Package 'clusterProfiler'

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Type Package

for genes and gene clusters
Version 3.6.0
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Description This package implements methods to analyze and visualize functional profiles (GO and KEGG) of gene and gene clusters.
Depends R (>= 3.3.1), DOSE (>= 3.3.2)
Imports AnnotationDbi, ggplot2, GO.db, GOSemSim (>= 2.0.0), magrittr, methods, plyr, qvalue, rvcheck, stats, stats4, tidyr, utils
Suggests AnnotationHub, GSEABase, KEGG.db, knitr, org.Hs.eg.db, prettydoc, pathview, ReactomePA, testthat, topGO
VignetteBuilder knitr
ByteCompile true
License Artistic-2.0
<pre>URL https://guangchuangyu.github.io/clusterProfiler</pre>
BugReports https://github.com/GuangchuangYu/clusterProfiler/issues
biocViews Annotation, Clustering, GeneSetEnrichment, GO, KEGG, MultipleComparison, Pathways, Reactome, Visualization
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Author Guangchuang Yu [aut, cre], Li-Gen Wang [ctb], Giovanni Dall'Olio [ctb] (formula interface of compareCluster)
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clusterProfiler-package

statistical analysis and visualization of functional profiles for genes and gene clusters The package implements methods to analyze and visualize functional profiles of gene and gene clusters.

Description

statistical analysis and visualization of functional profiles for genes and gene clusters The package implements methods to analyze and visualize functional profiles of gene and gene clusters.

bitr 3

bitr bitr

Description

Biological Id TRanslator

Usage

```
bitr(geneID, fromType, toType, OrgDb, drop = TRUE)
```

Arguments

geneID input gene id
fromType input id type
toType output id type
OrgDb annotation db
drop drop NA or not

Value

data.frame

Author(s)

Guangchuang Yu

bitr_kegg bitr_kegg

Description

convert biological ID using KEGG API

Usage

```
bitr_kegg(geneID, fromType, toType, organism, drop = TRUE)
```

Arguments

geneID input gene id fromType input id type toType output id type

organism supported organism, can be search using search_kegg_organism function

drop NA or not

Value

data.frame

4 buildGOmap

Author(s)

Guangchuang Yu

browseKEGG

browseKEGG

Description

open KEGG pathway with web browser

Usage

```
browseKEGG(x, pathID)
```

Arguments

x an instance of enrichResult or gseaResult

pathID pathway ID

Value

url

Author(s)

Guangchuang Yu

buildGOmap

buildGOmap

Description

building GO mapping files

Usage

buildGOmap(gomap)

Arguments

gomap

data.frame with two columns of GO and gene ID

Details

provided by a data.frame of GO (column 1) and gene (column 2) direct annotation this function will building gene to GO and GO to gene mapping, with directly and undirectly (ancestor GO term) annotation.

Value

data.frame, GO annotation with indirect annotation

compareCluster 5

Author(s)

Yu Guangchuang

compareCluster

Compare gene clusters functional profile

Description

Given a list of gene set, this function will compute profiles of each gene cluster.

Usage

```
compareCluster(geneClusters, fun = "enrichGO", data = "", ...)
```

Arguments

```
geneClusters a list of entrez gene id. Alternatively, a formula of type Entrez~group

fun One of "groupGO", "enrichGO", "enrichKEGG", "enrichDO" or "enrichPathway".

data if geneClusters is a formula, the data from which the clusters must be extracted.

Other arguments.
```

Value

A clusterProfResult instance.

Author(s)

```
Guangchuang Yu https://guangchuangyu.github.io
```

See Also

```
compareClusterResult-class, groupGO enrichGO
```

Examples

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compareClusterResult-class

Class "compareClusterResult" This class represents the comparison result of gene clusters by GO categories at specific level or GO enrichment analysis.

Description

Class "compareClusterResult" This class represents the comparison result of gene clusters by GO categories at specific level or GO enrichment analysis.

Slots

```
compareClusterResult cluster comparing result geneClusters a list of genes fun one of groupGO, enrichGO and enrichKEGG .call function call
```

Author(s)

```
Guangchuang Yu https://guangchuangyu.github.io
```

See Also

groupGOResult enrichResult compareCluster

DataSet

Datasets gcSample contains a sample of gene clusters.

