

Package ‘dcGOR’

September 25, 2014

Type Package

Title Analysis of ontologies and protein domain annotations.

Version 1.0.3

Date 2014-9-25

Author Hai Fang and Julian Gough

Maintainer Hai Fang <hfang@cs.bris.ac.uk>

Depends R (>= 3.1.0), Matrix, igraph, dnet

Imports methods

Suggests foreach, doMC

Description The package is used to analyse domain-centric ontologies and annotations, particularly those in the dcGO database. The dcGO (<http://supfam.org/SUPERFAMILY/dcGO>) is a comprehensive domain-centric database for annotating protein domains using a panel of ontologies including Gene Ontology. With the package, users are expected to analyse and visualise domain-centric ontologies and annotations. Supported analyses include but are not limited to: easy access to a wide range of ontologies and their domain-centric annotations; able to build customised ontologies and annotations; domain-based enrichment analysis and visualisation; construction of a domain (semantic similarity) network according to ontology annotations; significance analysis for estimating a contact (statistical significance) network via Random Walk with Restart; and high-performance parallel computing.

URL <http://supfam.org/dcGOR>

Collate 'ClassMethod-dcGOR.r' 'dcDAGannotate.r' 'dcRDataLoader.r'
'dcEnrichment.r' 'visEnrichment.r' 'dcDAGdomainSim.r'
'dcRWRpipeline.r' 'dcConverter.r' 'dcBuildInfoDataFrame.r' 'dcBuildAnno.r' 'dcBuildOnto.r'

License GPL-2

biocViews Bioinformatics

NeedsCompilation no

Repository CRAN

Date/Publication 2014-09-25 13:41:30

R topics documented:

| | |
|--------------------------------|----|
| AdjData-class | 3 |
| Ancestral_domainome | 4 |
| Anno-class | 5 |
| Anno-method | 7 |
| AnnoData-class | 8 |
| Cnetwork-class | 9 |
| Cnetwork-method | 10 |
| Coutput-class | 11 |
| Coutput-method | 13 |
| dcBuildAnno | 15 |
| dcBuildInfoDataFrame | 16 |
| dcBuildOnto | 17 |
| dcConverter | 18 |
| dcDAGannotate | 20 |
| dcDAGdomainSim | 21 |
| dcEnrichment | 26 |
| dcRDataLoader | 32 |
| dcRWRpipeline | 34 |
| Dnetwork-class | 37 |
| Dnetwork-method | 39 |
| Eoutput-class | 40 |
| Eoutput-method | 42 |
| eTOL | 44 |
| InfoDataFrame-class | 45 |
| InfoDataFrame-method | 47 |
| InterPro | 48 |
| InterPro2GOBP | 49 |
| InterPro2GOCC | 50 |
| InterPro2GOMF | 51 |
| Onto-class | 52 |
| Onto-method | 54 |
| onto.DO | 55 |
| onto.GOBP | 56 |
| onto.GOCC | 56 |
| onto.GOMF | 57 |
| onto.HPMI | 58 |
| onto.HPON | 59 |
| onto.HPPA | 60 |
| onto.MP | 61 |
| Pfam | 61 |
| Pfam2GOBP | 62 |
| Pfam2GOCC | 63 |
| Pfam2GOMF | 64 |
| Rfam | 65 |
| Rfam2GOBP | 66 |
| Rfam2GOCC | 67 |

| | |
|-------------------------|----|
| Rfam2GOMF | 68 |
| SCOP.fa | 69 |
| SCOP.fa2DO | 70 |
| SCOP.fa2GOBP | 71 |
| SCOP.fa2GOCC | 72 |
| SCOP.fa2GOMF | 73 |
| SCOP.fa2HPMI | 74 |
| SCOP.fa2HPON | 75 |
| SCOP.fa2HPPA | 76 |
| SCOP.fa2MP | 77 |
| SCOP.sf | 78 |
| SCOP.sf2DO | 79 |
| SCOP.sf2GOBP | 80 |
| SCOP.sf2GOCC | 81 |
| SCOP.sf2GOMF | 82 |
| SCOP.sf2HPMI | 83 |
| SCOP.sf2HPON | 84 |
| SCOP.sf2HPPA | 85 |
| SCOP.sf2MP | 86 |
| visEnrichment | 87 |

| | |
|--------------|-----------|
| Index | 92 |
|--------------|-----------|

| | |
|---------------|--|
| AdjData-class | <i>Definition for VIRTUAL S4 class AdjData</i> |
|---------------|--|

Description

AdjData is union of other classes: either matrix or dgCMatrix (a sparse matrix in the package Matrix). It is used as a virtual class

Value

Class AdjData

See Also

[Onto-class](#)

Ancestral_domainome *Ancestral superfamily domain repertoires in Eukaryotes*

Description

An object of class "Anno" that contains information about domain repertoires (a complete set of domains: domain-ome) in Eukaryotes (including extant and ancestral genomes). This data is prepared based on 1) SUPERFAMILY database which provides domain and architecture assignments to all completely sequenced genomes including eukaryotic genomes; 2) ancestral domain architecture repertoires inferred by applying Dollo parsimony to eukaryotic part of species tree of life (sTOL), from which ancestral superfamily domain and architecture repertoires at all branching points in eukaryotic evolution are inferred. This allows us to list ancestral domain and architecture repertoires that were present at these points. Based on the observed/inferred domain and architecture repertoires, we also define genome-specific plasticity potential for an individual domain as how many different architectures (or architecture diversity) it can be formed in an extant/ancestral genome. As a result, for each genome, domain repertoires (domainome) are represented as a profile of states on domains, where non-zero entry indicates a domain for which how many different architectures have occurred in the genome.

Usage

```
data(Ancestral_domainome)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of 2019 domains X 875 terms/genomes (including 438 extant genomes and 437 ancestral genomes), with each entry telling how many different architectures a domain has in a genome. Note: zero entry also means that this domain is absent in the genome
- termData: variables describing terms/genomes (i.e. columns in annoData), including extant/ancestral genome information: "left_id" (unique and used as internal id), "right_id" (used in combination with "left_id" to define the post-ordered binary tree structure), "taxon_id" (NCBI taxonomy id, if matched), "genome" (2-letter genome identifiers used in SUPERFAMILY, if being extant), "name" (NCBI taxonomy name, if matched), "rank" (NCBI taxonomy rank, if matched), "branchlength" (branch length in relevance to the parent), and "common_name" (NCBI taxonomy common name, if matched and existed)
- domainData: variables describing domains (i.e. rows in annoData), including information about domains: "sunid" for SCOP id, "level" for SCOP level, "classification" for SCOP classification, "description" for SCOP description

References

Fang et al. (2013) A daily-updated tree of (sequenced) life as a reference for genome research. *Scientific reports*, 3:2015.

Morais et al. (2011) SUPERFAMILY 1.75 including a domain-centric gene ontology method. *Nucleic Acids Res*, 39(Database issue):D427-34.

Andreeva et al. (2008) Data growth and its impact on the SCOP database: new developments. *Nucleic Acids Res*, 36(Database issue):D419-425

Examples

```
data(Ancestral_domainome)
Ancestral_domainome
# retrieve info on terms/genomes
termData(Ancestral_domainome)
# retrieve info on SCOP domains
domainData(Ancestral_domainome)
# retrieve the first 5 rows and columns of data
x <- annoData(Ancestral_domainome)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

Anno-class

Definition for S4 class Anno

Description

Anno has 3 slots: annoData, termData and domainData

Value

Class Anno

Slots

annoData An object of S4 class [AnnoData](#), containing data matrix with the column number equal to nrow(termData) and the row number equal to nrow(domainData).

termData An object of S4 class [InfoDataFrame](#), describing information on columns in annoData.

domainData An object of S4 class [InfoDataFrame](#), describing information on rows in annoData.

Creation

An object of this class can be created via: `new("Anno", annoData, termData, domainData)`

Methods

Class-specific methods:

- `dim()`: retrieve the dimension in the object
- `annoData()`: retrieve the slot 'annoData' in the object
- `termData()`: retrieve the slot 'termData' (as class [InfoDataFrame](#)) in the object

- `domainData()`: retrieve the slot 'domainData' (as class `InfoDataFrame`) in the object
- `tData()`: retrieve `termData` (as `data.frame`) in the object
- `dData()`: retrieve `domainData` (as `data.frame`) in the object
- `termNames()`: retrieve term names (ie, row names of `termData`) in the object
- `domanNames()`: retrieve domain names (ie, row names of `domainData`) in the object

Standard generic methods:

- `str()`: compact display of the content in the object
- `show()`: abbreviated display of the object
- `as(matrix, "Anno")`: convert a matrix to an object of class `Anno`
- `as(dgCMatrix, "Anno")`: convert a `dgCMatrix` (a sparse matrix) to an object of class `Anno`
- `[i, j]`: get the subset of the same class

Access

Ways to access information on this class:

- `showClass("Anno")`: show the class definition
- `showMethods(classes="Anno")`: show the method definition upon this class
- `getSlots("Anno")`: get the name and class of each slot in this class
- `slotNames("Anno")`: get the name of each slot in this class
- `selectMethod(f, signature="Anno")`: retrieve the definition code for the method 'f' defined in this class

See Also

[Anno-method](#)

Examples

```
# create an object of class Anno, only given a matrix
annoData <- matrix(runif(50),nrow=10,ncol=5)
as(annoData, "Anno")

# create an object of class Anno, given a matrix plus information on its columns/rows
# 1) create termData: an object of class InfoDataFrame
data <- data.frame(x=1:5, y=I(LETTERS[1:5]), row.names=paste("Term",
1:5, sep="_"))
termData <- new("InfoDataFrame", data=data)
termData

# 2) create domainData: an object of class InfoDataFrame
data <- data.frame(x=1:10, y=I(LETTERS[1:10]),
row.names=paste("Domain", 1:10, sep="_"))
domainData <- new("InfoDataFrame", data=data)
domainData

# 3) create an object of class Anno
# VERY IMPORTANT: make sure having consistent names between annoData and domainData (and termData)
```

```
annoData <- matrix(runif(50),nrow=10,ncol=5)
rownames(annoData) <- rowNames(domainData)
colnames(annoData) <- rowNames(termData)
x <- new("Anno", annoData=annoData, domainData=domainData,
termData=termData)
x
# 4) look at various methods defined on class Anno
dim(x)
annoData(x)
termData(x)
tData(x)
domainData(x)
dData(x)
termNames(x)
domainNames(x)
# 5) get the subset
x[1:3,1:2]
```

Anno-method

Methods defined for S4 class Anno

Description

Methods defined for class Anno.

