# Package 'clusterProfiler'

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Type Package

Title A universal enrichment tool for interpreting omics data

Version 4.4.4

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Description This package supports functional characteristics of both coding and non-coding genomics data for thousands of species with up-to-date gene annotation. It provides a univeral interface for gene functional annotation from a variety of sources and thus can be applied in diverse scenarios. It provides a tidy interface to access, manipulate, and visualize enrichment results to help users achieve efficient data interpretation. Datasets obtained from multiple treatments and time points can be analyzed and compared in a single run, easily revealing functional consensus and differences among distinct conditions.

**Depends** R (>= 3.5.0)

**Imports** AnnotationDbi, downloader, DOSE (>= 3.13.1), dplyr, enrichplot (>= 1.9.3), GO.db, GOSemSim, magrittr, methods, plyr, qvalue, rlang, stats, tidyr, utils, yulab.utils

**Suggests** AnnotationHub, knitr, rmarkdown, org.Hs.eg.db, prettydoc, ReactomePA, testthat

VignetteBuilder knitr

ByteCompile true License Artistic-2.0 Encoding UTF-8

URL https://yulab-smu.top/biomedical-knowledge-mining-book/ (docs),
 https://doi.org/10.1016/j.xinn.2021.100141 (paper)

BugReports https://github.com/GuangchuangYu/clusterProfiler/issues

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## 

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clusterProfiler-package

statistical analysis and visualization of functional profiles for genes and gene clusters

## Description

The package implements methods to analyze and visualize functional profiles of gene and gene clusters.

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 NA or not

## Value

data.frame

## Author(s)

4 browseKEGG

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

## 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

buildGOmap 5

#### 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

## Author(s)

Yu Guangchuang

6 compareCluster

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 = "",
  source_from = NULL,
   ...
)
```

## **Arguments**

a list of entrez gene id. Alternatively, a formula of type Entrez~group or a formula of type Entrez | logFC ~ group for "gseGO", "gseKEGG" and "GSEA".

One of "groupGO", "enrichGO", "enrichKEGG", "enrichDO" or "enrichPathway" . Users can also supply their own function.

data if geneClusters is a formula, the data from which the clusters must be extracted. source\_from If using a custom function in "fun", provide the source package as a string here. Otherwise, the function will be obtained from the global environment.

Other arguments.

#### Value

A clusterProfResult instance.

#### Author(s)

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

#### See Also

```
{\tt compareClusterResult-class, groupGO~enrichGO}
```

#### **Examples**

DataSet 7

```
# plot(xx, type="dot", caption="KEGG Enrichment Comparison")
dotplot(xx)
## formula interface
mydf <- data.frame(Entrez=c('1', '100', '1000', '100101467',</pre>
                             '100127206', '100128071'),
                   logFC = c(1.1, -0.5, 5, 2.5, -3, 3),
                   group = c('A', 'A', 'A', 'B', 'B', 'B'),
                   othergroup = c('good', 'good', 'bad', 'bad', 'good', 'bad'))
xx.formula <- compareCluster(Entrez~group, data=mydf,</pre>
                             fun='groupGO', OrgDb='org.Hs.eg.db')
as.data.frame(xx.formula)
## formula interface with more than one grouping variable
xx.formula.twogroups <- compareCluster(Entrez~group+othergroup, data=mydf,
                                        fun='groupGO', OrgDb='org.Hs.eg.db')
as.data.frame(xx.formula.twogroups)
## End(Not run)
```

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

Datasets DE\_GSE8057 contains differential epxressed genes obtained from GSE8057 dataset

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 8 enrichDAVID

## Value

list

#### 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

enrichDAVID 9

#### 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
)
```

## Arguments

gene input gene idType id type

universe background genes. If missing, the all genes listed in the database (eg TERM2GENE

table) will be used as background.

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

annotation david annotation

pvalueCutoff adjusted pvalue cutoff on enrichment tests to report

 $\verb|pAdjustMethod| one of "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none" | learning the statement of t$ 

qvalueCutoff qvalue cutoff on enrichment tests to report as significant. Tests must pass i)

pvalueCutoff on unadjusted pvalues, ii) pvalueCutoff on adjusted pvalues

and iii) qvalueCutoff on qvalues to be reported.

species species
david.user david user

#### Value

A enrichResult instance

#### Author(s)

10 enricher

ner	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	adjusted pvalue cutoff on enrichment tests to report
pAdjustMethod	one of "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none"
universe	background genes. If missing, the all genes listed in the database (eg TERM2GENE table) will be used as background.
minGSSize	minimal size of genes annotated for testing
maxGSSize	maximal size of genes annotated for testing
qvalueCutoff	qvalue cutoff on enrichment tests to report as significant. Tests must pass i) pvalueCutoff on unadjusted pvalues, ii) pvalueCutoff on adjusted pvalues and iii) qvalueCutoff on qvalues to be reported.
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

