG pathway id, such as 'hsa00010' |
| result_name | the name of figure file generated by KEGGprofile. The default name is pathway_name+'_profile_'+type+'.png', such as 'hsa04110_profile_lines.png' |
| KEGG_database | the matrix returned by function parse_XMLfile, which contains genes in this pathway, and their names, locations etc |
| groups | a character used to indicate expression values from different types of samples |
| bg_col | background color for gene rectangles in the pathway map |
| text_col | the colors for text in the pathway map. A color matrix generated by function col_by_value can be used here |
| line_col | line color for expression in different samples in the pathway map, valid when type='lines' |
| border_col | border color for gene rectangles in the pathway map. A color matrix generated by function col_by_value can be used here |
| text_cex | cex for text in the pathway map. A color matrix generated by function col_by_value can be used here |
| magnify | the coefficient used to magnify the gene rectangles |
| type | the type of pathway map visualization, could be 'bg' or 'lines'. Default is 'bg'. See also 'Details' |
| pathway_min | The pathways with number of annotated genes less than pathway_min would be ignored |
| genes_kept | methods used for choosing genes when several genes corresponding to one location in pathway map. Default is 'foldchange', which kept the gene with largest fold changes. 'first' kept the first gene. 'random' chosed gene random. 'var' kept the gene with largest variation. 'abs' kept the gene with largest absolute value |

| | |
|---------------|--|
| species | the species id in KEGG database, 'hsa' means human, 'mmu' means mouse, 'rno' means rat, etc |
| database_dir | the directory where the XML files and png files are located |
| max_dist | The expression changes that represented by the distance from the bottom to the top of gene rectangle, valid when type='lines'. This param is used to ensure the dynamic changes of lines in different gene polygon represent equal variation. It would be calculated from the maximum changes of genes in this pathway by default. If max_dist=NA, then the lines would be plotted from top to bottom in each gene rectangle |
| lwd | The line width when type='lines' |
| speciesRefMap | Logical, use the species specific figure as reference map. if set as FALSE, the reference pathway figure without species information will be used |

Details

There are two visualization methods to represent gene expression profiles: 'background' and 'lines'. The first one is applicable for analysis with only one sample or one type of data, which divides the gene polygon into several sub-polygons to represent different time points. And each sub-polygon has a specific background color to represent expression changes in that time point. The second method plots lines with different colors in the gene polygon to represent different samples or different types of data. The dynamic changes of lines mean the profiles of genes in different time points.

Value

a matrix containing genes mapped in this pathway, and their names, expressions

Examples

```
XML2database<-parse_XMLfile(pathway_id="04110",species="hsa",database_dir=system.file("extdata",package="KEGG_KEGG"),
data(pro_pho_expr)
temp<-plot_profile(pro_pho_expr,pathway_name="hsa04110",KEGG_database=XML2database,line_col=c("brown1","sea
```

| | |
|--------------|--|
| pro_pho_expr | <i>expression profiles in proteome and phosphoproteome</i> |
|--------------|--|

Description

This data set is from a previously published data of proteome and phosphoproteome analysis in different cell phase. The column 1-6 are proteome data and column 7-12 are phosphoproteome data in this data.frame. The 6 time points are G1, G1/S, Early S, Late S, G2, Mitosis.

Usage

```
pro_pho_expr
```

Source

Olsen, J.V., et al. (2010) Quantitative phosphoproteomics reveals widespread full phosphorylation site occupancy during mitosis, Sci Signal, 3, ra3.

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