Usage

```
## S4 method for signature 'Anno'
dim(x)
```

```
## S4 method for signature 'Anno'
annoData(x)
```

```
## S4 method for signature 'Anno'
termData(x)
```

```
## S4 method for signature 'Anno'
domainData(x)
```

```
## S4 method for signature 'Anno'
tData(object)
```

```
## S4 method for signature 'Anno'
dData(object)
```

```
## S4 method for signature 'Anno'
termNames(object)
```

```
## S4 method for signature 'Anno'
domainNames(object)

## S4 method for signature 'Anno'
show(object)

## S4 method for signature 'Anno,ANY,ANY,ANY'
x[i, j, ..., drop = FALSE]
```

Arguments

| | |
|--------|---|
| x | an object of class Anno |
| object | an object of class Anno |
| i | an index |
| j | an index |
| ... | additional parameters |
| drop | For matrices and arrays. If TRUE the result is coerced to the lowest possible dimension (see the examples). This only works for extracting elements, not for the replacement. See drop for further details. |

See Also

[Anno-class](#)

AnnoData-class

Definition for VIRTUAL S4 class AnnoData

Description

AnnoData is union of other classes: either matrix or dgCMatrix (a sparse matrix in the package Matrix). It is used as a virtual class

Value

Class AnnoData

See Also

[Anno-class](#)

| | |
|----------------|---|
| Cnetwork-class | <i>Definition for S4 class Cnetwork</i> |
|----------------|---|

Description

Cnetwork is an S4 class to store a contact network, such as the one from RWR-based contact between samples/terms by [dcRWRpipeline](#). It has 2 slots: nodeInfo and adjMatrix

Value

Class Cnetwork

Slots

nodeInfo An object of S4 class [InfoDataFrame](#), describing information on nodes/domains.

adjMatrix An object of S4 class [AdjData](#), containing symmetric adjacency data matrix for an indirect domain network

Creation

An object of this class can be created via: `new("Cnetwork", nodeInfo, adjMatrix)`

Methods

Class-specific methods:

- `dim()`: retrieve the dimension in the object
- `adjMatrix()`: retrieve the slot 'adjMatrix' in the object
- `nodeInfo()`: retrieve the slot 'nodeInfo' (as class [InfoDataFrame](#)) in the object
- `nInfo()`: retrieve nodeInfo (as `data.frame`) in the object
- `nodeNames()`: retrieve node/term names (ie, row names of nodeInfo) in the object

Standard generic methods:

- `str()`: compact display of the content in the object
- `show()`: abbreviated display of the object
- `as(matrix, "Cnetwork")`: convert a matrix to an object of class Cnetwork
- `as(dgCMatrix, "Cnetwork")`: convert a `dgCMatrix` (a sparse matrix) to an object of class Cnetwork
- `[i]`: get the subset of the same class

Access

Ways to access information on this class:

- `showClass("Cnetwork")`: show the class definition
- `showMethods(classes="Cnetwork")`: show the method definition upon this class
- `getSlots("Cnetwork")`: get the name and class of each slot in this class
- `slotNames("Cnetwork")`: get the name of each slot in this class
- `selectMethod(f, signature="Cnetwork")`: retrieve the definition code for the method 'f' defined in this class

See Also

[Cnetwork-method](#)

Examples

```
# create an object of class Cnetwork, only given a matrix
adjM <- matrix(runif(25),nrow=5,ncol=5)
as(adjM, "Cnetwork")

# create an object of class Cnetwork, given a matrix plus information on nodes
# 1) create nodeI: an object of class InfoDataFrame
data <- data.frame(id=paste("Domain", 1:5, sep="_"),
  level=rep("SCOP",5), description=I(LETTERS[1:5]),
  row.names=paste("Domain", 1:5, sep="_"))
nodeI <- new("InfoDataFrame", data=data)
nodeI
# 2) create an object of class Cnetwork
# VERY IMPORTANT: make sure having consistent names between nodeInfo and adjMatrix
adjM <- matrix(runif(25),nrow=5,ncol=5)
colnames(adjM) <- rownames(adjM) <- rowNames(nodeI)
x <- new("Cnetwork", adjMatrix=adjM, nodeInfo=nodeI)
x
# 3) look at various methods defined on class Cnetwork
dim(x)
adjMatrix(x)
nodeInfo(x)
nInfo(x)
nodeNames(x)
# 4) get the subset
x[1:2]
```

Cnetwork-method

Methods defined for S4 class Cnetwork

Description

Methods defined for class Cnetwork.

Usage

```
## S4 method for signature 'Cnetwork'
dim(x)

## S4 method for signature 'Cnetwork'
adjMatrix(x)

## S4 method for signature 'Cnetwork'
nodeInfo(x)

## S4 method for signature 'Cnetwork'
nInfo(object)

## S4 method for signature 'Cnetwork'
nodeNames(object)

## S4 method for signature 'Cnetwork'
show(object)

## S4 method for signature 'Cnetwork,ANY,ANY,ANY'
x[i, j, ..., drop = FALSE]
```

Arguments

| | |
|--------|---|
| x | an object of class Cnetwork |
| object | an object of class Cnetwork |
| i | an index |
| j | an index |
| ... | additional parameters |
| drop | For matrices and arrays. If TRUE the result is coerced to the lowest possible dimension (see the examples). This only works for extracting elements, not for the replacement. See drop for further details. |

See Also

[Cnetwork-class](#)

Coutput-class

Definition for S4 class Coutput

Description

Coutput is an S4 class to store output by [dcRWRpipeline](#).

Value

Class Coutput

Slots

`ratio` A symmetrix matrix, containing ratio
`zscore` A symmetrix matrix, containing z-scores
`pvalue` A symmetrix matrix, containing p-values
`adjp` A symmetrix matrix, containing adjusted p-values
`cnetwork` An object of S4 class [Cnetwork](#), storing contact network.

Creation

An object of this class can be created via: `new("Coutput", ratio, zscore, pvalue, adjp, cnetwork)`

Methods

Class-specific methods:

- `ratio()`: retrieve the slot 'ratio' in the object
- `zscore()`: retrieve the slot 'zscore' in the object
- `pvalue()`: retrieve the slot 'pvalue' in the object
- `adjp()`: retrieve the slot 'adjp' in the object
- `cnetwork()`: retrieve the slot 'cnetwork' in the object
- `write()`: write the object into a local file

Standard generic methods:

- `str()`: compact display of the content in the object
- `show()`: abbreviated display of the object

Access

Ways to access information on this class:

- `showClass("Coutput")`: show the class definition
- `showMethods(classes="Coutput")`: show the method definition upon this class
- `getSlots("Coutput")`: get the name and class of each slot in this class
- `slotNames("Coutput")`: get the name of each slot in this class
- `selectMethod(f, signature="Coutput")`: retrieve the definition code for the method 'f' defined in this class

See Also

[Coutput-method](#)

Examples

```
## Not run:
# 1) load onto.GOMF (as 'Onto' object)
g <- dcRDataLoader('onto.GOMF')

# 2) load SCOP superfamilies annotated by GOMF (as 'Anno' object)
Anno <- dcRDataLoader('SCOP.sf2GOMF')

# 3) prepare for ontology appended with annotation information
dag <- dcDAGannotate(g, annotations=Anno, path.mode="shortest_paths",
verbose=TRUE)

# 4) calculate pair-wise semantic similarity between 10 randomly chosen domains
alldomains <- unique(unlist(nInfo(dag)$annotations))
domains <- sample(alldomains,10)
dnetwork <- dcDAGdomainSim(g=dag, domains=domains,
method.domain="BM.average", method.term="Resnik", parallel=FALSE,
verbose=TRUE)
dnetwork

# 5) estimate RWR dating based sample/term relationships
# define sets of seeds as data
# each seed with equal weight (i.e. all non-zero entries are '1')
data <- data.frame(aSeeds=c(1,0,1,0,1), bSeeds=c(0,0,1,0,1))
rownames(data) <- id(dnetwork)[1:5]
# calculate their two contact graph
coutput <- dcRWRpipeline(data=data, g=dnetwork, parallel=FALSE)
coutput

# 6) write into the file 'Coutput.txt' in your local directory
write(coutput, file='Coutput.txt', saveBy="adjp")

# 7) retrieve several slots directly
ratio(coutput)
zscore(coutput)
pvalue(coutput)
adjp(coutput)
cnetwork(coutput)

## End(Not run)
```

Coutput-method

Methods defined for S4 class Coutput

Description

Methods defined for S4 class Coutput.

Usage

```
## S4 method for signature 'Coutput'
show(object)

## S4 method for signature 'Coutput'
ratio(x)

## S4 method for signature 'Coutput'
zscore(x)

## S4 method for signature 'Coutput'
pvalue(x)

## S4 method for signature 'Coutput'
adjp(x)

## S4 method for signature 'Coutput'
cnetwork(x)

## S4 method for signature 'Coutput'
write(x, file = "Coutput.txt", saveBy = c("adjp",
"pvalue", "zscore", "ratio"), verbose = T)
```

Arguments

| | |
|---------|--|
| object | an object of S4 class Coutput. Usually this is an output from dcRWRpipeline |
| x | an object of S4 class Coutput. Usually this is part of the output from dcRWRpipeline |
| saveBy | which statistics will be saved. It can be "pvalue" for p value, "adjp" for adjusted p value, "zscore" for z-score, "ratio" for ratio |
| file | a character specifying a file name written into. By default, it is 'Coutput.txt' |
| verbose | logical to indicate whether the messages will be displayed in the screen. By default, it sets to true for display |

Value

write(x) also returns a symmetrix matrix storing the specific statistics

See Also

[Coutput-class](#)

`dcBuildAnno`*Function to build an object of the S4 class Anno from input files*

Description

`dcBuildAnno` is supposed to build an object of the S4 class `Anno`, given input files. These input files include 1) a file containing domain information, 2) a file containing term information, and 3) a file containing associations between domains and terms.

Usage

```
dcBuildAnno(domain_info.file, term_info.file, association.file,  
output.file = "Anno.RData")
```

Arguments

`domain_info.file`

an input file containing domain information. For example, a file containing InterPro domains (InterPro) can be found in <http://supfam.org/dcGOR/data/InterPro/InterPro.txt>. As seen in this example, the input file must contain the header (in the first row), and entries in the first column intend to be domain ID (and must be unique)

`term_info.file`

an input file containing term information. For example, a file containing Gene Ontology (GO) terms can be found in <http://supfam.org/dcGOR/data/InterPro/GO.txt>. As seen in this example, the input file must contain the header (in the first row) and four columns: 1st column for term ID (must be unique), 2nd column for term name, 3rd column for term namespace, and 4th column for term distance. These four columns must be provided, but the content for the last column can be arbitrary (if it is hard to prepare)

`association.file`

an input file containing associations between domains and terms. For example, a file containing associations between InterPro domains and GO Molecular Function (GOMF) terms can be found in <http://supfam.org/dcGOR/data/InterPro/Domain2GOMF.txt>. As seen in this example, the input file must contain the header (in the first row) and two columns: 1st column for domain ID (corresponding to the first column in `'domain_info.file'`), 2nd column for term ID (corresponding to the first column in `'term_info.file'`). If there are additional columns, these columns will be ignored

`output.file`

an output file used to save the built object as an RData-formatted file. If NULL, this file will be saved into "Anno.RData" in the current working local directory

Value

Any use-specified variable that is given on the right side of the assignment sign `'<-'`, which contains the built `Anno` object. Also, an RData file specified in `"output.file"` is saved in the local directory.

Note

If there are no use-specified variable that is given on the right side of the assignment sign '<-', then no object will be loaded onto the working environment.

See Also

[Anno](#)

Examples

```
## Not run:
# build an "Anno" object that contains SCOP domain superfamilies (sf) annotated by GOBP terms
InterPro2GOMF <-
dcBuildAnno(domain_info.file="http://supfam.org/dcGOR/data/InterPro/InterPro.txt",
term_info.file="http://supfam.org/dcGOR/data/InterPro/GO.txt",
association.file="http://supfam.org/dcGOR/data/InterPro/Domain2GOMF.txt",
output.file="InterPro2GOMF.RData")
InterPro2GOMF

## End(Not run)
```

dcBuildInfoDataFrame *Function to build an object of the S4 class InfoDataframe from an input file*

Description

dcBuildInfoDataFrame is supposed to build an object of of the S4 class [InfoDataFrame](#), given an input file. This input file can, for example, contain the domain information.

Usage

```
dcBuildInfoDataFrame(input.file, output.file = "InfoDataFrame.RData")
```

Arguments

| | |
|-------------|--|
| input.file | an input file used to build the object. For example, a file containing InterPro domains (InterPro) can be found in http://supfam.org/dcGOR/data/InterPro/InterPro.txt . As seen in this example, the input file must contain the header (in the first row), and entries in the first column intend to be domain identities (and must be unique) |
| output.file | an output file used to save the built object as an RData-formatted file. If NULL, this file will be saved into "InfoDataFrame.RData" in the current working local directory |

Value

Any use-specified variable that is given on the right side of the assignment sign '<-', which contains the built dcBuildInfoDataFrame object. Also, an RData file specified in "output.file" is saved in the local directory.

Note

If there are no use-specified variable that is given on the right side of the assignment sign '<-', then no object will be loaded onto the working environment.

See Also

[InfoDataFrame](#)

Examples

```
## Not run:
# build an "InfoDataFrame" object that contains information on InterPro domains (InterPro)
InterPro <-
dcBuildInfoDataFrame(input.file="http://supfam.org/dcGOR/data/InterPro/InterPro.txt",
output.file="InterPro.RData")
InterPro

## End(Not run)
```

dcBuildOnto

Function to build an object of the S4 class Onto from input files

Description

dcBuildOnto is supposed to build an object of the S4 class [Onto](#), given input files. These input files include 1) a file containing term relations, and 2) a file containing term/node information.

