# Package 'DOSE'

March 24, 2022

```
Title Disease Ontology Semantic and Enrichment analysis
Version 3.20.1
Maintainer Guangchuang Yu <guangchuangyu@gmail.com>
Description This package implements five methods proposed by
     Resnik, Schlicker, Jiang, Lin and Wang respectively
     for measuring semantic similarities among DO terms and
     gene products. Enrichment analyses including hypergeometric
     model and gene set enrichment analysis are also implemented
     for discovering disease associations of high-throughput
     biological data.
Depends R (>= 3.5.0)
Imports AnnotationDbi, BiocParallel, DO.db, fgsea, ggplot2, GOSemSim
     (>= 2.0.0), methods, qvalue, reshape2, stats, utils
Suggests prettydoc, clusterProfiler, knitr, rmarkdown, org.Hs.eg.db,
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VignetteBuilder knitr
ByteCompile true
License Artistic-2.0
Encoding UTF-8
URL https://yulab-smu.top/biomedical-knowledge-mining-book/
BugReports https://github.com/GuangchuangYu/DOSE/issues
biocViews Annotation, Visualization, MultipleComparison,
     GeneSetEnrichment, Pathways, Software
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Type Package

**27** 

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DOSE-package

Disease Ontology Semantic and Enrichment analysis Implemented five methods proposed by Resnik, Schlicker, Jiang, Lin and Wang respectively for measuring DO semantic similarities, and hypergeometric test for enrichment analysis.

## **Description**

This package is designed to estimate DO-based semantic similarity measurement and enrichment analysis.

## **Details**

Package: DOSE
Type: Package
Version: 2.3.5
Date: 2-27-2012

biocViews: Bioinformatics, Annotation

Depends:

Imports: methods, AnnotationDbi, DO.db Suggests: clusterProfiler, GOSemSim

License: Artistic-2.0

#### Author(s)

Guangchuang Yu, Li-Gen Wang

Maintainer: Guangchuang Yu <guangchuangyu@gmail.com>

## See Also

enrichResult

clusterSim clusterSim

## **Description**

semantic similarity between two gene clusters

## Usage

```
clusterSim(cluster1, cluster2, measure = "Wang", combine = "BMA")
```

#### **Arguments**

cluster1 a vector of gene IDs

cluster2 another vector of gene IDs

measure One of "Resnik", "Lin", "Rel", "Jiang" and "Wang" methods.

combine One of "max", "avg", "rcmax", "BMA" methods, for combining

#### **Details**

given two gene clusters, this function calculates semantic similarity between them.

#### Value

similarity

#### Author(s)

Yu Guangchuang

#### **Examples**

```
cluster1 <- c("835", "5261","241", "994")
cluster2 <- c("307", "308", "317", "321", "506", "540", "378", "388", "396")
clusterSim(cluster1, cluster2)</pre>
```

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 gene2Symbol gene ID to Symbol keytype Gene ID type readable logical flag of gene ID in symbol or not. .call function call
```

computeIC 5

```
termsim Similarity between term
method method of calculating the similarity between nodes
dr dimension reduction result
```

## Author(s)

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

#### See Also

enrichResult

computeIC

compute information content

## Description

compute information content

## Usage

```
computeIC(ont = "DO", organism = "human")
```

## Arguments

ont "DO" organism "human"

## Author(s)

Guangchuang Yu http://guangchuangyu.github.io

DataSet

Datasets

## Description

Information content and DO term to entrez gene IDs mapping

6 enrichDGN

|--|--|--|

## Description

measuring similarities between two DO term vectors.

## Usage

```
doSim(DOID1, DOID2, measure = "Wang")
```

## Arguments

| DOID1 | DO term vector |
|-------|----------------|
| DOID2 | DO term vector |

measure one of "Wang", "Resnik", "Rel", "Jiang", and "Lin".

#### **Details**

provide two DO term vectors, this function will calculate their similarities.

