

Package ‘ReactomePA’

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Type Package

Title Reactome Pathway Analysis

Version 1.8.1

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Description This package provides functions for pathway analysis based on REACTOME pathway database. It will implement enrichment analysis, gene set enrichment analysis and functional modules detection.

Dependes R (>= 2.10)

Imports AnnotationDbi, reactome.db, org.Hs.eg.db, DOSE, igraph,graphite

Suggests clusterProfiler, GOSemSim, ChIPseeker, knitr

VignetteBuilder knitr

License GPL-2

biocViews Pathways, Visualization, Annotation

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ReactomePA-package *Reactome Pathway Analysis*

Description

This package is designed for reactome pathway analysis.

Details

Package: ReactomePA
Type: Package
Version: 0.2.1
Date: 02-09-2012
biocViews: Bioinformatics, Pathway, Visualization
Depends: AnnotationDbi, org.Hs.eg.db, igraph, plyr, methods, stats, qvalue, reactome.db
Suggests: GOSemSim, DOSE, clusterProfiler
License: GPL-2

Author(s)

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See Also

[enrichResult](#)

DataSet *Datasets sample contains a sample of gene IDs.*

Description

Datasets sample contains a sample of gene IDs.

enrichPathway	<i>Pathway Enrichment Analysis of a gene set. Given a vector of genes, this function will return the enriched pathways with FDR control.</i>
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Description

Pathway Enrichment Analysis of a gene set. Given a vector of genes, this function will return the enriched pathways with FDR control.

Usage

```
enrichPathway(gene, organism = "human", pvalueCutoff = 0.05,  
              pAdjustMethod = "BH", qvalueCutoff = 0.2, universe, minGSSize = 5,  
              readable = FALSE)
```

Arguments

gene	a vector of entrez gene id.
organism	one of "human", "rat", "mouse", "celegans", "zebrafish", "fly".
pvalueCutoff	Cutoff value of pvalue.
pAdjustMethod	one of "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none"
qvalueCutoff	Cutoff value of qvalue
universe	background genes
minGSSize	minimal size of genes annotated by Ontology term for testing.
readable	whether mapping gene ID to gene Name

Value

A `enrichResult` instance.

Author(s)

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See Also

[enrichResult-class](#)

Examples

```
gene <- c("11171", "8243", "112464", "2194",  
         "9318", "79026", "1654", "65003",  
         "6240", "3476", "6238", "3836",  
         "4176", "1017", "249")  
yy = enrichPathway(gene, pvalueCutoff=0.05)  
head(summary(yy))  
#plot(yy)
```

 gseAnalyzer

gseAnalyzer

Description

Gene Set Enrichment Analysis of Reactome Pathway

Usage

```
gseAnalyzer(geneList, organism = "human", exponent = 1, nPerm = 1000,
  minGSSize = 10, pvalueCutoff = 0.05, pAdjustMethod = "BH",
  verbose = TRUE)
```

Arguments

geneList	order ranked geneList
organism	organism
exponent	weight of each step
nPerm	permutation numbers
minGSSize	minimal size of each geneSet for analyzing
pvalueCutoff	pvalue Cutoff
pAdjustMethod	pvalue adjustment method
verbose	print message or not

Value

gseaResult object

Author(s)

Yu Guangchuang

viewPathway

viewPathway

Description

view reactome pathway

Usage

```
viewPathway(pathName, organism = "human", readable = TRUE,
  foldChange = NULL, ...)
```

Arguments

pathName	pathway Name
organism	supported organism
readable	logical
foldChange	fold change
...	additional parameter

Details

plotting reactome pathway

Value

plot

Author(s)

Yu Guangchuang

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