Package 'GOSemSim'

May 27, 2019

Version 2.10.0 Maintainer Guangchuang Yu <guangchuangyu@gmail.com> **Description** The semantic comparisons of Gene Ontology (GO) annotations provide quantitative ways to compute similarities between genes and gene groups, and have became important basis for many bioinformatics analysis approaches. GOSemSim is an R package for semantic similarity computation among GO terms, sets of GO terms, gene products and gene clusters. GOSemSim implemented five methods proposed by Resnik, Schlicker, Jiang, Lin and Wang respectively. **Depends** R (>= 3.4.0) LinkingTo Rcpp Imports AnnotationDbi, GO.db, methods, utils Suggests AnnotationHub, BiocManager, clusterProfiler, DOSE, knitr, org.Hs.eg.db, prettydoc, testthat VignetteBuilder knitr ByteCompile true License Artistic-2.0 **Encoding** UTF-8 URL https://guangchuangyu.github.io/software/GOSemSim BugReports https://github.com/YuLab-SMU/GOSemSim/issues biocViews Annotation, GO, Clustering, Pathways, Network, Software RoxygenNote 6.1.1 git_url https://git.bioconductor.org/packages/GOSemSim git_branch RELEASE_3_9 git_last_commit 5db1ecb git_last_commit_date 2019-05-02 **Date/Publication** 2019-05-26 Author Guangchuang Yu [aut, cre],

Type Package

Title GO-terms Semantic Similarity Measures

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GOSer	Sim-package Gene Ontology-based Sematic Similarity Measures	_

Description

Implementation of semantic similarity measures to estimate the functional similarities among Gene Ontology terms and gene products

Details

Quantitative measure of functional similarities among gene products is important for post-genomics study. and widely used in gene function prediction, cluster analysis and pathway modeling. This package is designed to estimate the GO terms' and genes' semantic similarities. Implemented five methods proposed by Resnik, Schlicker, Jiang, Lin and Wang respectively. Support many species, including Anopheles, Arabidopsis, Bovine, Canine, Chicken, Chimp, E coli strain K12 and strain Sakai, Fly, Human, Malaria, Mouse, Pig, Rhesus, Rat, Worm, Xenopus, Yeast, Zebrafish.

Package: GOSemSim Type: Package Version: 2.0.0 Date: 09-11-2012

biocViews: GO, Clustering, Pathways, Anopheles_gambiae, Arabidopsis_thaliana, Bos_taurus, Caenorhabditis_elegans

Depends:

Imports: methods, AnnotationDbi, GO.db

Suggests: clusterProfiler, DOSE

License: Artistic-2.0

Author(s)

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References

Yu et al. (2010) GOSemSim: an R package for measuring semantic similarity among GO terms and gene products *Bioinformatics* (Oxford, England), 26:7 976–978, April 2010. ISSN 1367-4803 http://bioinformatics.oxfordjournals.org/cgi/content/abstract/26/7/976 PMID: 20179076

See Also

 $\verb"goSim mgoSim" geneSim mgeneSim clusterSim mclusterSim"$

clusterSim

Semantic Similarity Between Two Gene Clusters

Description

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

Usage

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

Arguments

cluster1	A set of gene IDs.
cluster2	Another set of gene IDs.
semData	GOSemSimDATA object
measure	One of "Resnik", "Lin", "Rel", "Jiang" and "Wang" methods.
drop	A set of evidence codes based on which certain annotations are dropped. Use NULL to keep all GO annotations.
combine	One of "max", "avg", "rcmax", "BMA" methods, for combining semantic similarity scores of multiple GO terms associated with protein or multiple proteins assiciated with protein cluster.

Value

similarity

References

Yu et al. (2010) GOSemSim: an R package for measuring semantic similarity among GO terms and gene products *Bioinformatics* (Oxford, England), 26:7 976–978, April 2010. ISSN 1367-4803 http://bioinformatics.oxfordjournals.org/cgi/content/abstract/26/7/976 PMID: 20179076

See Also

goSim mgoSim geneSim mgeneSim mclusterSim

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Examples

```
d <- godata('org.Hs.eg.db', ont="MF", computeIC=FALSE)
  cluster1 <- c("835", "5261","241", "994")
cluster2 <- c("307", "308", "317", "321", "506", "540", "378", "388", "396")
clusterSim(cluster1, cluster2, semData=d, measure="Wang")</pre>
```

combineScores

combining similarity matrix to similarity score

Description

Functions for combining similarity matrix to similarity score

Usage

```
combineScores(SimScores, combine)
```

Arguments

SimScores similarity matrix combine combine method

Value

similarity value

Author(s)