Description

Datasets gcSample contains a sample of gene clusters.

Datasets kegg_species contains kegg species information

```
{\tt dotplot}, {\tt compareClusterResult-method} \\ {\tt dotplot}
```

Description

dot plot method

Usage

```
## S4 method for signature 'compareClusterResult'
dotplot(object, x = ~Cluster,
   colorBy = "p.adjust", showCategory = 5, by = "geneRatio",
   split = NULL, includeAll = TRUE, font.size = 12, title = "")
```

Arguments

object compareClusterResult object

x x variable

colorBy one of pvalue or p.adjust

showCategory category numbers

by one of geneRatio, Percentage or count

split ONTOLOGY or NULL

includeAll logical font.size font size title figure title

download_KEGG

 $download_KEGG$

Description

download the latest version of KEGG pathway/module

Usage

```
download_KEGG(species, keggType = "KEGG", keyType = "kegg")
```

Arguments

species species

keggType one of 'KEGG' or 'MKEGG' keyType supported keyType, see bitr_kegg

Value

list

8 enrichDAVID

Author(s)

Guangchuang Yu

dropG0

dropGO

Description

drop GO term of specific level or specific terms (mostly too general).

Usage

```
dropGO(x, level = NULL, term = NULL)
```

Arguments

x an instance of 'enrichResult' or 'compareClusterResult'

level GO level term GO term

Value

modified version of x

Author(s)

Guangchuang Yu

enrichDAVID

enrichDAVID

Description

```
enrichment analysis by DAVID
```

Usage

```
enrichDAVID(gene, idType = "ENTREZ_GENE_ID", universe, minGSSize = 10,
  maxGSSize = 500, annotation = "GOTERM_BP_FAT", pvalueCutoff = 0.05,
  pAdjustMethod = "BH", qvalueCutoff = 0.2, species = NA, david.user)
```

enricher 9

Arguments

gene input gene idType id type

universe background genes

minGSSize minimal size of genes annotated for testing maxGSSize maximal size of genes annotated for testing

annotation david annotation pvalueCutoff pvalueCutoff

pAdjustMethod one of "BH" and "bonferroni"

qvalueCutoff
species species
david.user david user

Value

A enrichResult instance

Author(s)

Guangchuang Yu

enricher enricher

Description

A universal enrichment analyzer

Usage

```
enricher(gene, pvalueCutoff = 0.05, pAdjustMethod = "BH", universe,
  minGSSize = 10, maxGSSize = 500, qvalueCutoff = 0.2, TERM2GENE,
  TERM2NAME = NA)
```

Arguments

gene a vector of gene id pvalueCutoff pvalue cutoff

pAdjustMethod one of "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none"

universe background genes

minGSSize minimal size of genes annotated for testing maxGSSize maximal size of genes annotated for testing

qvalueCutoff qvalue cutoff

TERM2GENE user input annotation of TERM TO GENE mapping, a data.frame of 2 column

with term and gene

TERM2NAME user input of TERM TO NAME mapping, a data frame of 2 column with term

and name

10 enrichGO

Value

A enrichResult instance

Author(s)

Guangchuang Yu

enrichGO GO Enrichment Analysis of a gene set. Given a vector of genes, this

function will return the enrichment GO categories after FDR control.

Description

GO Enrichment Analysis of a gene set. Given a vector of genes, this function will return the enrichment GO categories after FDR control.

Usage

```
enrichGO(gene, OrgDb, keyType = "ENTREZID", ont = "MF",
   pvalueCutoff = 0.05, pAdjustMethod = "BH", universe, qvalueCutoff = 0.2,
   minGSSize = 10, maxGSSize = 500, readable = FALSE, pool = FALSE)
```

Arguments

gene a vector of entrez gene id.

OrgDb OrgDb

keyType keytype of input gene

ont One of "MF", "BP", and "CC" subontologies.

pvalueCutoff Cutoff value of pvalue.

pAdjustMethod one of "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none"

universe background genes qvalueCutoff qvalue cutoff

minGSSize minimal size of genes annotated by Ontology term for testing.

maxGSSize maximal size of genes annotated for testing readable whether mapping gene ID to gene Name

pool If ont='ALL', whether pool 3 GO sub-ontologies

Value

An enrichResult instance.