## Value

score matrix

## Author(s)

 $Guang chuang \ Yu \ \texttt{https://guangchuangyu.github.io}$ 

| enrichDGN | Enrichment                | analysis | based | on | the | DisGeNET | (http://www. |
|-----------|---------------------------|----------|-------|----|-----|----------|--------------|
|           | <pre>disgenet.org/)</pre> |          |       |    |     |          |              |

## Description

given a vector of genes, this function will return the enrichment NCG categories with FDR control

enrichDGNv 7

#### Usage

```
enrichDGN(
   gene,
   pvalueCutoff = 0.05,
   pAdjustMethod = "BH",
   universe,
   minGSSize = 10,
   maxGSSize = 500,
   qvalueCutoff = 0.2,
   readable = FALSE
)
```

#### Arguments

gene a vector of entrez 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 by NCG category for testing

maxGSSize maximal size of each geneSet for analyzing

qvalueCutoff qvalue cutoff

readable whether mapping gene ID to gene Name

#### Value

A enrichResult instance

#### Author(s)

Guangchuang Yu

#### References

Janet et al. (2015) DisGeNET: a discovery platform for the dynamical exploration of human diseases and their genes. *Database* bav028 http://database.oxfordjournals.org/content/2015/bav028.long

enrichDGNv enrichDGN

## Description

Enrichment analysis based on the DisGeNET (http://www.disgenet.org/)

8 enrichDGNv

#### Usage

```
enrichDGNv(
   snp,
   pvalueCutoff = 0.05,
   pAdjustMethod = "BH",
   universe,
   minGSSize = 10,
   maxGSSize = 500,
   qvalueCutoff = 0.2,
   readable = FALSE
)
```

#### **Arguments**

a vector of SNP snp pvalueCutoff pvalue cutoff pAdjustMethod one of "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none" background genes universe minGSSize minimal size of genes annotated by NCG category for testing maxGSSize maximal size of each geneSet for analyzing qvalueCutoff qvalue cutoff readable whether mapping gene ID to gene Name

#### **Details**

given a vector of genes, this function will return the enrichment NCG categories with FDR control

#### Value

A enrichResult instance

## Author(s)

Guangchuang Yu

## References

Janet et al. (2015) DisGeNET: a discovery platform for the dynamical exploration of human diseases and their genes. *Database* bav028 http://database.oxfordjournals.org/content/2015/bav028.long

enrichDO 9

enrichDO DO Enrichment Analysis

## Description

Given a vector of genes, this function will return the enrichment DO categories with FDR control.

## Usage

```
enrichDO(
   gene,
   ont = "DO",
   pvalueCutoff = 0.05,
   pAdjustMethod = "BH",
   universe,
   minGSSize = 10,
   maxGSSize = 500,
   qvalueCutoff = 0.2,
   readable = FALSE
)
```

#### **Arguments**

```
a vector of entrez gene id
gene
ont
                 one of DO or DOLite.
pvalueCutoff
                 pvalue cutoff
pAdjustMethod
                 one of "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none"
universe
                 background genes
minGSSize
                  minimal size of genes annotated by NCG category for testing
maxGSSize
                  maximal size of each geneSet for analyzing
qvalueCutoff
                  qvalue cutoff
readable
                  whether mapping gene ID to gene Name
```

#### Value

A enrichResult instance.

## Author(s)

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

#### See Also

```
enrichResult-class
```

10 enricher\_internal

#### **Examples**

```
data(geneList)
gene = names(geneList)[geneList > 1]
yy = enrichDO(gene, pvalueCutoff=0.05)
summary(yy)
```

enricher\_internal

enrich.internal

## Description

interal method for enrichment analysis

## Usage

```
enricher_internal(
  gene,
  pvalueCutoff,
  pAdjustMethod = "BH",
  universe = NULL,
  minGSSize = 10,
  maxGSSize = 500,
  qvalueCutoff = 0.2,
  USER_DATA
)
```