Usage

```
dcBuildOnto(relations.file, nodes.file, output.file = "Onto.RData")
```

Arguments

`relations.file` an input file containing term relations (i.e. edges from parent terms to child terms). For example, a file containing relations between GO Molecular Function (GOMF) terms can be found in http://supfam.org/dcGOR/data/onto/igraph_GOMF_edges.txt. As seen in this example, the input file must contain the header (in the first row) and two columns: 1st column for parent term ID, and 2nd column for child term ID

| | |
|-------------|---|
| nodes.file | an input file containing term/node information. For example, a file containing GO Molecular Function (GOMF) terms can be found in http://supfam.org/dcGOR/data/onto/igraph_GOMF_nodes.txt . As seen in this example, the input file must contain the header (in the first row) and five columns: 1st column 'name' for node names (actually term ID; must be unique), 2nd column 'term_id' for term ID, 3rd 'term_name' for term name, 4th column 'term_namespace' for term namespace, and 5th column 'term_distance' for term distance. These five columns must be provided, the content in the first two columns are identical, and the content for the last column can be arbitrary (if it is hard to prepare) |
| output.file | an output file used to save the built object as an RData-formatted file. If NULL, this file will be saved into "Onto.RData" in the current working local directory |

Value

Any use-specified variable that is given on the right side of the assignment sign '<-', which contains the built Onto object. Also, an RData file specified in "output.file" is saved in the local directory.

Note

If there are no use-specified variable that is given on the right side of the assignment sign '<-', then no object will be loaded onto the working environment.

See Also

[Onto](#)

Examples

```
## Not run:
# build an "Onto" object for GO Molecular Function
onto.GOMF <-
dcBuildOnto(relations.file="http://supfam.org/dcGOR/data/onto/igraph_GOMF_edges.txt",
nodes.file="http://supfam.org/dcGOR/data/onto/igraph_GOMF_nodes.txt",
output.file="onto.GOMF.RData")
onto.GOMF

## End(Not run)
```

dcConverter

Function to convert an object between graph classes

Description

dcConverter is supposed to convert an object between classes 'Onto' and 'igraph', or between 'Dnetwork' and 'igraph', or between 'Cnetwork' and 'igraph'.

Usage

```
dcConverter(obj, from = c("Onto", "igraph", "Dnetwork", "Cnetwork"),
to = c("igraph", "Onto", "Dnetwork", "Cnetwork"), verbose = TRUE)
```

Arguments

| | |
|---------|---|
| obj | an object of class "Onto", "igraph", "Dnetwork" or "Cnetwork" |
| from | a character specifying the class converted from. It can be one of "Onto", "igraph", "Dnetwork" and "Dnetwork" |
| to | a character specifying the class converted to. It can be one of "Onto", "igraph", "Dnetwork" and "Dnetwork" |
| verbose | logical to indicate whether the messages will be displayed in the screen. By default, it sets to true for display |

Value

an object of class "Onto", "igraph", "Dnetwork" or "Cnetwork"

Note

Conversion is also supported between classes 'Onto' and 'igraph', or between 'Dnetwork' and 'igraph', or between 'Cnetwork' and 'igraph'

See Also

[dcRDataLoader](#), [Onto-class](#), [Dnetwork-class](#), [Cnetwork-class](#)

Examples

```
## Not run:
# 1) conversion between 'Onto' and 'igraph'
# 1a) load onto.GOMF (as 'Onto' object)
on <- dcRDataLoader('onto.GOMF')
on
# 1b) convert the object from 'Onto' to 'igraph' class
ig <- dcConverter(on, from='Onto', to='igraph')
ig
# 1c) convert the object from 'igraph' to 'Onto' class
dcConverter(ig, from='igraph', to='Onto')

# 2) conversion between 'Dnetwork' and 'igraph'
# 2a) computer a domain semantic network (as 'Dnetwork' object)
g <- dcRDataLoader('onto.GOMF')
Anno <- dcRDataLoader('SCOP.sf2GOMF')
dag <- dcDAGannotate(g, annotations=Anno, path.mode="shortest_paths",
verbose=FALSE)
alldomains <- unique(unlist(nInfo(dag)$annotations))
domains <- sample(alldomains,5) # randomly sample 5 domains
dnetwork <- dcDAGdomainSim(g=dag, domains=domains,
method.domain="BM.average", method.term="Resnik", parallel=FALSE,
```

```

verbose=FALSE)
dnetwork
# 2b) convert the object from 'Dnetwork' to 'igraph' class
ig <- dcConverter(dnetwork, from='Dnetwork', to='igraph')
ig
# 2c) convert the object from 'igraph' to 'Dnetwork' class
dcConverter(ig, from='igraph', to='Dnetwork')

## End(Not run)

```

| | |
|---------------|---|
| dcDAGannotate | <i>Function to generate a subgraph of a direct acyclic graph (DAG) induced by the input annotation data</i> |
|---------------|---|

Description

dcDAGannotate is supposed to produce a subgraph induced by the input annotation data, given a direct acyclic graph (DAG; an ontology). The input is a graph of "igraph" or "Onto" object, a list of the vertices containing annotation data, and the mode defining the paths to the root of DAG. The induced subgraph contains vertices (with annotation data) and their ancestors along with the defined paths to the root of DAG. The annotations at these vertices (including their ancestors) are also updated according to the true-path rule: a domain annotated to a term should also be annotated by its all ancestor terms.

Usage

```

dcDAGannotate(g, annotations, path.mode = c("all_paths",
"shortest_paths",
"all_shortest_paths"), verbose = TRUE)

```

Arguments

| | |
|-------------|---|
| g | an object of class "igraph" or Onto |
| annotations | an object of class Anno , that is, the vertices/nodes for which annotation data are provided |
| path.mode | the mode of paths induced by vertices/nodes with input annotation data. It can be "all_paths" for all possible paths to the root, "shortest_paths" for only one path to the root (for each node in query), "all_shortest_paths" for all shortest paths to the root (i.e. for each node, find all shortest paths with the equal lengths) |
| verbose | logical to indicate whether the messages will be displayed in the screen. By default, it sets to true for display |

Value

- subg: an induced subgraph, an object of class "igraph" or "Onto" (the same as input). In addition to the original attributes to nodes and edges, the return subgraph is also appended by new node attributes: "annotations", which contains a list of domains either as original annotations or inherited annotations; "IC", which stands for information content defined as negative 10-based log-transformed frequency of domains annotated to that term.

Note

For the mode "shortest_paths", the induced subgraph is the most concise, and thus informative for visualisation when there are many nodes in query, while the mode "all_paths" results in the complete subgraph.

See Also

[dcRDataLoader](#), [dcEnrichment](#), [dcDAGdomainSim](#), [dcConverter](#)

Examples

```
## Not run:
# 1) load onto.GOMF (as 'Onto' object)
g <- dcRDataLoader('onto.GOMF')

# 2) load SCOP superfamilies annotated by GOMF (as 'Anno' object)
Anno <- dcRDataLoader('SCOP.sf2GOMF')

# 3) prepare for annotation data
# randomly select 5 terms vertices (and their annotation data)
annotations <- Anno[,sample(1:dim(Anno)[2], 5)]

# 4) obtain the induced subgraph according to the input annotation data
# 4a) based on all possible paths (i.e. the complete subgraph induced)
dcDAGannotate(g, annotations, path.mode="all_paths", verbose=TRUE)
# 4b) based on shortest paths (i.e. the most concise subgraph induced)
dag <- dcDAGannotate(g, annotations, path.mode="shortest_paths",
verbose=TRUE)

# 5) color-code nodes/terms according to the number of annotations
if(class(dag)=='Onto') dag <- dcConverter(dag, from='Onto',
to='igraph')
data <- sapply(V(dag)$annotations, length)
names(data) <- V(dag)$name
dnet::visDAG(g=dag, data=data, node.info="both")

## End(Not run)
```

dcDAGdomainSim

Function to calculate pair-wise semantic similarity between domains based on a direct acyclic graph (DAG) with annotated data

Description

dcDAGdomainSim is supposed to calculate pair-wise semantic similarity between domains based on a direct acyclic graph (DAG) with annotated data. It first calculates semantic similarity between terms and then derives semantic similarity between domains from terms-term semantic similarity. Parallel computing is also supported for Linux or Mac operating systems.

Usage

```
dcDAGdomainSim(g, domains = NULL, method.domain = c("BM.average",
"BM.max",
"BM.complete", "average", "max"), method.term = c("Resnik", "Lin",
"Schlicker", "Jiang", "Pesquita"), force = TRUE, fast = TRUE,
parallel = TRUE, multicores = NULL, verbose = TRUE)
```

Arguments

| | |
|----------------------------|---|
| <code>g</code> | an object of class "igraph" or Onto . It must contain a node attribute called 'annotations' for storing annotation data (see example for howto) |
| <code>domains</code> | the domains between which pair-wise semantic similarity is calculated. If NULL, all domains annotatable in the input dag will be used for calculation, which is very prohibitively expensive! |
| <code>method.domain</code> | the method used for how to derive semantic similarity between domains from semantic similarity between terms. It can be "average" for average similarity between any two terms (one from domain 1, the other from domain 2), "max" for the maximum similarity between any two terms, "BM.average" for best-matching (BM) based average similarity (i.e. for each term of either domain, first calculate maximum similarity to any term in the other domain, then take average of maximum similarity; the final BM-based average similarity is the pre-calculated average between two domains in pair), "BM.max" for BM based maximum similarity (i.e. the same as "BM.average", but the final BM-based maximum similarity is the maximum of the pre-calculated average between two domains in pair), "BM.complete" for BM-based complete-linkage similarity (inspired by complete-linkage concept: the least of any maximum similarity between a term of one domain and a term of the other domain). When comparing BM-based similarity between domains, "BM.average" and "BM.max" are sensitive to the number of terms involved; instead, "BM.complete" is much robust in this aspect. By default, it uses "BM.average". |
| <code>method.term</code> | the method used to measure semantic similarity between terms. It can be "Resnik" for information content (IC) of most informative information ancestor (MICA) (see http://arxiv.org/pdf/cmp-1g/9511007.pdf), "Lin" for $2 * IC$ at MICA divided by the sum of IC at pairs of terms (see http://webdocs.cs.ualberta.ca/~lindek/papers/sim.pdf), "Schlicker" for weighted version of 'Lin' by the $1 - \text{prob}(\text{MICA})$ (see http://www.ncbi.nlm.nih.gov/pubmed/16776819), "Jiang" for $1 - \text{difference between the sum of IC at pairs of terms and } 2 * IC \text{ at MICA}$ (see http://arxiv.org/pdf/cmp-1g/9709008.pdf), "Pesquita" for graph information content similarity related to Tanimoto-Jacard index (ie. summed information content of common ancestors divided by summed information content of all ancestors of term1 and term2 (see http://www.ncbi.nlm.nih.gov/pubmed/18460186)) |
| <code>force</code> | logical to indicate whether the only most specific terms (for each domain) will be used. By default, it sets to true. It is always advisable to use this since it is computationally fast but without compromising accuracy (considering the fact that true-path-rule has been applied when running dcDAGannotate) |

| | |
|------------|---|
| fast | logical to indicate whether a vectorised fast computation is used. By default, it sets to true. It is always advisable to use this vectorised fast computation; since the conventional computation is just used for understanding scripts |
| parallel | logical to indicate whether parallel computation with multicores is used. By default, it sets to true, but not necessarily does so. Partly because parallel backends available will be system-specific (now only Linux or Mac OS). Also, it will depend on whether these two packages "foreach" and "doMC" have been installed. It can be installed via: <code>source("http://bioconductor.org/biocLite.R"); biocLite(c("foreach", "doMC"))</code> . If not yet installed, this option will be disabled |
| multicores | an integer to specify how many cores will be registered as the multicore parallel backend to the 'foreach' package. If NULL, it will use a half of cores available in a user's computer. This option only works when parallel computation is enabled |
| verbose | logical to indicate whether the messages will be displayed in the screen. By default, it sets to true for display |

Value

an object of S4 class [Dnetwork](#). It is a weighted and undirect graph, with following slots:

- nodeInfo: an object of S4 class, describing information on nodes/domains
- adjMatrix: an object of S4 class [AdjData](#), containing symmetric adjacency data matrix for pair-wise semantic similarity between domains

Note

For the mode "shortest_paths", the induced subgraph is the most concise, and thus informative for visualisation when there are many nodes in query, while the mode "all_paths" results in the complete subgraph.

