

Package ‘clusterProfiler’

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Type Package

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

Version 1.8.0

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Description The package implements methods to analyze and visualize functional profiles (GO and KEGG) of gene and gene clusters.

Depends R (>= 2.10), ggplot2

Imports methods, stats4, DBI, plyr, AnnotationDbi, GO.db, KEGG.db, org.Hs.eg.db, DOSE

Suggests GOSemSim, ReactomePA

biocViews Clustering, GO, Pathways, Visualization, MultipleComparisons, GeneSetEnrichment

Collate 'AllGenerics.R' 'clusterProfiler-package.R' 'compareCluster.R' 'enrichGO.R' 'enrichKEGG.R' 'GFFparser.R' 'groupGO.R' 'utilities.R' 'zzz.R'

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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

This package is designed to compare gene clusters functional profiles.

Details

Package:	clusterProfiler
Type:	Package
Version:	1.0.0
Date:	03-15-2011
biocViews:	GO, Clustering, Visualization
Depends:	AnnotationDbi, GO.db, org.Hs.eg.db, ggplot2, plyr, methods
Suggests:	GOSemSim
License:	Artistic-2.0

Author(s)

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See Also

[compareClusterResult](#), [groupGOResult](#) [enrichResult](#)

`buildGOMap``buildGOMap`

Description

building GO mapping files

Usage

```
buildGOMap(gomap, compress = TRUE)
```

Arguments

gomap	data.frame with two columns names "entrezgene", and "go_accession"
compress	logical, indicate file save in compress or not.

Details

provided by a data.frame of gene and GO directly annotation file this function will building gene to GO and GO to gene mapping, with directly and indirectly annotation.

Value

files save in the the working directory

Author(s)

Yu Guangchuang

`compareCluster`

Compare gene clusters functional profile Given a list of gene set, this function will compute profiles of each gene cluster.

Description

Compare gene clusters functional profile Given a list of gene set, this function will compute profiles of each gene cluster.

Usage

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

Arguments

- geneClusters a list of entrez gene id.
- fun One of "groupGO", "enrichGO", "enrichKEGG", "enrichDO" or "enrichPathway".
- ... Other arguments.

Value

A clusterProfResult instance.

Author(s)

Guangchuang Yu <http://ygc.name>

See Also

[compareClusterResult-class](#), [groupGO](#) [enrichGO](#)

Examples

```
data(gcSample)
xx <- compareCluster(gcSample, fun="enrichKEGG", organism="human", pvalueCutoff=0.05)
#summary(xx)
#plot(xx, type="dot", caption="KEGG Enrichment Comparison")
```

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.

Author(s)

Guangchuang Yu <http://ygc.name>

See Also

[groupGOResult](#) [enrichResult](#) [compareCluster](#)

DataSet	<i>Datasets gcSample contains a sample of gene clusters.</i>
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Description

Datasets gcSample contains a sample of gene clusters.

enrichGO	<i>GO Enrichment Analysis of a gene set. Given a vector of genes, this function will return the enrichment GO categories with FDR control.</i>
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Description

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

Usage

```
enrichGO(gene, organism = "human", ont = "MF",
          pvalueCutoff = 0.05, qvalueCutoff = 0.05,
          readable = FALSE)
```

Arguments

gene	a vector of entrez gene id.
organism	Currently, only "human", "mouse" and "yeast" supported.
ont	One of "MF", "BP", and "CC" subontologies.
pvalueCutoff	Cutoff value of pvalue.
qvalueCutoff	Cutoff value of qvalue.
readable	if readable is TRUE, the gene IDs will mapping to gene symbols.

Value

A enrichResult instance.

Author(s)

Guangchuang Yu <http://ygc.name>

See Also

[enrichResult-class](#), [compareCluster](#)

Examples

```
#data(gcSample)
#yy <- enrichGO(gcSample[[1]], organism="human", ont="BP", pvalueCutoff=0.01)
#head(summary(yy))
#plot(yy)
```

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 = "human", pvalueCutoff = 0.05,
           qvalueCutoff = 0.05, readable = FALSE)
```

Arguments

- | | |
|--------------|---|
| gene | a vector of entrez gene id. |
| organism | Currently, only "human" and "mouse" supported. |
| pvalueCutoff | Cutoff value of pvalue. |
| qvalueCutoff | Cutoff value of qvalue. |
| readable | if readable is TRUE, the gene IDs will mapping to gene symbols. |

Value

A `enrichResult` instance.

Author(s)

Guangchuang Yu <http://ygc.name>

See Also

[enrichResult-class](#), [compareCluster](#)

Examples

```
data(gcSample)
yy = enrichKEGG(gcSample[[5]], pvalueCutoff=0.01)
head(summary(yy))
#plot(yy)
```

<code>getGOLevel</code>	<i>get GOIDs at a specific level</i>
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Description

query GOIDs at a specific level.

Usage

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

Arguments

ont	Ontology
level	GO level

Value

a vector of GOIDs

Author(s)

Guangchuang Yu <http://ygc.name>

<code>Gff2GeneTable</code>	<i>Gff2GeneTable</i>
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Description

read GFF file and build gene information table

Usage

```
Gff2GeneTable(gffFile)
```

Arguments

gffFile	GFF file
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Details

given the GFF file, this function will extract information and save it in working directory

Value

file save.

Author(s)

Yu Guangchuang

groupGO	<i>Functional Profile of a gene set at specific GO level. Given a vector of genes, this function will return the GO profile at specific level.</i>
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Description

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

Usage

```
groupGO(gene, organism = "human", ont = "CC", level = 2,
readable = FALSE)
```

Arguments

- | | |
|----------|---|
| gene | a vector of entrez gene id. |
| organism | Currently, only "human" and "mouse" supported. |
| 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

[groupGOResult-class](#), [compareCluster](#)

Examples

```
data(gcSample)
yy <- groupGO(gcSample[[1]], organism="human", ont="BP", level=2)
head(summary(yy))
#plot(yy)
```

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

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

Author(s)

Guangchuang Yu <http://ygc.name>

See Also

[compareClusterResult](#) [compareCluster](#) [groupGO](#)

plot	<i>plot method</i>
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Description

plot method generics

Arguments

... Additional argument list

Value

plot

Author(s)

Guangchuang Yu <http://ygc.name>

`plotting.clusterProfile`
plotting-clusterProfile

Description

Internal plot function for plotting compareClusterResult

Usage

```
plotting.clusterProfile(clProf.reshape.df, type = "dot",
by = "percentage", title = "", font.size = 12)
```

Arguments

<code>clProf.reshape.df</code>	data frame of compareCluster result
<code>type</code>	one of dot and bar
<code>by</code>	one of percentage and count
<code>title</code>	graph title
<code>font.size</code>	graph font size

Value

ggplot object

Author(s)

Guangchuang Yu <http://ygc.name>

`show` *show method*

Description

show method for compareClusterResult instance
 show method for groupGOResult instance

Arguments

<code>object</code>	A compareClusterResult instance.
<code>object</code>	A groupGOResult instance

Value

message

message

Author(s)

Guangchuang Yu <http://ygc.name>

Guangchuang Yu <http://ygc.name>

summary

summary method

Description

summary method for compareClusterResult instance

Arguments

object A compareClusterResult instance.

Value

A data frame

Author(s)

Guangchuang Yu <http://ygc.name>

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