## **Arguments**

gene a vector of entrez gene id. 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 each geneSet for analyzing

qvalueCutoff cutoff of qvalue
USER\_DATA ontology information

#### **Details**

using the hypergeometric model

## Value

A enrichResult instance.

enrichNCG 11

#### Author(s)

Guangchuang Yu http://guangchuangyu.github.io

enrichNCG enrichNCG

#### **Description**

Enrichment analysis based on the Network of Cancer Genes database (http://ncg.kcl.ac.uk/)

## Usage

```
enrichNCG(
   gene,
   pvalueCutoff = 0.05,
   pAdjustMethod = "BH",
   universe,
   minGSSize = 10,
   maxGSSize = 500,
   qvalueCutoff = 0.2,
   readable = FALSE
)
```

## **Arguments**

```
a vector of entrez gene id
gene
pvalueCutoff
                 pvalue cutoff
pAdjustMethod
                 one of "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none"
universe
                 background genes
minGSSize
                 minimal size of genes annotated by NCG category for testing
maxGSSize
                 maximal size of each geneSet for analyzing
qvalueCutoff
                 qvalue cutoff
readable
                 whether mapping gene ID to gene Name
```

#### **Details**

given a vector of genes, this function will return the enrichment NCG categories with FDR control

#### Value

A enrichResult instance

#### Author(s)

Guangchuang Yu

12 enrichResult-class

enrichResult-class

Class "enrichResult" This class represents the result of enrichment analysis.

## **Description**

Class "enrichResult" This class represents the result of enrichment analysis.

#### **Slots**

```
result enrichment analysis
pvalueCutoff pvalueCutoff
pAdjustMethod pvalue adjust method
qvalueCutoff qvalueCutoff
organism only "human" supported
ontology biological ontology
gene Gene IDs
keytype Gene ID type
universe background gene
gene2Symbol mapping gene to Symbol
geneSets gene sets
readable logical flag of gene ID in symbol or not.
termsim Similarity between term
method method of calculating the similarity between nodes
dr dimension reduction result
```

#### Author(s)

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

#### See Also

enrichD0

EXTID2NAME 13

EXTID2NAME

EXTID2NAME

## Description

mapping gene ID to gene Symbol

## Usage

```
EXTID2NAME(OrgDb, geneID, keytype)
```

## Arguments

OrgDb OrgDb

geneID entrez gene ID

keytype keytype

## Value

gene symbol

## Author(s)

Guangchuang Yu http://guangchuangyu.github.io

gene2D0

convert Gene ID to DO Terms

## Description

provide gene ID, this function will convert to the corresponding DO Terms

## Usage

gene2DO(gene)

## **Arguments**

gene

entrez gene ID

#### Value

DO Terms

## Author(s)

Guangchuang Yu http://guangchuangyu.github.io

14 geneInCategory

geneID

geneID generic

## Description

```
geneID generic
```

## Usage

geneID(x)

## Arguments

Х

enrichResult object

#### Value

'geneID' return the 'geneID' column of the enriched result which can be converted to data.frame via 'as.data.frame'

## **Examples**

```
data(geneList, package="DOSE")
de <- names(geneList)[1:100]
x <- enrichDO(de)
geneID(x)</pre>
```

geneInCategory

geneInCategory generic

## Description

```
geneInCategory generic
```

## Usage

```
geneInCategory(x)
```

## **Arguments**

Χ

enrichResult

#### Value

'geneInCategory' return a list of genes, by spliting the input gene vector to enriched functional categories

geneSim 15

#### **Examples**

```
data(geneList, package="DOSE")
de <- names(geneList)[1:100]
x <- enrichDO(de)
geneInCategory(x)</pre>
```

geneSim

geneSim

## Description

measuring similarities bewteen two gene vectors.