See Also

[dcRDataLoader](#), [dcDAGannotate](#), [dcConverter](#), [Dnetwork-class](#)

Examples

```
## Not run:
# 1) Semantic similarity between SCOP domain superfamilies (sf)
## 1a) load onto.GOMF (as 'Onto' object)
g <- dcRDataLoader('onto.GOMF')
## 1b) load SCOP superfamilies annotated by GOMF (as 'Anno' object)
Anno <- dcRDataLoader('SCOP.sf2GOMF')
## 1c) prepare for ontology appended with annotation information
dag <- dcDAGannotate(g, annotations=Anno, path.mode="shortest_paths",
verbose=FALSE)
## 1d) calculate pair-wise semantic similarity between 8 randomly chosen domains
alldomains <- unique(unlist(nInfo(dag)$annotations))
domains <- sample(alldomains,8)
dnetwork <- dcDAGdomainSim(g=dag, domains=domains,
method.domain="BM.average", method.term="Resnik", parallel=FALSE,
```

```

verbose=TRUE)
dnetwork
## 1e) convert it to an object of class 'igraph'
ig <- dcConverter(dnetwork, from='Dnetwork', to='igraph')
ig
## 1f) visualise the domain network
### extract edge weight (with 2-digit precision)
x <- signif(E(ig)$weight, digits=2)
### rescale into an interval [1,4] as edge width
edge.width <- 1 + (x-min(x))/(max(x)-min(x))*3
### do visualisation
dnet::visNet(g=ig, vertex.shape="sphere", edge.width=edge.width,
edge.label=x, edge.label.cex=0.7)

#####
# 2) Semantic similarity between Pfam domains (Pfam)
## 2a) load onto.GOMF (as 'Onto' object)
g <- dcRDataLoader('onto.GOMF')
## 2b) load Pfam domains annotated by GOMF (as 'Anno' object)
Anno <- dcRDataLoader('Pfam2GOMF')
## 2c) prepare for ontology appended with annotation information
dag <- dcDAGannotate(g, annotations=Anno, path.mode="shortest_paths",
verbose=FALSE)
## 2d) calculate pair-wise semantic similarity between 8 randomly chosen domains
alldomains <- unique(unlist(nInfo(dag)$annotations))
domains <- sample(alldomains,8)
dnetwork <- dcDAGdomainSim(g=dag, domains=domains,
method.domain="BM.average", method.term="Resnik", parallel=FALSE,
verbose=TRUE)
dnetwork
## 2e) convert it to an object of class 'igraph'
ig <- dcConverter(dnetwork, from='Dnetwork', to='igraph')
ig
## 2f) visualise the domain network
### extract edge weight (with 2-digit precision)
x <- signif(E(ig)$weight, digits=2)
### rescale into an interval [1,4] as edge width
edge.width <- 1 + (x-min(x))/(max(x)-min(x))*3
### do visualisation
dnet::visNet(g=ig, vertex.shape="sphere", edge.width=edge.width,
edge.label=x, edge.label.cex=0.7)

#####
# 3) Semantic similarity between InterPro domains (InterPro)
## 3a) load onto.GOMF (as 'Onto' object)
g <- dcRDataLoader('onto.GOMF')
## 3b) load InterPro domains annotated by GOMF (as 'Anno' object)
Anno <- dcRDataLoader('InterPro2GOMF')
## 3c) prepare for ontology appended with annotation information
dag <- dcDAGannotate(g, annotations=Anno, path.mode="shortest_paths",
verbose=FALSE)
## 3d) calculate pair-wise semantic similarity between 8 randomly chosen domains
alldomains <- unique(unlist(nInfo(dag)$annotations))

```



```

domains <- sample(alldomains,8)
dnetwork <- dcDAGdomainSim(g=dag, domains=domains,
method.domain="BM.average", method.term="Resnik", parallel=FALSE,
verbose=TRUE)
dnetwork
## 3e) convert it to an object of class 'igraph'
ig <- dcConverter(dnetwork, from='Dnetwork', to='igraph')
ig
## 3f) visualise the domain network
### extract edge weight (with 2-digit precision)
x <- signif(E(ig)$weight, digits=2)
### rescale into an interval [1,4] as edge width
edge.width <- 1 + (x-min(x))/(max(x)-min(x))*3
### do visualisation
dnet::visNet(g=ig, vertex.shape="sphere", edge.width=edge.width,
edge.label=x, edge.label.cex=0.7)

#####
# 4) Semantic similarity between Rfam RNA families (Rfam)
## 4a) load onto.GOBP (as 'Onto' object)
g <- dcRDataLoader('onto.GOBP')
## 4b) load Rfam families annotated by GOBP (as 'Anno' object)
Anno <- dcRDataLoader('Rfam2GOBP')
## 4c) prepare for ontology with annotation information
dag <- dcDAGannotate(g, annotations=Anno, path.mode="shortest_paths",
verbose=FALSE)
## 4d) calculate pair-wise semantic similarity between 8 randomly chosen RNAs
alldomains <- unique(unlist(nInfo(dag)$annotations))
domains <- sample(alldomains,8)
dnetwork <- dcDAGdomainSim(g=dag, domains=domains,
method.domain="BM.average", method.term="Resnik", parallel=FALSE,
verbose=TRUE)
dnetwork
## 4e) convert it to an object of class 'igraph'
ig <- dcConverter(dnetwork, from='Dnetwork', to='igraph')
ig
## 4f) visualise the domain network
### extract edge weight (with 2-digit precision)
x <- signif(E(ig)$weight, digits=2)
### rescale into an interval [1,4] as edge width
edge.width <- 1 + (x-min(x))/(max(x)-min(x))*3
### do visualisation
dnet::visNet(g=ig, vertex.shape="sphere", edge.width=edge.width,
edge.label=x, edge.label.cex=0.7)

#####
# 5) Advanced usage: customised data for ontology and annotations
# 5a) customise ontology
g <-
dcBuildOnto(relations.file="http://supfam.org/dcGOR/data/onto/igraph_GOMF_edges.txt",
nodes.file="http://supfam.org/dcGOR/data/onto/igraph_GOMF_nodes.txt",
output.file="ontology.RData")
# 5b) customise Anno

```

```

Anno <-
dcBuildAnno(domain_info.file="http://supfam.org/dcGOR/data/InterPro/InterPro.txt",
term_info.file="http://supfam.org/dcGOR/data/InterPro/GO.txt",
association.file="http://supfam.org/dcGOR/data/InterPro/Domain2GOMF.txt",
output.file="annotations.RData")
## 5c) prepare for ontology appended with annotation information
dag <- dcDAGannotate(g, annotations=Anno, path.mode="shortest_paths",
verbose=FALSE)
## 5d) calculate pair-wise semantic similarity between 8 randomly chosen domains
alldomains <- unique(unlist(nInfo(dag)$annotations))
domains <- sample(alldomains,8)
dnetwork <- dcDAGdomainSim(g=dag, domains=domains,
method.domain="BM.average", method.term="Resnik", parallel=FALSE,
verbose=TRUE)
dnetwork
## 5e) convert it to an object of class 'igraph'
ig <- dcConverter(dnetwork, from='Dnetwork', to='igraph')
ig
## 5f) visualise the domain network
### extract edge weight (with 2-digit precision)
x <- signif(E(ig)$weight, digits=2)
### rescale into an interval [1,4] as edge width
edge.width <- 1 + (x-min(x))/(max(x)-min(x))*3
### do visualisation
dnet::visNet(g=ig, vertex.shape="sphere", edge.width=edge.width,
edge.label=x, edge.label.cex=0.7)

## End(Not run)

```

dcEnrichment

Function to conduct ontology enrichment analysis given a group of domains

Description

dcEnrichment is supposed to conduct enrichment analysis for an input group of domains using a specified ontology. It returns an object of S4 class "Eoutput". Enrichment analysis is based on either Fisher's exact test or Hypergeometric test. The test can respect the hierarchy of the ontology. The user can customise the background domains; otherwise, the function will use all annotatable domains as the test background

Usage

```

dcEnrichment(data, background = NULL, domain = c(NA, "SCOP.sf",
"SCOP.fa",
"Pfam", "InterPro", "Rfam"), ontology = c(NA, "GOBP", "GOMF", "GOCC",
"DO",
"HPPA", "HPMI", "HPON", "MP", "EC", "KW", "UP"), sizeRange = c(10,
1000),

```

```

min.overlap = 3, which_distance = NULL, test = c("HypergeoTest",
"FisherTest", "BinomialTest"), p.adjust.method = c("BH", "BY",
"bonferroni",
"holm", "hochberg", "hommel"), ontology.algorithm = c("none", "pc",
"elim",
"lea"), elim.pvalue = 0.01, lea.depth = 2, verbose = T,
domain.RData = NULL, ontology.RData = NULL, annotations.RData = NULL,
RData.location = "http://supfam.org/dcGOR/data")

```

Arguments

| | |
|----------------|--|
| data | an input vector. It contains id for a list of domains, for example, sunids for SCOP domains |
| background | a background vector. It contains id for a list of background domains, for example, sunids for SCOP domains. If NULL, by default all annotatable domains are used as background |
| domain | the domain identity. It can be one of 'SCOP.sf' for SCOP superfamilies, 'SCOP.fa' for SCOP families, 'Pfam' for Pfam domains, 'InterPro' for InterPro domains, 'Rfam' for Rfam RNA families |
| ontology | the ontology identity. It can be "GOBP" for Gene Ontology Biological Process, "GOMF" for Gene Ontology Molecular Function, "GOCC" for Gene Ontology Cellular Component, "DO" for Disease Ontology, "HPPA" for Human Phenotype Phenotypic Abnormality, "HPMI" for Human Phenotype Mode of Inheritance, "HPON" for Human Phenotype ONset and clinical course, "MP" for Mammalian Phenotype, "EC" for Enzyme Commission, "KW" for UniProtKB KeyWords, "UP" for UniProtKB UniPathway. For details on the eligibility for pairs of input domain and ontology, please refer to the online Documentations at http://supfam.org/dcGOR/docs.html |
| sizeRange | the minimum and maximum size of members of each term in consideration. By default, it sets to a minimum of 10 but no more than 1000 |
| min.overlap | the minimum number of overlaps. Only those terms that overlap with input data at least min.overlap (3 domains by default) will be processed |
| which_distance | which distance of terms in the ontology is used to restrict terms in consideration. By default, it sets to 'NULL' to consider all distances |
| test | the statistic test used. It can be "FisherTest" for using fisher's exact test, "HypergeoTest" for using hypergeometric test, or "BinomialTest" for using binomial test. Fisher's exact test is to test the independence between domain group (domains belonging to a group or not) and domain annotation (domains annotated by a term or not), and thus compare sampling to the left part of background (after sampling without replacement). Hypergeometric test is to sample at random (without replacement) from the background containing annotated and non-annotated domains, and thus compare sampling to background. Unlike hypergeometric test, binomial test is to sample at random (with replacement) from the background with the constant probability. In terms of the ease of finding the significance, they are in order: hypergeometric test > binomial test > fisher's exact test. In other words, in terms of the calculated p-value, hypergeometric test < binomial test < fisher's exact test |