Author(s)

```
Guangchuang Yu https://guangchuangyu.github.io
```

See Also

```
enrichResult-class, compareCluster
```

enrichKEGG 11

Examples

```
## Not run:
    data(geneList)
de <- names(geneList)[1:100]
yy <- enrichGO(de, 'org.Hs.eg.db', ont="BP", pvalueCutoff=0.01)
head(yy)
## End(Not run)</pre>
```

enrichKEGG

KEGG Enrichment Analysis of a gene set. Given a vector of genes, this function will return the enrichment KEGG categories with FDR control.

Description

KEGG Enrichment Analysis of a gene set. Given a vector of genes, this function will return the enrichment KEGG categories with FDR control.

Usage

```
enrichKEGG(gene, organism = "hsa", keyType = "kegg", pvalueCutoff = 0.05,
  pAdjustMethod = "BH", universe, minGSSize = 10, maxGSSize = 500,
  qvalueCutoff = 0.2, use_internal_data = FALSE)
```

Arguments

gene a vector of entrez gene id.

organism supported organism listed in 'http://www.genome.jp/kegg/catalog/org_list.html'

keyType one of "kegg", 'ncbi-geneid', 'ncib-proteinid' and 'uniprot'

pvalueCutoff Cutoff value of pvalue.

pAdjustMethod one of "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none"

universe background genes

minGSSize minimal size of genes annotated by Ontology term for testing.

maxGSSize maximal size of genes annotated for testing

qvalueCutoff qvalue cutoff

use_internal_data

logical, use KEGG.db or latest online KEGG data

Value

A enrichResult instance.

Author(s)

```
Guangchuang Yu http://ygc.name
```

See Also

```
enrichResult-class, compareCluster
```

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Examples

```
data(geneList)
    de <- names(geneList)[1:100]
yy <- enrichKEGG(de, pvalueCutoff=0.01)
head(yy)
#plot(yy)</pre>
```

enrichMKEGG KEGG Module Enrichment Analysis of a gene set. Given a vector of

genes, this function will return the enrichment KEGG Module cate-

gories with FDR control.

Description

KEGG Module Enrichment Analysis of a gene set. Given a vector of genes, this function will return the enrichment KEGG Module categories with FDR control.

Usage

```
enrichMKEGG(gene, organism = "hsa", keyType = "kegg", pvalueCutoff = 0.05,
   pAdjustMethod = "BH", universe, minGSSize = 10, maxGSSize = 500,
   qvalueCutoff = 0.2)
```

Arguments

gene a vector of entrez gene id.

organism supported organism listed in 'http://www.genome.jp/kegg/catalog/org_list.html'

keyType one of "kegg", 'ncbi-geneid', 'ncib-proteinid' and 'uniprot'

pvalueCutoff Cutoff value of pvalue.

pAdjustMethod one of "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none"

universe background genes

minGSSize minimal size of genes annotated by Ontology term for testing.

maxGSSize maximal size of genes annotated for testing

qvalueCutoff qvalue cutoff

Value

A enrichResult instance.

Description

convert compareClusterResult to a data.frame that ready for plot

Usage

```
## $3 method for class 'compareClusterResult'
fortify(model, data, showCategory = 5,
   by = "geneRatio", split = NULL, includeAll = TRUE)
```

Arguments

model compareClusterResult object

data not use here

showCategory category numbers

by one of geneRatio, Percentage or count

split ONTOLOGY or NULL

includeAll logical

Value

data.frame

Author(s)

Guangchuang Yu

getGOLevel

get GOIDs at a specific level

Description

```
query GOIDs at a specific level.
```

Usage

```
getGOLevel(ont, level)
```

Arguments

ont Ontology level GO level go2ont

Value

a vector of GOIDs

Author(s)

Guangchuang Yu http://guangchuangyu.github.io

Gff2GeneTable

Gff2GeneTable

Description

read GFF file and build gene information table

Usage

```
Gff2GeneTable(gffFile, compress = TRUE)
```

Arguments

gffFile GFF file

compress file or not

Details

given a GFF file, this function extracts information from it and save it in working directory

Value

file save.