## Usage

```
geneSim(geneID1, geneID2 = NULL, measure = "Wang", combine = "BMA")
```

## Arguments

```
geneID1 entrez gene vector

geneID2 entrez gene vector

measure one of "Wang", "Resnik", "Rel", "Jiang", and "Lin".

combine One of "max", "avg", "rcmax", "BMA" methods, for combining semantic similarity scores of multiple DO terms associated with gene/protein.
```

## **Details**

provide two entrez gene vectors, this function will calculate their similarity.

#### Value

score matrix

## Author(s)

Guangchuang Yu http://ygc.name

GSEA\_internal

gseaResult-class

Class "gseaResult" This class represents the result of GSEA analysis

## **Description**

Class "gseaResult" This class represents the result of GSEA analysis

#### **Slots**

```
result GSEA anaysis
organism organism
setType setType
geneSets geneSets
geneList order rank geneList
keytype ID type of gene
permScores permutation scores
params parameters
gene2Symbol gene ID to Symbol
readable whether convert gene ID to symbol
dr dimension reduction result
```

## Author(s)

Guangchuang Yu https://guangchuangyu.github.io

GSEA\_internal

 $GSEA\_internal$ 

## **Description**

generic function for gene set enrichment analysis

## Usage

```
GSEA_internal(
   geneList,
   exponent,
   minGSSize,
   maxGSSize,
   eps,
   pvalueCutoff,
   pAdjustMethod,
```

gseDGN 17

```
verbose,
seed = FALSE,
USER_DATA,
by = "fgsea",
...
)
```

## **Arguments**

geneList order ranked geneList
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 p value Cutoff

pAdjustMethod p value adjustment method

verbose print message or not

seed set seed inside the function to make result reproducible. FALSE by default.

USER\_DATA annotation data

by one of 'fgsea' or 'DOSE'

... other parameter

## Value

gseaResult object

## Author(s)

Yu Guangchuang

gseDGN

DisGeNET Gene Set Enrichment Analysis

## **Description**

perform gsea analysis

18 gseDO

## Usage

```
gseDGN(
  geneList,
  exponent = 1,
  minGSSize = 10,
  maxGSSize = 500,
  pvalueCutoff = 0.05,
  pAdjustMethod = "BH",
  verbose = TRUE,
  seed = FALSE,
  by = "fgsea",
  ...
)
```

## Arguments

geneList order ranked geneList exponent weight of each step

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

pvalueCutoff pvalue Cutoff

pAdjustMethod p value adjustment method

verbose print message or not

seed logical

by one of 'fgsea' or 'DOSE'

... other parameter

#### Value

gseaResult object

#### Author(s)

Yu Guangchuang

gseD0

DO Gene Set Enrichment Analysis

## Description

perform gsea analysis

gseNCG 19

## Usage

```
gseDO(
  geneList,
  exponent = 1,
  minGSSize = 10,
  maxGSSize = 500,
  pvalueCutoff = 0.05,
  pAdjustMethod = "BH",
  verbose = TRUE,
  seed = FALSE,
  by = "fgsea",
  ...
)
```

## Arguments

geneList order ranked geneList exponent weight of each step

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

pvalueCutoff pvalue Cutoff

 ${\tt pAdjustMethod} \quad p \ value \ adjustment \ method$ 

verbose print message or not

seed logical

by one of 'fgsea' or 'DOSE'

... other parameter

#### Value

gseaResult object

#### Author(s)

Yu Guangchuang

gseNCG

NCG Gene Set Enrichment Analysis

## Description

perform gsea analysis

20 gsfilter

#### Usage

```
gseNCG(
  geneList,
  exponent = 1,
  minGSSize = 10,
  maxGSSize = 500,
  pvalueCutoff = 0.05,
  pAdjustMethod = "BH",
  verbose = TRUE,
  seed = FALSE,
  by = "fgsea",
  ...
)
```