| | |
|---------------------------------|--|
| <code>p.adjust.method</code> | the method used to adjust p-values. It can be one of "BH", "BY", "bonferroni", "holm", "hochberg" and "hommel". The first two methods "BH" (widely used) and "BY" control the false discovery rate (FDR: the expected proportion of false discoveries amongst the rejected hypotheses); the last four methods "bonferroni", "holm", "hochberg" and "hommel" are designed to give strong control of the family-wise error rate (FWER). Notes: FDR is a less stringent condition than FWER |
| <code>ontology.algorithm</code> | the algorithm used to account for the hierarchy of the ontology. It can be one of "none", "pc", "elim" and "lea". For details, please see 'Note' |
| <code>elim.pvalue</code> | the parameter only used when "ontology.algorithm" is "elim". It is used to control how to declare a significantly enriched term (and subsequently all domains in this term are eliminated from all its ancestors) |
| <code>lea.depth</code> | the parameter only used when "ontology.algorithm" is "lea". It is used to control how many maximum depth is used to consider the children of a term (and subsequently all domains in these children term are eliminated from the use for the recalculation of the significance at this term) |
| <code>verbose</code> | logical to indicate whether the messages will be displayed in the screen. By default, it sets to TRUE for display |
| <code>domain.RData</code> | a file name for RData-formatted file containing an object of S4 class 'InfoDataFrame' (i.g. domain). By default, it is NULL. It is only needed when the user wants to customise enrichment analysis using their own data |
| <code>ontology.RData</code> | a file name for RData-formatted file containing an object of S4 class 'Onto' (i.g. ontology). By default, it is NULL. It is only needed when the user wants to customise enrichment analysis using their own data |
| <code>annotations.RData</code> | a file name for RData-formatted file containing an object of S4 class 'Anno' (i.g. annotations). By default, it is NULL. It is only needed when the user wants to customise enrichment analysis using their own data |
| <code>RData.location</code> | the characters to tell the location of built-in RData files. By default, it remotely locates at "http://supfam.org/dcGOR/data" or "https://github.com/hfangbristol/dcGOR/data". For the user equipped with fast internet connection, this option can be just left as default. But it is always advisable to download these files locally. Especially when the user needs to run this function many times, there is no need to ask the function to remotely download every time (also it will unnecessarily increase the runtime). For examples, these files (as a whole or part of them) can be first downloaded into your current working directory, and then set this option as: <code>RData.location = "."</code> . If RData to load is already part of package itself, this parameter can be ignored (since this function will try to load it via function data first) |

Value

an object of S4 class `Eoutput`, with following slots:

- `domain`: a character specifying the domain identity

- **ontology**: a character specifying the ontology used
- **term_info**: a matrix of nTerm X 5 containing term information, where nTerm is the number of terms in consideration, and the 5 columns are "term_id" (i.e. "Term ID"), "term_name" (i.e. "Term Name"), "namespace" (i.e. "Term Namespace"), "distance" (i.e. "Term Distance") and "IC" (i.e. "Information Content for the term based on annotation frequency by it")
- **anno**: a list of terms, each storing annotated domain members (also within the background domains). Always, terms are identified by "term_id" and domain members identified by their ids (e.g. sunids for SCOP domains)
- **data**: a vector containing input data in consideration. It is not always the same as the input data as only those mappable and annotatable are retained
- **background**: a vector containing background in consideration. It is not always the same as the input background as only those mappable/annotatable are retained
- **overlap**: a list of terms, each storing domains overlapped between domains annotated by a term and domains in the input data (i.e. the domains of interest). Always, terms are identified by "term_id" and domain members identified by their IDs (e.g. sunids for SCOP domains)
- **zscore**: a vector containing z-scores
- **pvalue**: a vector containing p-values
- **adjp**: a vector containing adjusted p-values. It is the p value but after being adjusted for multiple comparisons

Note

The interpretation of the algorithms used to account for the hierarchy of the ontology is:

- "none": does not consider the ontology hierarchy at all.
- "lea": computers the significance of a term in terms of the significance of its children at the maximum depth (e.g. 2). Precisely, once domains are already annotated to any children terms with a more significance than itself, then all these domains are eliminated from the use for the recalculation of the significance at that term. The final p-values takes the maximum of the original p-value and the recalculated p-value.
- "elim": computers the significance of a term in terms of the significance of its all children. Precisely, once domains are already annotated to a significantly enriched term under the cutoff of e.g. $pvalue < 1e-2$, all these domains are eliminated from the ancestors of that term).
- "pc": requires the significance of a term not only using the whole domains as background but also using domains annotated to all its direct parents/ancestors as background. The final p-value takes the maximum of both p-values in these two calculations.
- "Notes": the order of the number of significant terms is: "none" > "lea" > "elim" > "pc".

See Also

[dcRDataLoader](#), [dcDAGannotate](#), [Eoutput-class](#), [visEnrichment](#), [dcConverter](#)

Examples

```

## Not run:
# 1) Enrichment analysis for SCOP domain superfamilies (sf)
## 1a) load SCOP.sf (as 'InfoDataFrame' object)
SCOP.sf <- dcRDataLoader('SCOP.sf')
### randomly select 50 domains as a list of domains of interest
data <- sample(rowNames(SCOP.sf), 50)
## 1b) perform enrichment analysis, producing an object of S4 class 'Eoutput'
eoutput <- dcEnrichment(data, domain="SCOP.sf", ontology="GOMF")
eoutput
## 1c) view the top 10 significance terms
view(eoutput, top_num=10, sortBy="pvalue", details=TRUE)
## 1d) visualise the top 10 significant terms in the ontology hierarchy
### color-coded according to 10-based negative logarithm of adjusted p-values (adjp)
visEnrichment(eoutput)
## 1e) the same as above but using a customised background
### randomly select 500 domains as background
background <- sample(rowNames(SCOP.sf), 500)
### perform enrichment analysis, producing an object of S4 class 'Eoutput'
eoutput <- dcEnrichment(data, background=background, domain="SCOP.sf",
ontology="GOMF")
eoutput
### view the top 10 significance terms
view(eoutput, top_num=10, sortBy="pvalue", details=TRUE)
### visualise the top 10 significant terms in the ontology hierarchy
### color-coded according to 10-based negative logarithm of adjusted p-values (adjp)
visEnrichment(eoutput)

#####
# 2) Enrichment analysis for Pfam domains (Pfam)
## 2a) load Pfam (as 'InfoDataFrame' object)
Pfam <- dcRDataLoader('Pfam')
### randomly select 100 domains as a list of domains of interest
data <- sample(rowNames(Pfam), 100)
## 2b) perform enrichment analysis, producing an object of S4 class 'Eoutput'
eoutput <- dcEnrichment(data, domain="Pfam", ontology="GOMF")
eoutput
## 2c) view the top 10 significance terms
view(eoutput, top_num=10, sortBy="pvalue", details=TRUE)
## 2d) visualise the top 10 significant terms in the ontology hierarchy
### color-coded according to 10-based negative logarithm of adjusted p-values (adjp)
visEnrichment(eoutput)
## 2e) the same as above but using a customised background
### randomly select 1000 domains as background
background <- sample(rowNames(Pfam), 1000)
### perform enrichment analysis, producing an object of S4 class 'Eoutput'
eoutput <- dcEnrichment(data, background=background, domain="Pfam",
ontology="GOMF")
eoutput
### view the top 10 significance terms
view(eoutput, top_num=10, sortBy="pvalue", details=TRUE)
### visualise the top 10 significant terms in the ontology hierarchy

```

```
### color-coded according to 10-based negative logarithm of adjusted p-values (adjp)
visEnrichment(eoutput)
```

```
#####
# 3) Enrichment analysis for InterPro domains (InterPro)
## 3a) load InterPro (as 'InfoDataFrame' object)
InterPro <- dcRDataLoader('InterPro')
### randomly select 100 domains as a list of domains of interest
data <- sample(rowNames(InterPro), 100)
## 3b) perform enrichment analysis, producing an object of S4 class 'Eoutput'
eoutput <- dcEnrichment(data, domain="InterPro", ontology="GOMF")
eoutput
## 3c) view the top 10 significance terms
view(eoutput, top_num=10, sortBy="pvalue", details=TRUE)
## 3d) visualise the top 10 significant terms in the ontology hierarchy
### color-coded according to 10-based negative logarithm of adjusted p-values (adjp)
visEnrichment(eoutput)
## 3e) the same as above but using a customised background
### randomly select 1000 domains as background
background <- sample(rowNames(InterPro), 1000)
### perform enrichment analysis, producing an object of S4 class 'Eoutput'
eoutput <- dcEnrichment(data, background=background, domain="InterPro",
ontology="GOMF")
eoutput
### view the top 10 significance terms
view(eoutput, top_num=10, sortBy="pvalue", details=TRUE)
### visualise the top 10 significant terms in the ontology hierarchy
### color-coded according to 10-based negative logarithm of adjusted p-values (adjp)
visEnrichment(eoutput)
```

```
#####
# 4) Enrichment analysis for Rfam RNA families (Rfam)
## 4a) load Rfam (as 'InfoDataFrame' object)
Rfam <- dcRDataLoader('Rfam')
### randomly select 100 RNAs as a list of RNAs of interest
data <- sample(rowNames(Rfam), 100)
## 4b) perform enrichment analysis, producing an object of S4 class 'Eoutput'
eoutput <- dcEnrichment(data, domain="Rfam", ontology="GOBP")
eoutput
## 4c) view the top 10 significance terms
view(eoutput, top_num=10, sortBy="pvalue", details=FALSE)
## 4d) visualise the top 10 significant terms in the ontology hierarchy
### color-coded according to 10-based negative logarithm of adjusted p-values (adjp)
visEnrichment(eoutput)
## 4e) the same as above but using a customised background
### randomly select 1000 RNAs as background
background <- sample(rowNames(Rfam), 1000)
### perform enrichment analysis, producing an object of S4 class 'Eoutput'
eoutput <- dcEnrichment(data, background=background, domain="Rfam",
ontology="GOBP")
eoutput
### view the top 10 significance terms
view(eoutput, top_num=10, sortBy="pvalue", details=FALSE)
```

```

### visualise the top 10 significant terms in the ontology hierarchy
### color-coded according to 10-based negative logarithm of adjusted p-values (adjp)
visEnrichment(eoutput)

#####
# 5) Advanced usage: customised data for domain, ontology and annotations
# 5a) create domain, ontology and annotations
## for domain
domain <-
dcBuildInfoDataFrame(input.file="http://supfam.org/dcGOR/data/InterPro/InterPro.txt",
output.file="domain.RData")
## for ontology
dcBuildOnto(relations.file="http://supfam.org/dcGOR/data/onto/igraph_GOMF_edges.txt",
nodes.file="http://supfam.org/dcGOR/data/onto/igraph_GOMF_nodes.txt",
output.file="ontology.RData")
## for annotations
dcBuildAnno(domain_info.file="http://supfam.org/dcGOR/data/InterPro/InterPro.txt",
term_info.file="http://supfam.org/dcGOR/data/InterPro/GO.txt",
association.file="http://supfam.org/dcGOR/data/InterPro/Domain2GOMF.txt",
output.file="annotations.RData")
## 5b) prepare data and background
### randomly select 100 domains as a list of domains of interest
data <- sample(rowNames(domain), 100)
### randomly select 1000 domains as background
background <- sample(rowNames(domain), 1000)
## 5c) perform enrichment analysis, producing an object of S4 class 'Eoutput'
eoutput <- dcEnrichment(data, background=background,
domain.RData='domain.RData', ontology.RData='ontology.RData',
annotations.RData='annotations.RData')
eoutput
## 5d) view the top 10 significance terms
view(eoutput, top_num=10, sortBy="pvalue", details=TRUE)
### visualise the top 10 significant terms in the ontology hierarchy
### color-coded according to 10-based negative logarithm of adjusted p-values (adjp)
visEnrichment(eoutput)

## End(Not run)

```

dcRDataLoader

Function to load dcGOR built-in RData

Description

dcRDataLoader is supposed to load RData that are used by package dcGOR.