## Value

A enrichResult instance

## Author(s)

enrichGO 11

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

minGSSize

maxGSSize

readable

pool

OrgDb	OrgDb
keyType	keytype of input gene
ont	One of "BP", "MF", and "CC" subontologies, or "ALL" for all three.
pvalueCutoff	adjusted pvalue cutoff on enrichment tests to report
pAdjustMethod	one of "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none"
universe	background genes. If missing, the all genes listed in the database (eg TERM2GENE table) will be used as background.
qvalueCutoff	qvalue cutoff on enrichment tests to report as significant. Tests must pass i) pvalueCutoff on unadjusted pvalues, ii) pvalueCutoff on adjusted pvalues and iii) qvalueCutoff on qvalues to be reported.

minimal size of genes annotated by Ontology term for testing.

maximal size of genes annotated for testing

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

whether mapping gene ID to gene Name

a vector of entrez gene id.

12 enrichKEGG

#### Value

An enrichResult instance.

#### Author(s)

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

#### See Also

```
enrichResult-class, compareCluster
```

#### **Examples**

```
## Not run:
    data(geneList, package = "DOSE")
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.

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

enrichKEGG 13

#### **Arguments**

gene a vector of entrez gene id.

organism supported organism listed in 'https://www.genome.jp/kegg/catalog/org\_list.html'

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

pvalueCutoff adjusted pvalue cutoff on enrichment tests to report

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

universe background genes. If missing, the all genes listed in the database (eg TERM2GENE

table) will be used as background.

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

maxGSSize maximal size of genes annotated for testing

qvalueCutoff qvalue cutoff on enrichment tests to report as significant. Tests must pass i)

pvalueCutoff on unadjusted pvalues, ii) pvalueCutoff on adjusted pvalues

and iii) qvalueCutoff on qvalues to be reported.

use\_internal\_data

logical, use KEGG.db or latest online KEGG data

#### Value

A enrichResult instance.

#### Author(s)

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

#### See Also

```
enrichResult-class, compareCluster
```

#### **Examples**

```
## Not run:
    data(geneList, package='DOSE')
    de <- names(geneList)[1:100]
    yy <- enrichKEGG(de, pvalueCutoff=0.01)
    head(yy)
## End(Not run)</pre>
```

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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 'https://www.genome.jp/kegg/catalog/org_list.html'
keyType	one of "kegg", 'ncbi-geneid', 'ncib-proteinid' and 'uniprot'
pvalueCutoff	adjusted pvalue cutoff on enrichment tests to report
pAdjustMethod	one of "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none"
universe	background genes. If missing, the all genes listed in the database (eg TERM2GENE table) will be used as background.
minGSSize	minimal size of genes annotated by Ontology term for testing.
maxGSSize	maximal size of genes annotated for testing
qvalueCutoff	qvalue cutoff on enrichment tests to report as significant. Tests must pass i) pvalueCutoff on unadjusted pvalues, ii) pvalueCutoff on adjusted pvalues and iii) qvalueCutoff on qvalues to be reported.

## Value

A enrichResult instance.

enrichWP 15

## Description

ORA analysis for WikiPathways

## Usage

```
enrichWP(gene, organism, ...)
```

## Arguments

gene a vector of entrez gene id

organism supported organisms, which can be accessed via the get\_wp\_organisms() func-

tion

... additional parameters, see also the parameters supported by the enricher() func-

tion

#### **Details**

This function performs over-representation analysis using WikiPathways

## Value

A enrichResult instance

## Author(s)

Guangchuang Yu

get\_wp\_organisms
get\_wp\_organism

## Description

list supported organism of WikiPathways

#### Usage

```
get_wp_organisms()
```

#### **Details**

This function extracts information from 'https://data.wikipathways.org/current/gmt/' and lists all supported organisms

16 Gff2GeneTable

## Value

supported organism list

## Author(s)

Guangchuang Yu

Gff2GeneTable

Gff2GeneTable

## Description

read GFF file and build gene information table

## Usage

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

## Arguments

gffFile GFF file

compress 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 17

go2ont

go2ont

## Description

convert goid to ontology (BP, CC, MF)

## Usage

go2ont(goid)

## Arguments

goid

a vector of GO IDs

## 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)

18 groupGO

gofilter

gofilter

#### **Description**

filter GO enriched result at specific level

#### Usage

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

## Arguments

X

output from enrichGO or compareCluster

level

GO level

#### Value

updated object

#### Author(s)

Guangchuang Yu

groupG0

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.

## 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.

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

groupGOResult-class 19

#### **Arguments**

gene a vector of entrez gene id.

OrgDb OrgDb

keyType key type 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://guangchuangyu.github.io
```