Author(s)

Yu Guangchuang

go2ont

go2ont

Description

convert goid to ontology (BP, CC, MF)

Usage

go2ont(goid)

Arguments

goid

a vector of GO IDs

go2term 15

Value

data.frame

Author(s)

Guangchuang Yu

go2term

go2term

Description

convert goid to descriptive term

Usage

go2term(goid)

Arguments

goid

a vector of GO IDs

Value

data.frame

Author(s)

Guangchuang Yu

gofilter

gofilter

Description

filter GO enriched result at specific level

Usage

```
gofilter(x, level = 4)
```

Arguments

Х

output from enrichGO or compareCluster

level

GO level

Value

updated object

Author(s)

Guangchuang Yu

16 groupGO

groupGO	Functional Profile of a gene set at specific GO level. Given a vector of genes, this function will return the GO profile at a specific level.
	genes, in a function with return the Go profile at a specific teres.

Description

Functional Profile of a gene set at specific GO level. Given a vector of genes, this function will return the GO profile at a specific level.

Usage

```
groupGO(gene, OrgDb, keytype = "ENTREZID", ont = "CC", level = 2,
  readable = FALSE)
```

Arguments

gene a vector of entrez gene id.

OrgDb OrgDb

keytype keytype of input gene

ont One of "MF", "BP", and "CC" subontologies.

level Specific GO Level.

readable if readable is TRUE, the gene IDs will mapping to gene symbols.

Value

A groupGOResult instance.

Author(s)

```
Guangchuang Yu http://ygc.name
```

See Also

```
{\tt group GORe sult-class, compare Cluster}
```

Examples

```
data(gcSample)
yy <- groupGO(gcSample[[1]], 'org.Hs.eg.db', ont="BP", level=2)
head(summary(yy))
#plot(yy)</pre>
```

groupGOResult-class 17

groupGOResult-class	Class "groupGOResult" This class represents the result of functional
	Profiles of a set of gene at specific GO level.

Description

Class "groupGOResult" This class represents the result of functional Profiles of a set of gene at specific GO level.

Slots

```
result GO classification result
ontology Ontology
level GO level
organism one of "human", "mouse" and "yeast"
gene Gene IDs
readable logical flag of gene ID in symbol or not.
```

Author(s)

```
Guangchuang Yu http://ygc.name
```

See Also

 ${\tt compareClusterResult\ compareCluster\ groupGO}$

```
GSEA GSEA
```

Description

a universal gene set enrichment analysis tools

Usage

```
GSEA(geneList, exponent = 1, nPerm = 1000, minGSSize = 10,
maxGSSize = 500, pvalueCutoff = 0.05, pAdjustMethod = "BH", TERM2GENE,
TERM2NAME = NA, verbose = TRUE, seed = FALSE, by = "fgsea")
```

Arguments

```
geneList order ranked geneList
exponent weight of each step
nPerm number of permutations
minGSSize minimal size of each geneS
```

minGSSize minimal size of each geneSet for analyzing maxGSSize maximal size of genes annotated for testing

pvalueCutoff pvalue cutoff

18 gseGO

pAdjustMethod p value adjustment method

TERM2GENE user input annotation of TERM TO GENE mapping, a data.frame of 2 column

with term and gene

TERM2NAME user input of TERM TO NAME mapping, a data.frame of 2 column with term

and name

verbose logical seed logical

by one of 'fgsea' or 'DOSE'

Value

gseaResult object

Author(s)

Guangchuang Yu

gseGO gseGO

Description

Gene Set Enrichment Analysis of Gene Ontology

Usage

```
gseGO(geneList, ont = "BP", OrgDb, keyType = "ENTREZID", exponent = 1,
    nPerm = 1000, minGSSize = 10, maxGSSize = 500, pvalueCutoff = 0.05,
    pAdjustMethod = "BH", verbose = TRUE, seed = FALSE, by = "fgsea")
```

Arguments

geneList order ranked geneList

ont one of "BP", "MF", "CC" or "GO"