Usage

```

dcRDataLoader(RData = c(NA, "SCOP.sf", "SCOP.fa", "Pfam", "InterPro",
"Rfam",
"onto.GOBP", "onto.GOMF", "onto.GOCC", "onto.DO", "onto.HPPA",

```



```

"onto.HPMI",
"onto.HPON", "onto.MP", "onto.EC", "onto.KW", "onto.UP",
"SCOP.sf2GOBP",
"SCOP.sf2GOMF", "SCOP.sf2GOCC", "SCOP.sf2DO", "SCOP.sf2HPPA",
"SCOP.sf2HPMI",
"SCOP.sf2HPON", "SCOP.sf2MP", "SCOP.sf2EC", "SCOP.sf2KW", "SCOP.sf2UP",
"SCOP.fa2GOBP", "SCOP.fa2GOMF", "SCOP.fa2GOCC", "SCOP.fa2DO",
"SCOP.fa2HPPA",
"SCOP.fa2HPMI", "SCOP.fa2HPON", "SCOP.fa2MP", "SCOP.fa2EC",
"SCOP.fa2KW",
"SCOP.fa2UP", "Pfam2GOBP", "Pfam2GOMF", "Pfam2GOCC", "InterPro2GOBP",
"InterPro2GOMF", "InterPro2GOCC", "Rfam2GOBP", "Rfam2GOMF",
"Rfam2GOCC",
"Ancestral_domainome", "eTOL"), domain = c(NA, "SCOP.sf", "SCOP.fa",
"Pfam",
"InterPro", "Rfam"), ontology = c(NA, "GOBP", "GOMF", "GOCC", "DO",
"HPPA",
"HPMI", "HPON", "MP", "EC", "KW", "UP"), verbose = T,
RData.location = "http://supfam.org/dcGOR/data")

```

Arguments

| | |
|----------------|--|
| RData | which built-in RData to load. If NOT NA, this RData will be always loaded. It can be: domains/RNAs (including 'SCOP.sf', 'SCOP.fa', 'Pfam', 'InterPro', 'Rfam'), ontologies (including 'onto.GOBP', 'onto.GOMF', 'onto.GOCC', 'onto.DO', 'onto.HPPA', 'onto.HPMI', 'onto.HPON', 'onto.MP', 'onto.EC', 'onto.KW', 'onto.UP'), annotations (including 'SCOP.sf2GOBP', 'SCOP.sf2GOMF', 'SCOP.sf2GOCC', 'SCOP.sf2DO', 'SCOP.sf2HPPA', 'SCOP.sf2HPMI', 'SCOP.sf2HPON', 'SCOP.sf2MP', 'SCOP.sf2EC', 'SCOP.sf2KW', 'SCOP.sf2UP', 'SCOP.fa2GOBP', 'SCOP.fa2GOMF', 'SCOP.fa2GOCC', 'SCOP.fa2DO', 'SCOP.fa2HPPA', 'SCOP.fa2HPMI', 'SCOP.fa2HPON', 'SCOP.fa2MP', 'SCOP.fa2EC', 'SCOP.fa2KW', 'SCOP.fa2UP', 'Pfam2GOBP', 'Pfam2GOMF', 'Pfam2GOCC', 'InterPro2GOBP', 'InterPro2GOMF', 'InterPro2GOCC', 'Rfam2GOBP', 'Rfam2GOMF', 'Rfam2GOCC'), and domainome in eukaryotic genomes (including 'Ancestral_domainome', 'eTOL'). On the meanings, please refer to the Documentations |
| domain | domain part of annotation RData to load. When RData is NA and this plus next are NOT NA, then this plus next one are used to specify which annotation RData to load. In addition to NA, it can also be: 'SCOP.sf', 'SCOP.fa', 'Pfam' and 'InterPro' |
| ontology | ontology part of annotation RData to load. This only works together with the previous 'domain' parameter. In addition to NA, it can also be: 'GOBP', 'GOMF', 'GOCC', 'DO', 'HPPA', 'HPMI', 'HPON', 'MP', 'EC', 'KW', 'UP' |
| verbose | logical to indicate whether the messages will be displayed in the screen. By default, it sets to TRUE for display |
| RData.location | the characters to tell the location of built-in RData files. By default, it remotely locates at "http://supfam.org/dcGOR/data" or "https://github.com/hfangbristol/dcGOR/data". For the user equipped with fast internet connection, this option can be just left as default. But it is always advisable to download these |

files locally. Especially when the user needs to run this function many times, there is no need to ask the function to remotely download every time (also it will unnecessarily increase the runtime). For examples, these files (as a whole or part of them) can be first downloaded into your current working directory, and then set this option as: `RData.location = "."`. If RData to load is already part of package itself, this parameter can be ignored (since this function will try to load it via function data first)

Value

any use-specified variable that is given on the right side of the assignment sign '`<-`', which contains the loaded RData.

Note

If there are no use-specified variable that is given on the right side of the assignment sign '`<-`', then no RData will be loaded onto the working environment.

See Also

[dcEnrichment](#)

Examples

```
# Always, load from specified RData directly
SCOP.sf <- dcRDataLoader(RData='SCOP.sf')
Pfam <- dcRDataLoader(RData='Pfam')
InterPro <- dcRDataLoader(RData='InterPro')
Rfam <- dcRDataLoader(RData='Rfam')
onto.GOMF <- dcRDataLoader(RData='onto.GOMF')
# But for annotation data, there are two ways to do so:
# 1) in a direct way
SCOP.sf2GOMF <- dcRDataLoader(RData='SCOP.sf2GOMF')
# 2) in an indirect way: specify both domain and ontology
SCOP.sf2GOMF <- dcRDataLoader(domain='SCOP.sf', ontology='GOMF')
```

dcRWRpipeline

Function to setup a pipeline to estimate RWR-based contact strength between samples from an input domain-sample data matrix and an input graph

Description

dcRWRpipeline is supposed to estimate sample relationships (ie. contact strength between samples) from an input domain-sample matrix and an input graph (such as a domain-domain semantic network). The pipeline includes: 1) random walk restart (RWR) of the input graph using the input matrix as seeds; 2) calculation of contact strength (inner products of RWR-smoothed columns of input matrix); 3) estimation of the contact significance by a randomisation procedure. It supports

two methods how to use RWR: 'direct' for directly applying RWR in the given seeds; 'indirectly' for first pre-computing affinity matrix of the input graph, and then deriving the affinity score. Parallel computing is also supported for Linux or Mac operating systems.

Usage

```
dcRWRpipeline(data, g, method = c("indirect", "direct"),
normalise = c("laplacian", "row", "column", "none"), restart = 0.75,
normalise.affinity.matrix = c("none", "quantile"),
permutation = c("random", "degree"), num.permutation = 100,
p.adjust.method = c("BH", "BY", "bonferroni", "holm", "hochberg",
"hommel"),
adjp.cutoff = 0.05, parallel = TRUE, multicores = NULL, verbose = T)
```

Arguments

| | |
|---------------------------|--|
| data | an input domain-sample data matrix used for seeds. Each value in input domain-sample matrix does not necessarily have to be binary (non-zeros will be used as a weight, but should be non-negative for easy interpretation). |
| g | an object of class "igraph" or Dnetwork |
| method | the method used to calculate RWR. It can be 'direct' for directly applying RWR, 'indirect' for indirectly applying RWR (first pre-compute affinity matrix and then derive the affinity score) |
| normalise | the way to normalise the adjacency matrix of the input graph. It can be 'laplacian' for laplacian normalisation, 'row' for row-wise normalisation, 'column' for column-wise normalisation, or 'none' |
| restart | the restart probability used for RWR. The restart probability takes the value from 0 to 1, controlling the range from the starting nodes/seeds that the walker will explore. The higher the value, the more likely the walker is to visit the nodes centered on the starting nodes. At the extreme when the restart probability is zero, the walker moves freely to the neighbors at each step without restarting from seeds, i.e., following a random walk (RW) |
| normalise.affinity.matrix | the way to normalise the output affinity matrix. It can be 'none' for no normalisation, 'quantile' for quantile normalisation to ensure that columns (if multiple) of the output affinity matrix have the same quantiles |
| permutation | how to do permutation. It can be 'degree' for degree-preserving permutation, 'random' for permutation in random |
| num.permutation | the number of permutations used to for generating the distribution of contact strength under randomisation |
| p.adjust.method | the method used to adjust p-values. It can be one of "BH", "BY", "bonferroni", "holm", "hochberg" and "hommel". The first two methods "BH" (widely used) and "BY" control the false discovery rate (FDR: the expected proportion of false discoveries amongst the rejected hypotheses); the last four methods "bonferroni", "holm", "hochberg" and "hommel" are designed to give strong control of |

| | |
|-------------|---|
| | the family-wise error rate (FWER). Notes: FDR is a less stringent condition than FWER |
| adjp.cutoff | the cutoff of adjusted pvalue to construct the contact graph |
| parallel | logical to indicate whether parallel computation with multicores is used. By default, it sets to true, but not necessarily does so. Partly because parallel backends available will be system-specific (now only Linux or Mac OS). Also, it will depend on whether these two packages "foreach" and "doMC" have been installed. It can be installed via: <code>source("http://bioconductor.org/biocLite.R"); biocLite(c("foreach", "doMC"))</code> . If not yet installed, this option will be disabled |
| multicores | an integer to specify how many cores will be registered as the multicore parallel backend to the 'foreach' package. If NULL, it will use a half of cores available in a user's computer. This option only works when parallel computation is enabled |
| verbose | logical to indicate whether the messages will be displayed in the screen. By default, it sets to true for display |

Value

an object of class "iContact", a list with following components:

- ratio: a symmetric matrix storing ratio (the observed against the expected) between pairwise samples
- zscore: a symmetric matrix storing zscore between pairwise samples
- pval: a symmetric matrix storing pvalue between pairwise samples
- adjpval: a symmetric matrix storing adjusted pvalue between pairwise samples
- icontact: the constructed contact graph (as an 'igraph' object) under the cutoff of adjusted value
- Amatrix: a pre-computed affinity matrix when using 'indirect' method; NULL otherwise
- call: the call that produced this result

Note

The choice of which method to use RWR depends on the number of seed sets and the number of permutations for statistical test. If the total product of both numbers are huge, it is better to use 'indirect' method (for a single run).

See Also

[dcRDataLoader](#), [dcDAGannotate](#), [dcDAGdomainSim](#), [dcConverter](#)

Examples

```
## Not run:
# 1) load onto.GOMF (as 'Onto' object)
g <- dcRDataLoader('onto.GOMF')

# 2) load SCOP superfamilies annotated by GOMF (as 'Anno' object)
```

```

Anno <- dcRDataLoader('SCOP.sf2GOMF')

# 3) prepare for ontology appended with annotation information
dag <- dcDAGannotate(g, annotations=Anno, path.mode="shortest_paths",
verbose=TRUE)

# 4) calculate pair-wise semantic similarity between 10 randomly chosen domains
alldomains <- unique(unlist(nInfo(dag)$annotations))
domains <- sample(alldomains,10)
dnetwork <- dcDAGdomainSim(g=dag, domains=domains,
method.domain="BM.average", method.term="Resnik", parallel=FALSE,
verbose=TRUE)
dnetwork

# 5) estimate RWR dating based sample/term relationships
# define sets of seeds as data
# each seed with equal weight (i.e. all non-zero entries are '1')
data <- data.frame(aSeeds=c(1,0,1,0,1), bSeeds=c(0,0,1,0,1))
rownames(data) <- id(dnetwork)[1:5]
# calculate their two contact graph
output <- dcRWRpipeline(data=data, g=dnetwork, parallel=FALSE)
output

## End(Not run)

```

Dnetwork-class

Definition for S4 class Dnetwork

Description

Dnetwork is an S4 class to store a domain network, such as the one from semantic similarity between pairs of domains by [dcDAGdomainSim](#). It has 2 slots: nodeInfo and adjMatrix

Value

Class Dnetwork

Slots

nodeInfo An object of S4 class [InfoDataFrame](#), describing information on nodes/domains.

adjMatrix An object of S4 class [AdjData](#), containing symmetric adjacency data matrix for an indirect domain network

Creation

An object of this class can be created via: `new("Dnetwork", nodeInfo, adjMatrix)`

Methods

Class-specific methods:

- `dim()`: retrieve the dimension in the object
- `adjMatrix()`: retrieve the slot 'adjMatrix' in the object
- `nodeInfo()`: retrieve the slot 'nodeInfo' (as class `InfoDataFrame`) in the object
- `nInfo()`: retrieve nodeInfo (as `data.frame`) in the object
- `nodeNameNames()`: retrieve node/term names (ie, row names of `nodeInfo`) in the object
- `id()`: retrieve domain id (ie, column 'id' of `nodeInfo`) in the object, if any
- `level()`: retrieve domain level (ie, column 'level' of `nodeInfo`) in the object, if any
- `description()`: retrieve domain description (ie, column 'description' of `nodeInfo`) in the object, if any