#### See Also

```
groupGOResult-class, compareCluster
```

## **Examples**

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

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.
```

20 GSEA

#### Author(s)

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

#### See Also

compareClusterResult compareCluster groupGO

GSEA GSEA

## **Description**

a universal gene set enrichment analysis tools

#### Usage

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

#### **Arguments**

geneList order ranked geneList exponent weight of each step minGSSize minimal size of each geneSet for analyzing maximal size of genes annotated for testing maxGSSize This parameter sets the boundary for calculating the p value. eps pvalueCutoff adjusted pvalue cutoff pAdjustMethod p value adjustment method TERM2GENE user input annotation of TERM TO GENE mapping, a data.frame of 2 column with term and gene user input of TERM TO NAME mapping, a data.frame of 2 column with term TERM2NAME and name

gseGO 21

```
verbose logical seed logical by one of 'fgsea' or 'DOSE' ... other parameter
```

#### 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,
  minGSSize = 10,
  maxGSSize = 500,
  eps = 1e-10,
  pvalueCutoff = 0.05,
  pAdjustMethod = "BH",
  verbose = TRUE,
  seed = FALSE,
  by = "fgsea",
  ...
)
```

## Arguments

```
geneList order ranked geneList
ont one of "BP", "MF", and "CC" subontologies, or "ALL" for all three.
OrgDb OrgDb
keyType keytype of gene
```

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exponent weight of each step

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

eps This parameter sets the boundary for calculating the p value.

pvalueCutoff pvalue Cutoff

pAdjustMethod pvalue adjustment method verbose print message or not

seed logical

by one of 'fgsea' or 'DOSE'

... other parameter

#### Value

gseaResult object

#### Author(s)

Yu Guangchuang

gseKEGG gseKEGG

#### **Description**

Gene Set Enrichment Analysis of KEGG

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

gseMKEGG 23

#### **Arguments**

geneList order ranked geneList
organism supported organism listed in 'https://www.genome.jp/kegg/catalog/org\_list.html'
keyType one of "kegg", 'ncbi-geneid', 'ncib-proteinid' and 'uniprot'
exponent weight of each step
minGSSize minimal size of each geneSet for analyzing
maxGSSize maximal size of genes annotated for testing
eps This parameter sets the boundary for calculating the p value.

pvalueCutoff pvalue 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'

... other parameter

#### Value

gseaResult object

#### Author(s)

Yu Guangchuang

gseMKEGG gseMKEGG

#### Description

Gene Set Enrichment Analysis of KEGG Module

```
gseMKEGG(
  geneList,
  organism = "hsa",
  keyType = "kegg",
  exponent = 1,
  minGSSize = 10,
  maxGSSize = 500,
  eps = 1e-10,
  pvalueCutoff = 0.05,
```

24 gseWP

```
pAdjustMethod = "BH",
verbose = TRUE,
seed = FALSE,
by = "fgsea",
...
)
```

## Arguments

geneList order ranked geneList

organism supported organism listed in 'https://www.genome.jp/kegg/catalog/org\_list.html'

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

exponent weight of each step

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

eps This parameter sets the boundary for calculating the p value.

pvalueCutoff pvalue Cutoff

pAdjustMethod pvalue adjustment method

verbose print message or not

seed logical

by one of 'fgsea' or 'DOSE'

... other parameter

#### Value

gseaResult object

#### Author(s)

Yu Guangchuang

gseWP gseWP

#### **Description**

GSEA analysis for WikiPathways

```
gseWP(geneList, organism, ...)
```

idType 25

## Arguments

geneList ranked gene list

organism supported organisms, which can be accessed via the get\_wp\_organisms() func-

tion

... additional parameters, see also the parameters supported by the GSEA() func-

tion

#### **Details**

This function performs GSEA using WikiPathways

#### Value

A gseaResult instance

## Author(s)

Guangchuang Yu

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)

26 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 compareClusterResult instance

#### Author(s)

parse\_gff 27

```
parse_gff
```

parse\_gff

## Description

```
parse GAF files
```

## Usage

```
parse_gff(GafFile, nrows = -1)
```

## Arguments

```
GafFile GAF file nrows a parameter
```

## **Details**

given a GAF file, this function extracts the information from it

## Value

a list with two dataframes

plotGOgraph

plot GO graph

## Description

```
plot GO graph
```

```
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

read.gmt read.gmt

## Description

parse gmt file to a data.frame

## Usage

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

## Arguments

gmtfile gmt file

#### Value

data.frame

#### Author(s)

search\_kegg\_organism 29

```
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,
   use_internal_data = TRUE
)
```

#### **Arguments**

#### Value

data.frame

#### Author(s)

Guangchuang Yu

```
simplify simplify method
```

## Description

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

30 simplify

## Usage

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

uniprot\_get 31

#### Author(s)

Guangchuang Yu Gwang-Jin Kim and Guangchuang Yu Guangchuang Yu

#### References

```
issue #28 https://github.com/GuangchuangYu/clusterProfiler/issues/28
issue #162 https://github.com/GuangchuangYu/clusterProfiler/issues/162
```

uniprot\_get

uniprot\_get

## Description

retreve annotation data from uniprot

#### Usage

```
uniprot_get(taxID)
```

## Arguments

taxID

taxonomy ID

#### Value

gene table data frame

## Author(s)

guangchuang yu

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