# Package 'ReactomePA'

November 14, 2023

```
Type Package
Title Reactome Pathway Analysis
Version 1.46.0
Maintainer Guangchuang Yu <guangchuangyu@gmail.com>
Description This package provides functions for pathway analysis based on
     REACTOME pathway database. It implements enrichment analysis, gene set
     enrichment analysis and several functions for visualization.
Depends R (>= 3.4.0)
Imports AnnotationDbi, DOSE (>= 3.5.1), enrichplot, ggplot2 (>=
     3.3.5), ggraph, reactome.db, igraph, graphite, gson
Suggests BiocStyle, clusterProfiler, knitr, rmarkdown, org.Hs.eg.db,
     prettydoc, testthat
VignetteBuilder knitr
ByteCompile true
License GPL-2
URL https://yulab-smu.top/biomedical-knowledge-mining-book/
BugReports https://github.com/GuangchuangYu/ReactomePA/issues
biocViews Pathways, Visualization, Annotation, MultipleComparison,
     GeneSetEnrichment, Reactome
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```

## **R** topics documented:

2

ReactomePA-packag	ţе.																	2
DataSet																		3
enrichPathway																		3
getALLEG																		4
getDb																		5
gsePathway																		5
gson_Reactome																		
reexports																		7
viewPathway																		7
																		9

ReactomePA-package

Reactome Pathway Analysis

## Description

Index

This package is designed for reactome pathway analysis.

## **Details**

Package: ReactomePA
Type: Package
Version: 1.9.4
Date: 02-09-2012

biocViews: Bioinformatics, Pathway, Visulization

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 3

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

## **Description**

Datasets sample contains a sample of gene IDs.

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

#### 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 = 10,
   maxGSSize = 500,
   readable = FALSE
)
```

## **Arguments**

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

4 getALLEG

## Value

A enrichResult instance.

## Author(s)

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

## 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)</pre>
```

getALLEG

getALLEG

## Description

get all entrezgene ID of a specific organism

## Usage

```
getALLEG(organism)
```

## Arguments

organism

species

## Value

entrez gene ID vector

## Author(s)

Yu Guangchuang

getDb 5

 ${\tt getDb}$ 

getDb

## Description

mapping organism name to annotationDb package name

## Usage

```
getDb(organism)
```

## Arguments

organism

one of supported organism

#### Value

annotationDb name

## Author(s)

Yu Guangchuang

gsePathway

gsePathway

## **Description**

Gene Set Enrichment Analysis of Reactome Pathway

## Usage

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

gson\_Reactome

## **Arguments**

geneList order ranked geneList

organism organism

exponent weight of each step

minGSSize minimal size of each geneSet for analyzing maxGSSize maximal size of each geneSet for analyzing

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

gson\_Reactome gson\_Reactome

## Description

download the latest version of Reactome and stored in a 'GSON' object

## Usage

```
gson_Reactome(organism = "human")
```

## Arguments

```
organism one of "human", "rat", "mouse", "celegans", "yeast", "zebrafish", "fly".
```

#### Value

a 'GSON' object

reexports 7

## **Examples**

```
## Not run:
rec_gson <- gson_Reactome("human")
## End(Not run)</pre>
```

reexports

Objects exported from other packages

## **Description**

These objects are imported from other packages. Follow the links below to see their documentation.

```
DOSE geneID, geneInCategory
enrichplot cnetplot, dotplot, emapplot, gseaplot, heatplot, ridgeplot
```

viewPathway

viewPathway

## **Description**

view reactome pathway

## Usage

```
viewPathway(
  pathName,
  organism = "human",
  readable = TRUE,
  foldChange = NULL,
  keyType = "ENTREZID",
  layout = "kk"
)
```

## **Arguments**

pathName pathway Name organism supported organism

readable logical foldChange

keyType keyType of gene ID (i.e. names of foldChange, if available)

layout graph layout

8 viewPathway

## **Details**

plotting reactome pathway

## Value

plot

## Author(s)

Yu Guangchuang

# **Index**

```
* datasets
    DataSet, 3
* internal
    reexports, 7
* manip
    enrichPathway, 3
* package
    ReactomePA-package, 2
cnetplot, 7
cnetplot (reexports), 7
DataSet, 3
dotplot, 7
dotplot (reexports), 7
emapplot, 7
emapplot (reexports), 7
enrichPathway, 3
enrichResult, 2
geneID, 7
geneID (reexports), 7
geneInCategory, 7
geneInCategory (reexports), 7
getALLEG, 4
getDb, 5
gseaplot, 7
gseaplot (reexports), 7
gsePathway, 5
gson_Reactome, 6
heatplot, 7
heatplot (reexports), 7
ReactomePA (ReactomePA-package), 2
ReactomePA-package, 2
reexports, 7
ridgeplot, 7
ridgeplot (reexports), 7
viewPathway, 7
```