Standard generic methods:

- `str()`: compact display of the content in the object
- `show()`: abbreviated display of the object
- `as(matrix, "Dnetwork")`: convert a matrix to an object of class `Dnetwork`
- `as(dgCMatrix, "Dnetwork")`: convert a `dgCMatrix` (a sparse matrix) to an object of class `Dnetwork`
- `[i]`: get the subset of the same class

Access

Ways to access information on this class:

- `showClass("Dnetwork")`: show the class definition
- `showMethods(classes="Dnetwork")`: show the method definition upon this class
- `getSlots("Dnetwork")`: get the name and class of each slot in this class
- `slotNames("Dnetwork")`: get the name of each slot in this class
- `selectMethod(f, signature="Dnetwork")`: retrieve the definition code for the method 'f' defined in this class

See Also

[Dnetwork-method](#)

Examples

```
# create an object of class Dnetwork, only given a matrix
adjM <- matrix(runif(25),nrow=5,ncol=5)
as(adjM, "Dnetwork")
```

```
# create an object of class Dnetwork, given a matrix plus information on nodes
# 1) create nodeI: an object of class InfoDataFrame
data <- data.frame(id=paste("Domain", 1:5, sep="_"),
```

```

level=rep("SCOP",5), description=I(LETTERS[1:5]),
row.names=paste("Domain", 1:5, sep="_")
nodeI <- new("InfoDataFrame", data=data)
nodeI
# 2) create an object of class Dnetwork
# VERY IMPORTANT: make sure having consistent names between nodeInfo and adjMatrix
adjM <- matrix(runif(25),nrow=5,ncol=5)
colnames(adjM) <- rownames(adjM) <- rowNames(nodeI)
x <- new("Dnetwork", adjMatrix=adjM, nodeInfo=nodeI)
x
# 3) look at various methods defined on class Dnetwork
dim(x)
adjMatrix(x)
nodeInfo(x)
nInfo(x)
nodeNames(x)
id(x)
level(x)
description(x)
# 4) get the subset
x[1:2]

```

Dnetwork-method

Methods defined for S4 class Dnetwork

Description

Methods defined for class Dnetwork.

Usage

```
## S4 method for signature 'Dnetwork'
dim(x)
```

```
## S4 method for signature 'Dnetwork'
adjMatrix(x)
```

```
## S4 method for signature 'Dnetwork'
nodeInfo(x)
```

```
## S4 method for signature 'Dnetwork'
nInfo(object)
```

```
## S4 method for signature 'Dnetwork'
nodeNames(object)
```

```
## S4 method for signature 'Dnetwork'
id(object)
```

```
## S4 method for signature 'Dnetwork'
level(object)

## S4 method for signature 'Dnetwork'
description(object)

## S4 method for signature 'Dnetwork'
show(object)

## S4 method for signature 'Dnetwork,ANY,ANY,ANY'
x[i, j, ..., drop = FALSE]
```

Arguments

| | |
|--------|---|
| x | an object of class Dnetwork |
| object | an object of class Dnetwork |
| i | an index |
| j | an index |
| ... | additional parameters |
| drop | For matrices and arrays. If TRUE the result is coerced to the lowest possible dimension (see the examples). This only works for extracting elements, not for the replacement. See drop for further details. |

See Also

[Dnetwork-class](#)

Eoutput-class

Definition for S4 class Eoutput

Description

Eoutput is an S4 class to store output from enrichment analysis by [dcEnrichment](#).

Value

Class Eoutput

Slots

domain A character specifying the domain identity

ontology A character specifying the ontology identity

term_info A data.frame of nTerm X 5 containing term information, where nTerm is the number of terms in consideration, and the 5 columns are "term_id" (i.e. "Term ID"), "term_name" (i.e. "Term Name"), "namespace" (i.e. "Term Namespace"), "distance" (i.e. "Term Distance") and "IC" (i.e. "Information Content for the term based on annotation frequency by it")

- anno** A list of terms, each storing annotated domains (also within the background domains). Always, terms are identified by "term_id" and domain members identified by their ids (e.g. sunids for SCOP domains)
- data** A vector containing input data in [dcEnrichment](#). It is not always the same as the input data as only those mappable are retained
- background** A vector containing background in [dcEnrichment](#). It is not always the same as the input background (if provided by the user) as only those mappable are retained
- overlap** A list of terms, each storing domains overlapped between domains annotated by a term and domains in the input data (i.e. the domains of interest). Always, terms are identified by "term_id" and domain members identified by their ids (e.g. sunids for SCOP domains)
- zscore** A vector of terms, containing z-scores
- pvalue** A vector of terms, containing p-values
- adjp** A vector of terms, containing adjusted p-values. It is the p value but after being adjusted for multiple comparisons

Creation

An object of this class can be created via: `new("Eoutput", domain, ontology, term_info, anno, data, overlap, zscore, pvalue, adjp)`

Methods

Class-specific methods:

- `zscore()`: retrieve the slot 'zscore' in the object
- `pvalue()`: retrieve the slot 'pvalue' in the object
- `adjp()`: retrieve the slot 'adjp' in the object
- `view()`: retrieve an integrated data.frame used for viewing the object
- `write()`: write the object into a local file

Standard generic methods:

- `str()`: compact display of the content in the object
- `show()`: abbreviated display of the object

Access

Ways to access information on this class:

- `showClass("Eoutput")`: show the class definition
- `showMethods(classes="Eoutput")`: show the method definition upon this class
- `getSlots("Eoutput")`: get the name and class of each slot in this class
- `slotNames("Eoutput")`: get the name of each slot in this class
- `selectMethod(f, signature="Eoutput")`: retrieve the definition code for the method 'f' defined in this class

See Also[Eoutput-method](#)**Examples**

```
## Not run:
# 1) load SCOP.sf (as 'InfoDataFrame' object)
SCOP.sf <- dcRDataLoader('SCOP.sf')
# randomly select 20 domains
data <- sample(rowNames(SCOP.sf), 20)

# 2) perform enrichment analysis, producing an object of S4 class 'Eoutput'
eoutput <- dcEnrichment(data, domain="SCOP.sf", ontology="GOMF")
eoutput

# 3) write into the file 'Eoutput.txt' in your local directory
write(eoutput, file='Eoutput.txt')

# 4) view the top 5 significant terms
view(eoutput, top_num=5, sortBy="pvalue", details=TRUE)

# 4) retrieve several slots directly
zscore(eoutput)[1:5]
pvalue(eoutput)[1:5]
adjp(eoutput)[1:5]

## End(Not run)
```

Eoutput-method

Methods defined for S4 class Eoutput

Description

Methods defined for S4 class Eoutput.

Usage

```
## S4 method for signature 'Eoutput'
show(object)

## S4 method for signature 'Eoutput'
zscore(x)

## S4 method for signature 'Eoutput'
pvalue(x)

## S4 method for signature 'Eoutput'
adjp(x)
```

```
## S4 method for signature 'Eoutput'
view(x, top_num = 5, sortBy = c("pvalue", "adjp",
"zscore", "nAnno", "nOverlap", "none"), decreasing = NULL, details = T)

## S4 method for signature 'Eoutput'
write(x, file = "Eoutput.txt", verbose = T)
```

Arguments

| | |
|------------|--|
| object | an object of S4 class Eoutput. Usually this is an output from dcEnrichment |
| x | an object of S4 class Eoutput. Usually this is an output from dcEnrichment |
| top_num | the maximum number (5, by default) of terms will be viewed. If NULL or NA, all terms will be viewed (this can be used for the subsequent saving) |
| sortBy | which statistics will be used for sorting and viewing terms. It can be "pvalue" for p value, "adjp" for adjusted p value, "zscore" for enrichment z-score, "nAnno" for the number in domains annotated by a term, "nOverlap" for the number in overlaps, and "none" for ordering simply according to ID of terms |
| decreasing | logical to indicate whether to sort in a decreasing order. If it is null, by default it will be true for "zscore", "nAnno" or "nOverlap"; otherwise false |
| details | logical to indicate whether the detailed information of terms is also viewed. By default, it sets to TRUE for the inclusion |
| file | a character specifying a file name written into. By default, it is 'Eoutput.txt' |
| verbose | logical to indicate whether the messages will be displayed in the screen. By default, it sets to true for display |

Value

view(x) returns a data frame with following components:

- term_id: term ID
- nAnno: number in domains annotated by a term
- nGroup: number in domains from the input group
- nOverlap: number in overlaps
- zscore: enrichment z-score
- pvalue: p value
- adjp: adjusted p value
- term_name: term name
- term_namespace: term namespace; optional, it is only appended when "details" is true
- term_distance: term distance; optional, it is only appended when "details" is true
- members: members (represented as domain IDs) in overlaps; optional, it is only appended when "details" is true

write(x) also returns the same data frame as view(x), in addition to a specified local file.

See Also[Eoutput-class](#)

eTOL

*eukaryotic Tree Of Life (eTOL)***Description**

A 'phylo' object that contains information about eukaryotic part of species tree of life (eTOL). It is a rooted binary tree. Tips represent extant genomes. Since its reconstruction is guided under the NCBI taxonomy, each internal node is either mapped onto a unique taxonomic identifier or left empty (assumedly a hypothetical unknown ancestral genome).

Usage

```
data(eTOL)
```

Value

an object of class "phylo" with the following components:

- `Nnode`: the number of (internal) nodes
- `tip.label`: a vector giving the names of the tips (i.e., "left_id" to define the post-ordered binary tree structure)
- `node.label`: a vector giving the names of the internal nodes (i.e., "left_id" to define the post-ordered binary tree structure)
- `genome_info`: a matrix of all nodes (including tips and internal nodes) X 8, giving extant/ancestral genome information: "left_id" (unique and used as internal id), "right_id" (used in combination with "left_id" to define the post-ordered binary tree structure), "taxon_id" (NCBI taxonomy id, if matched), "genome" (2-letter genome identifiers used in SUPER-FAMILY, if being extant), "name" (NCBI taxonomy name, if matched), "rank" (NCBI taxonomy rank, if matched), "branchlength" (branch length in relevance to the parent), and "common_name" (NCBI taxonomy common name, if matched and existed)
- `edge`: a two-column matrix of mode numeric where each row represents an edge of the tree; the nodes and the tips are symbolized with numbers; the tips are numbered 1, 2, ..., and the internal nodes are numbered after the tips. For each row, the first column gives the ancestor
- `edge.length`: a numeric vector giving the lengths of the branches given by 'edge'
- `root.length`: a numeric value giving the length of the branch at the root
- `connectivity`: a matrix of internal nodes X all nodes (including tips and internal nodes), with 1 for the presence of a ancestor-descendant path, and 0 otherwise

References

Fang et al. (2013) A daily-updated tree of (sequenced) life as a reference for genome research. *Scientific reports*, 3:2015.

Examples

```

data(eTOL)
eTOL
# list all components
names(eTOL)
# extract information about the first 5 genomes
eTOL$genome_info[1:5,]
# look at the dimension of connectivity
dim(eTOL$connectivity)
## Not run:
# visualise the connectivity matrix
Ntip <- length(eTOL$tip.label) # number of tips
Nnode <- eTOL$Nnode # number of internal nodes
data <- eTOL$connectivity
visHeatmapAdv(data, Rowv=FALSE, Colv=FALSE, zlim=c(0,1),
  colormap="gray-black",
  add.expr=abline(v=c(1,Ntip+1,(Ntip+Nnode+1))-0.5, col="white"),
  key=FALSE, labRow=NA, labCol=NA)

## End(Not run)

```

InfoDataFrame-class *Definition for S4 class InfoDataFrame*

Description

InfoDataFrame has two slots: data and dimLabels.

Value

Class InfoDataFrame

Slots

data A data.frame containing terms (rows) and measured variables (columns).
dimLabels A character describing labels for rows and columns.