OrgDb OrgDb

keyTypekeytype of geneexponentweight of each stepnPermpermutation numbers

minGSSize minimal size of each geneSet for analyzing maxGSSize maximal size of genes annotated for testing

pvalueCutoff pvalue Cutoff

pAdjustMethod pvalue adjustment method verbose print message or not

seed logical

by one of 'fgsea' or 'DOSE'

gseKEGG 19

Value

gseaResult object

Author(s)

Yu Guangchuang

gseKEGG gseKEGG

Description

Gene Set Enrichment Analysis of KEGG

Usage

```
gseKEGG(geneList, organism = "hsa", keyType = "kegg", exponent = 1,
   nPerm = 1000, minGSSize = 10, maxGSSize = 500, pvalueCutoff = 0.05,
   pAdjustMethod = "BH", verbose = TRUE, use_internal_data = FALSE,
   seed = FALSE, by = "fgsea")
```

Arguments

geneList order ranked geneList

organism supported organism listed in 'http://www.genome.jp/kegg/catalog/org_list.html'

keyType one of "kegg", 'ncbi-geneid', 'ncib-proteinid' and 'uniprot'

exponent weight of each step nPerm permutation numbers

minGSSize minimal size of each geneSet for analyzing maxGSSize maximal size of genes annotated for testing

 ${\tt pvalueCutoff} \qquad {\tt pvalue} \ {\tt Cutoff}$

pAdjustMethod pvalue adjustment method verbose print message or not

use_internal_data

logical, use KEGG.db or latest online KEGG data

seed logical

by one of 'fgsea' or 'DOSE'

Value

gseaResult object

Author(s)

Yu Guangchuang

20 gseMKEGG

|--|

Description

Gene Set Enrichment Analysis of KEGG Module

Usage

```
gseMKEGG(geneList, organism = "hsa", keyType = "kegg", exponent = 1,
    nPerm = 1000, minGSSize = 10, maxGSSize = 500, pvalueCutoff = 0.05,
    pAdjustMethod = "BH", verbose = TRUE, seed = FALSE, by = "fgsea")
```

Arguments

geneList order ranked geneList

organism supported organism listed in 'http://www.genome.jp/kegg/catalog/org_list.html'

keyType one of "kegg", 'ncbi-geneid', 'ncib-proteinid' and 'uniprot'

exponent weight of each step

nPerm permutation numbers

minGSSize minimal size of each geneSet for analyzing
maxGSSize maximal size of genes annotated for testing

pvalueCutoff pvalue Cutoff

pAdjustMethod pvalue adjustment method

verbose print message or not

seed logical

by one of 'fgsea' or 'DOSE'

Value

gseaResult object

Author(s)

Yu Guangchuang

idType 21

idType

idType

Description

list ID types supported by annoDb

Usage

```
idType(OrgDb = "org.Hs.eg.db")
```

Arguments

OrgDb

annotation db

Value

character vector

Author(s)

Guangchuang Yu

KEGG_path2extid

KEGG_path2extid

Description

query all genes in a KEGG pathway or module

Usage

```
KEGG_path2extid(keggID, species = sub("\\d+$", "", keggID),
keggType = "Path", keyType = "kegg")
```

Arguments

keggID KEGG ID, path or module ID

species species

keggType one of 'Path' or 'Module'

keyType KEGG gene type, one of "ncbi-proteinid", "ncbi-geneid", "uniprot", or "kegg"

Value

extid vector

Author(s)

guangchuang yu

22 merge_result

ko2name

ko2name

Description

convert ko ID to descriptive name

Usage

ko2name(ko)

Arguments

ko

ko ID

Value

data.frame

Author(s)

guangchuang yu

merge_result

merge_result

Description

merge a list of enrichResult objects to compareClusterResult

Usage

```
merge_result(enrichResultList)
```

Arguments

```
enrichResultList
```

a list of enrichResult objects

Value

 $a\ compare Cluster Result\ in stance$

Author(s)

Guangchuang Yu

plot 23

plot plot method

Description

plot method generics

Usage

```
## S4 method for signature 'compareClusterResult,ANY'
plot(x, type = "dot",
   colorBy = "p.adjust", showCategory = 5, by = "geneRatio",
   split = NULL, includeAll = TRUE, font.size = 12, title = "")
```