Guangchuang Yu http://guangchuangyu.github.io

geneSim

Semantic Similarity Between two Genes

Description

Given two genes, this function will calculate the semantic similarity between them, and return their semantic similarity and the corresponding GO terms

Usage

```
geneSim(gene1, gene2, semData, measure = "Wang", drop = "IEA",
  combine = "BMA")
```

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Arguments

gene1 Entrez gene id.

gene2 Another entrez gene id. semData GOSemSimDATA object

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

drop A set of evidence codes based on which certain annotations are dropped. Use

NULL to keep all GO annotations.

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

ilarity scores of multiple GO terms associated with protein or multiple proteins

assiciated with protein cluster.

Value

list of similarity value and corresponding GO.

References

Yu et al. (2010) GOSemSim: an R package for measuring semantic similarity among GO terms and gene products *Bioinformatics* (Oxford, England), 26:7 976–978, April 2010. ISSN 1367-4803 http://bioinformatics.oxfordjournals.org/cgi/content/abstract/26/7/976 PMID: 20179076

See Also

goSim mgoSim mgeneSim clusterSim mclusterSim

Examples

```
d <- godata('org.Hs.eg.db', ont="MF", computeIC=FALSE)
geneSim("241", "251", semData=d, measure="Wang")</pre>
```

godata godata

Description

prepare GO DATA for measuring semantic similarity

Usage

```
godata(OrgDb = NULL, keytype = "ENTREZID", ont, computeIC = TRUE)
```

Arguments

OrgDb OrgDb object keytype keytype

ont one of 'BP', 'MF', 'CC' computeIC logical, whether computer IC

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Value

GOSemSimDATA object

Author(s)

Guangchuang Yu

GOSemSimDATA-class

Class "GOSemSimDATA" This class stores IC and gene to go mapping for semantic similarity measurement

Description

Class "GOSemSimDATA" This class stores IC and gene to go mapping for semantic similarity measurement

Slots

```
keys gene ID
ont ontology
IC IC data
geneAnno gene to GO mapping
metadata metadata
```

goSim

Semantic Similarity Between Two GO Terms

Description

Given two GO IDs, this function calculates their semantic similarity.

Usage

```
goSim(GOID1, GOID2, semData, measure = "Wang")
```

Arguments

GOID1 GO ID 1. GOID2 GO ID 2.

semData GOSemSimDATA object

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

Value

similarity

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References

Yu et al. (2010) GOSemSim: an R package for measuring semantic similarity among GO terms and gene products *Bioinformatics* (Oxford, England), 26:7 976–978, April 2010. ISSN 1367-4803 http://bioinformatics.oxfordjournals.org/cgi/content/abstract/26/7/976 PMID: 20179076

See Also

mgoSim geneSim mgeneSim clusterSim mclusterSim

Examples

```
d <- godata('org.Hs.eg.db', ont="MF", computeIC=FALSE)
goSim("GO:0004022", "GO:0005515", semData=d, measure="Wang")</pre>
```

go_term_table

Information content of GO terms

Description

These datasets are the information contents of GOterms.

References

Yu et al. (2010) GOSemSim: an R package for measuring semantic similarity among GO terms and gene products *Bioinformatics* (Oxford, England), 26:7 976–978, April 2010. ISSN 1367-4803 http://bioinformatics.oxfordjournals.org/cgi/content/abstract/26/7/976 PMID: 20179076

infoContentMethod

information content based methods

Description

Information Content Based Methods for semantic similarity measuring

Usage

```
infoContentMethod(ID1, ID2, method, godata)
```

Arguments

ID1	Ontology Term
ID2	Ontology Term

method one of "Resnik", "Jiang", "Lin" and "Rel".

godata GOSemSimDATA object

Details

implemented for methods proposed by Resnik, Jiang, Lin and Schlicker.

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Value

semantic similarity score

Author(s)

Guangchuang Yu https://guangchuangyu.github.io

load_OrgDb

load_OrgDb

Description

load OrgDb

Usage

load_OrgDb(OrgDb)

Arguments

OrgDb

OrgDb object or OrgDb name

Value

OrgDb object

Author(s)

Guangchuang Yu

mclusterSim

Pairwise Semantic Similarities for a List of Gene Clusters

Description

Given a list of gene clusters, this function calculates pairwise semantic similarities.