#### **Arguments**

geneList

exponent weight of each step
minGSSize minimal size of each geneSet for analyzing
maxGSSize maximal size of each geneSet for analyzing
pvalueCutoff pvalue Cutoff
pAdjustMethod p value adjustment method
verbose print message or not
seed logical

order ranked geneList

by one of 'fgsea' or 'DOSE'

... other parameter

#### Value

gseaResult object

## Author(s)

Yu Guangchuang

gsfilter gsfilter

## Description

filter enriched result by gene set size or gene count

## Usage

```
gsfilter(x, by = "GSSize", min = NA, max = NA)
```

mclusterSim 21

#### **Arguments**

x instance of enrichResult or compareClusterResult

by one of 'GSSize' or 'Count'

min minimal size
max maximal size

#### Value

update object

#### Author(s)

Guangchuang Yu

mclusterSim mclusterSim

## Description

Pairwise semantic similarity for a list of gene clusters

#### Usage

```
mclusterSim(clusters, measure = "Wang", combine = "BMA")
```

## Arguments

clusters A list of gene clusters

measure one of "Wang", "Resnik", "Rel", "Jiang", and "Lin".

combine One of "max", "avg", "rcmax", "BMA" methods, for combining semantic simi-

larity scores of multiple DO terms associated with gene/protein.

#### Value

similarity matrix

## Author(s)

Yu Guangchuang

## **Examples**

```
cluster1 <- c("835", "5261","241")
cluster2 <- c("578","582")
cluster3 <- c("307", "308", "317")
clusters <- list(a=cluster1, b=cluster2, c=cluster3)
mclusterSim(clusters, measure="Wang")</pre>
```

22 rebuildAnnoData

parse\_ratio

parse\_ratio

## **Description**

parse character ratio to double value, such as 1/5 to 0.2

## Usage

```
parse_ratio(ratio)
```

## **Arguments**

ratio

character vector of ratio to parse

#### Value

A numeric vector (double) of parsed ratio

## Author(s)

Guangchuang Yu

rebuildAnnoData

rebuiding annotation data

## Description

rebuilding entrez and DO mapping datasets

## Usage

```
rebuildAnnoData(file)
```

## Arguments

file

do\_rif.human.txt

## Author(s)

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

setReadable 23

setReadable

setReadable

## Description

mapping geneID to gene Symbol

## Usage

```
setReadable(x, OrgDb, keyType = "auto")
```

## Arguments

x enrichResult Object

OrgDb OrgDb

keyType keyType of gene

#### Value

enrichResult Object

## Author(s)

Yu Guangchuang

show

show method

## Description

```
show method for gseaResult instance show method for enrichResult instance
```

## Usage

```
show(object)
show(object)
```

## Arguments

object

A enrichResult instance.

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## Value

```
message
message
```

## Author(s)

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

simplot

simplot

## **Description**

plotting similarity matrix

## Usage

```
simplot(
    sim,
    xlab = "",
    ylab = "",
    color.low = "white",
    color.high = "red",
    labs = TRUE,
    digits = 2,
    labs.size = 3,
    font.size = 14
)
```

## Arguments

```
\operatorname{sim}
                   similarity matrix
xlab
                   xlab
                   ylab
ylab
color.low
                   color of low value
color.high
                   color of high value
labs
                   logical, add text label or not
                   round digit numbers
digits
labs.size
                   lable size
font.size
                   font size
```

## Value

ggplot object

summary 25

#### Author(s)

Yu Guangchuang

summary

summary method

## Description

```
summary method for gseaResult instance summary method for enrichResult instance
```

## Usage

```
summary(object, ...)
summary(object, ...)
```

## Arguments

```
object A enrichResult instance.
... additional parameter
```

#### Value

A data frame

A data frame

## Author(s)

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

theme\_dose

theme\_dose

## Description

```
ggplot theme of DOSE
```

## Usage

```
theme_dose(font.size = 14)
```

26 theme\_dose

## Arguments

font.size font size

## Value

ggplot theme

## Examples

```
library(ggplot2)
qplot(1:10) + theme_dose()
```

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