Arguments

x compareClusterResult object

type one of bar or dot

colorBy one of pvalue or p.adjust

showCategory category numbers

by one of geneRatio, Percentage or count

split ONTOLOGY or NULL

includeAll logical font.size font size title figure title

... Additional argument list

Value

plot

Author(s)

Guangchuang Yu http://ygc.name

plotGOgraph plotGOgraph

Description

```
plot GO graph
```

Usage

```
plotGOgraph(x, firstSigNodes = 10, useInfo = "all", sigForAll = TRUE,
    useFullNames = TRUE, ...)
```

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Arguments

x output of enrichGO or gseGO

firstSigNodes number of significant nodes (retangle nodes in the graph)

useInfo additional info

sigForAll if TRUE the score/p-value of all nodes in the DAG is shown, otherwise only

score will be shown

useFullNames logical

... additional parameter of showSigOfNodes, please refer to topGO

Value

GO DAG graph

Author(s)

Guangchuang Yu

See Also

showSigOfNodes

```
plotting.clusterProfile
```

plotting-clusterProfile

Description

Internal plot function for plotting compareClusterResult

Usage

```
plotting.clusterProfile(clProf.reshape.df, x = ~Cluster, type = "dot",
    colorBy = "p.adjust", by = "geneRatio", title = "", font.size = 12)
```

Arguments

clProf.reshape.df

data frame of compareCluster result

x x variable

type one of dot and bar

colorBy one of pvalue or p.adjust by one of percentage and count

title graph title font.size graph font size

Value

ggplot object

read.gmt 25

Author(s)

Guangchuang Yu http://ygc.name

read.gmt

read.gmt

Description

parse gmt file to a data.frame

Usage

```
read.gmt(gmtfile)
```

Arguments

gmtfile

gmt file

Value

data.frame

Author(s)

Guangchuang Yu

```
search_kegg_organism search_kegg_organism
```

Description

search kegg organism, listed in http://www.genome.jp/kegg/catalog/org_list.html

Usage

```
search_kegg_organism(str, by = "scientific_name", ignore.case = FALSE)
```

Arguments

str string

by one of 'kegg.code', 'scientific_name' and 'common_name'

ignore.case TRUE or FALSE

Value

data.frame

Author(s)

Guangchuang Yu

26 simplify

simplify simplify method

Description

simplify output from enrichGO by removing redundancy of enriched GO terms simplify output from compareCluster by removing redundancy of enriched GO terms

Usage

```
## S4 method for signature 'enrichResult'
simplify(x, cutoff = 0.7, by = "p.adjust",
    select_fun = min, measure = "Wang", semData = NULL)
## S4 method for signature 'compareClusterResult'
simplify(x, cutoff = 0.7, by = "p.adjust",
    select_fun = min, measure = "Wang", semData = NULL)
```

Arguments

x output of enrichGO cutoff similarity cutoff

by feature to select representative term, selected by 'select_fun' function

select_fun function to select feature passed by 'by' parameter

measure method to measure similarity semData GOSemSimDATA object

Value

```
updated enrichResult object
updated compareClusterResult object
```

Author(s)

Guangchuang Yu Guangchuang Yu

References

issue #28 https://github.com/GuangchuangYu/clusterProfiler/issues/28

viewKEGG 27

viewKEGG	viewKEGG function is for visualize KEGG pathways works with en-
VIEWKEGG	richResult object to visualize enriched KEGG pathway

Description

viewKEGG function is for visualize KEGG pathways works with enrichResult object to visualize enriched KEGG pathway

Usage

```
viewKEGG(obj, pathwayID, foldChange, color.low = "green",
  color.high = "red", kegg.native = TRUE, out.suffix = "clusterProfiler")
```

Arguments

obj enrichResult object
pathwayID pathway ID or index
foldChange fold change values
color.low color of low foldChange genes
color.high color of high foldChange genes
kegg.native logical
out.suffix suffix of output file

References

Luo et al. (2013) Pathview: an R/Bioconductor package for pathway-based data integration and visualization. *Bioinformatics* (Oxford, England), 29:14 1830–1831, 2013. ISSN 1367-4803 http://bioinformatics.oxfordjournals.org/content/abstract/29/14/1830.abstract PMID: 23740750

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