Usage

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

Arguments

clusters A list of gene clusters. semData GOSemSimDATA object

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

drop A set of evidence codes based on which certain annotations are dropped. Use

NULL to keep all GO annotations.

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

ilarity scores of multiple GO terms associated with protein or multiple proteins

assiciated with protein cluster.

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Value

similarity matrix

References

Yu et al. (2010) GOSemSim: an R package for measuring semantic similarity among GO terms and gene products *Bioinformatics* (Oxford, England), 26:7 976–978, April 2010. ISSN 1367-4803 http://bioinformatics.oxfordjournals.org/cgi/content/abstract/26/7/976 PMID: 20179076

See Also

```
goSim mgoSim geneSim mgeneSim clusterSim
```

Examples

```
d <- godata('org.Hs.eg.db', ont="MF", computeIC=FALSE)
cluster1 <- c("835", "5261","241")
cluster2 <- c("578","582")
cluster3 <- c("307", "308", "317")
clusters <- list(a=cluster1, b=cluster2, c=cluster3)
mclusterSim(clusters, semData=d, measure="Wang")</pre>
```

mgeneSim

Pairwise Semantic Similarity for a List of Genes

Description

Given a list of genes, this function calculates pairwise semantic similarities.

Usage

```
mgeneSim(genes, semData, measure = "Wang", drop = "IEA",
  combine = "BMA", verbose = TRUE)
```

Arguments

genes	A list of entrez gene IDs.
semData	GOSemSimDATA object

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

drop A set of evidence codes based on which certain annotations are dropped. Use

NULL to keep all GO annotations.

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

ilarity scores of multiple GO terms associated with protein or multiple proteins

assiciated with protein cluster.

verbose show progress bar or not.

Value

similarity matrix

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References

Yu et al. (2010) GOSemSim: an R package for measuring semantic similarity among GO terms and gene products *Bioinformatics* (Oxford, England), 26:7 976–978, April 2010. ISSN 1367-4803 http://bioinformatics.oxfordjournals.org/cgi/content/abstract/26/7/976 PMID: 20179076

See Also

goSim mgoSim geneSim clusterSim mclusterSim

Examples

```
d <- godata('org.Hs.eg.db', ont="MF", computeIC=FALSE)
mgeneSim(c("835", "5261","241"), semData=d, measure="Wang")</pre>
```

mgoSim

Semantic Similarity Between two GO terms lists

Description

Given two GO term sets, this function will calculate the semantic similarity between them, and return their semantic similarity

Usage

```
mgoSim(GO1, GO2, semData, measure = "Wang", combine = "BMA")
```

Arguments

GO1 A set of go terms.

GO2 Another set of go terms.

semData GOSemSimDATA object

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

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

ilarity scores of multiple GO terms associated with protein or multiple proteins

assiciated with protein cluster.

Value

similarity

References

Yu et al. (2010) GOSemSim: an R package for measuring semantic similarity among GO terms and gene products *Bioinformatics* (Oxford, England), 26:7 976–978, April 2010. ISSN 1367-4803 http://bioinformatics.oxfordjournals.org/cgi/content/abstract/26/7/976 PMID: 20179076

See Also

goSim geneSim mgeneSim clusterSim mclusterSim

termSim 11

Examples

```
d <- godata('org.Hs.eg.db', ont="MF", computeIC=FALSE)
go1 <- c("GO:0004022", "GO:0004024", "GO:0004023")
go2 <- c("GO:0009055", "GO:0020037")
mgoSim("GO:0003824", go2, semData=d, measure="Wang")
mgoSim(go1, go2, semData=d, measure="Wang")</pre>
```

termSim

termSim

Description

measuring similarities between two term vectors.

Usage

```
termSim(t1, t2, semData, method = c("Wang", "Resnik", "Rel", "Jiang",
   "Lin"))
```

Arguments

t1 term vectort2 term vectorsemData GOSemSimDATA object

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

Details

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

Value

score matrix

Author(s)

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

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 $wangMethod_internal$ wangMethod

Description

Method Wang for semantic similarity measuring

Usage

```
wangMethod_internal(ID1, ID2, ont = "BP")
```

Arguments

ID1	Ontology Term
ID2	Ontology Term
ont	Ontology

Value

semantic similarity score

Author(s)

Guangchuang Yu http://ygc.name

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