Creation

An object of this class can be created via: `new("InfoDataFrame", data, dimLabels)`

Methods

Class-specific methods:

- `dim()`: retrieve the dimension in the object
- `nrow()`: retrieve number of rows in the object
- `ncol()`: retrieve number of columns in the object

- `rowNames()`: retrieve names of rows in the object
- `colNames()`: retrieve names of columns in the object
- `dimLabels()`: retrieve the slot 'dimLabels', containing labels used for display of rows and columns in the object
- `Data()`: retrieve the slot 'data' in the object

Standard generic methods:

- `str()`: compact display of the content in the object
- `show()`: abbreviated display of the object
- `as(data.frame, "InfoDataFrame")`: convert a data.frame to an object of class InfoDataFrame
- `[i, j]`: get the subset of the same class

Access

Ways to access information on this class:

- `showClass("InfoDataFrame")`: show the class definition
- `showMethods(classes="InfoDataFrame")`: show the method definition upon this class
- `getSlots("InfoDataFrame")`: get the name and class of each slot in this class
- `slotNames("InfoDataFrame")`: get the name of each slot in this class
- `selectMethod(f, signature="InfoDataFrame")`: retrieve the definition code for the method 'f' defined in this class

See Also

[InfoDataFrame-method](#)

Examples

```
# generate data on domain information on
data <- data.frame(x=1:10, y=I(LETTERS[1:10]),
  row.names=paste("Domain", 1:10, sep="_"))
dimLabels <- c("rowLabels", "colLabels")
# create an object of class InfoDataFrame
x <- new("InfoDataFrame", data=data, dimLabels=dimLabels)
x
# alternatively, using coerce methods
x <- as(data, "InfoDataFrame")
x
# look at various methods defined on class Anno
dimLabels(x)
dim(x)
nrow(x)
ncol(x)
rowNames(x)
colNames(x)
Data(x)
x[1:3,]
```

InfoDataFrame-method *Methods defined for S4 class InfoDataFrame*

Description

Methods defined for class InfoDataFrame.

Usage

```
dimLabels(x)

## S4 method for signature 'InfoDataFrame'
dimLabels(x)

## S4 method for signature 'InfoDataFrame'
dim(x)

## S4 method for signature 'InfoDataFrame'
nrow(x)

## S4 method for signature 'InfoDataFrame'
ncol(x)

## S4 method for signature 'InfoDataFrame'
rowNames(x)

## S4 method for signature 'InfoDataFrame'
colNames(x)

## S4 method for signature 'InfoDataFrame'
Data(x)

## S4 method for signature 'InfoDataFrame'
show(object)

## S4 method for signature 'InfoDataFrame,ANY,ANY,ANY'
x[i, j, ..., drop = FALSE]
```

Arguments

| | |
|--------|----------------------------------|
| x | an object of class InfoDataFrame |
| object | an object of class InfoDataFrame |
| i | an index |
| j | an index |
| ... | additional parameters |

drop For matrices and arrays. If TRUE the result is coerced to the lowest possible dimension (see the examples). This only works for extracting elements, not for the replacement. See [drop](#) for further details.

See Also

[InfoDataFrame-class](#)

InterPro *InterPro domains (InterPro).*

Description

An object of class "InfoDataFrame" that contains information on InterPro domains (InterPro). This data is prepared based on <ftp://anonymous@ftp.ebi.ac.uk/pub/databases/interpro/Current/entry.list>.

Usage

```
data(InterPro)
```

Value

an object of class [InfoDataFrame](#). It has slots for data and dimLabels:

- data: a data.frame containing information about 11638 annotatable domains (in rows), with 3 columns ("id" for InterPro ID, and "level" always equals "InterPro", "description" for InterPro description)
- dimLabels: a character describing labels for rows and columns in data

References

Hunter et al. (2012) InterPro in 2011: new developments in the family and domain prediction database. *Nucleic Acids Res*, 40(Database issue):D306-12.

See Also

[InfoDataFrame-class](#)

Examples

```
# load data
data(InterPro)
InterPro
# retrieve the dimension
dim(InterPro)
# retrieve names of columns
colNames(InterPro)
# retrieve the first 5 rows of data
Data(InterPro)[1:5,]
```

| | |
|---------------|--|
| InterPro2GOBP | <i>Annotations of InterPro domains by Gene Ontology Biological Process (GOBP).</i> |
|---------------|--|

Description

An object of class "Anno" that contains associations between Gene Ontology Biological Process terms and InterPro domains. This data is prepared based on the InterPro database (see <http://www.ebi.ac.uk/interpro/>) and <ftp://anonymous@ftp.ebi.ac.uk/pub/databases/interpro/Current/interpro2go>.

Usage

```
data(InterPro2GOBP)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for InterPro ID, and "level" always equals "InterPro", "description" for InterPro description

References

Hunter et al. (2012) Manual GO annotation of predictive protein signatures: the InterPro approach to GO curation. *Database (Oxford)*, 2012:bar068.

See Also

[Anno-class](#)

Examples

```
# load data
data(InterPro2GOBP)
InterPro2GOBP
# retrieve info on ontology terms
termData(InterPro2GOBP)
# retrieve info on InterPro domains
domainData(InterPro2GOBP)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(InterPro2GOBP)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

| | |
|---------------|--|
| InterPro2GOCC | <i>Annotations of InterPro domains by Gene Ontology Cellular Component (GOCC).</i> |
|---------------|--|

Description

An object of class "Anno" that contains associations between Gene Ontology Cellular Component terms and InterPro domains. This data is prepared based on the InterPro database (see <http://www.ebi.ac.uk/interpro/>) and <ftp://anonymous@ftp.ebi.ac.uk/pub/databases/interpro/Current/interpro2go>.

Usage

```
data(InterPro2GOCC)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for InterPro ID, and "level" always equals "InterPro", "description" for InterPro description

References

Hunter et al. (2012) Manual GO annotation of predictive protein signatures: the InterPro approach to GO curation. *Database (Oxford)*, 2012:bar068.

See Also

[Anno-class](#)

Examples

```
# load data
data(InterPro2GOCC)
InterPro2GOCC
# retrieve info on ontology terms
termData(InterPro2GOCC)
# retrieve info on InterPro domains
domainData(InterPro2GOCC)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(InterPro2GOCC)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

| | |
|---------------|--|
| InterPro2GOMF | <i>Annotations of InterPro domains by Gene Ontology Molecular Function (GOMF).</i> |
|---------------|--|

Description

An object of class "Anno" that contains associations between Gene Ontology Molecular Function terms and InterPro domains. This data is prepared based on the InterPro database (see <http://www.ebi.ac.uk/interpro/>) and <ftp://anonymous@ftp.ebi.ac.uk/pub/databases/interpro/Current/interpro2go>.

Usage

```
data(InterPro2GOMF)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for InterPro ID, and "level" always equals "InterPro", "description" for InterPro description

References

Hunter et al. (2012) Manual GO annotation of predictive protein signatures: the InterPro approach to GO curation. *Database (Oxford)*, 2012:bar068.

See Also

[Anno-class](#)

Examples

```
# load data
data(InterPro2GOMF)
InterPro2GOMF
# retrieve info on ontology terms
termData(InterPro2GOMF)
# retrieve info on InterPro domains
domainData(InterPro2GOMF)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(InterPro2GOMF)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

Onto-class

Definition for S4 class Onto

Description

Onto has 2 slots: nodeInfo and adjMatrix

Value

Class Onto

Slots

nodeInfo An object of S4 class [InfoDataFrame](#), describing information on nodes/terms.

adjMatrix An object of S4 class [AdjData](#), containing adjacency data matrix (for a direct graph), with rows for parent (arrow-outbound) and columns for children (arrow-inbound)

Creation

An object of this class can be created via: `new("Onto", nodeInfo, adjMatrix)`

Methods

Class-specific methods:

- `dim()`: retrieve the dimension in the object
- `adjMatrix()`: retrieve the slot 'adjMatrix' in the object
- `nodeInfo()`: retrieve the slot 'nodeInfo' (as class [InfoDataFrame](#)) in the object
- `nInfo()`: retrieve nodeInfo (as `data.frame`) in the object
- `nodeName()`: retrieve node/term names (ie, row names of nodeInfo) in the object
- `term_id()`: retrieve term id (ie, column 'term_id' of nodeInfo) in the object, if any
- `term_name()`: retrieve term id (ie, column 'term_name' of nodeInfo) in the object, if any
- `term_namespace()`: retrieve term id (ie, column 'term_namespace' of nodeInfo) in the object, if any
- `term_distance()`: retrieve term id (ie, column 'term_distance' of nodeInfo) in the object, if any

Standard generic methods:

- `str()`: compact display of the content in the object
- `show()`: abbreviated display of the object
- `as(matrix, "Onto")`: convert a matrix to an object of class Onto
- `as(dgCMatrix, "Onto")`: convert a `dgCMatrix` (a sparse matrix) to an object of class Onto
- `[i]`: get the subset of the same class

Access

Ways to access information on this class:

- `showClass("Onto")`: show the class definition
- `showMethods(classes="Onto")`: show the method definition upon this class
- `getSlots("Onto")`: get the name and class of each slot in this class
- `slotNames("Onto")`: get the name of each slot in this class
- `selectMethod(f, signature="Onto")`: retrieve the definition code for the method 'f' defined in this class

See Also

[Onto-method](#)

Examples

```
# create an object of class Onto, only given a matrix
adjM <- matrix(runif(25),nrow=5,ncol=5)
as(adjM, "Onto")

# create an object of class Onto, given a matrix plus information on nodes
# 1) create nodeI: an object of class InfoDataFrame
data <- data.frame(term_id=paste("Term", 1:5, sep="_"),
  term_name=I(LETTERS[1:5]), term_namespace=rep("Namespace",5),
  term_distance=1:5, row.names=paste("Term", 1:5, sep="_"))
nodeI <- new("InfoDataFrame", data=data)
nodeI
# 2) create an object of class Onto
# VERY IMPORTANT: make sure having consistent names between nodeInfo and adjMatrix
adjM <- matrix(runif(25),nrow=5,ncol=5)
colnames(adjM) <- rownames(adjM) <- rowNames(nodeI)
x <- new("Onto", adjMatrix=adjM, nodeInfo=nodeI)
x
# 3) look at various methods defined on class Onto
dim(x)
adjMatrix(x)
nodeInfo(x)
nInfo(x)
nodeName(x)
term_id(x)
term_namespace(x)
term_distance(x)
# 4) get the subset
x[1:2]
```

Onto-method

Methods defined for S4 class Onto

Description

Methods defined for class Onto.

Usage

```
## S4 method for signature 'Onto'  
dim(x)  
  
## S4 method for signature 'Onto'  
adjMatrix(x)  
  
## S4 method for signature 'Onto'  
nodeInfo(x)  
  
## S4 method for signature 'Onto'  
nInfo(object)  
  
## S4 method for signature 'Onto'  
nodeName(object)  
  
## S4 method for signature 'Onto'  
term_id(object)  
  
## S4 method for signature 'Onto'  
term_name(object)  
  
## S4 method for signature 'Onto'  
term_namespace(object)  
  
## S4 method for signature 'Onto'  
term_distance(object)  
  
## S4 method for signature 'Onto'  
show(object)  
  
## S4 method for signature 'Onto,ANY,ANY,ANY'  
x[i, j, ..., drop = FALSE]
```

Arguments

| | |
|--------|-------------------------|
| x | an object of class Onto |
| object | an object of class Onto |

| | |
|------|---|
| i | an index |
| j | an index |
| ... | additional parameters |
| drop | For matrices and arrays. If TRUE the result is coerced to the lowest possible dimension (see the examples). This only works for extracting elements, not for the replacement. See drop for further details. |

See Also

[Onto-class](#)

| | |
|---------|-------------------------------|
| onto.DO | <i>Disease Ontology (DO).</i> |
|---------|-------------------------------|

Description

An R object that contains information on Gene Ontology Biological Process terms. These terms are organised as a direct acyclic graph (DAG), which is further stored as an object of the class 'igraph' (see <http://igraph.org/r/doc/aaa-igraph-package.html>). This data is prepared based on <http://sourceforge.net/p/diseaseontology/code/HEAD/tree/trunk/HumanDO.obo>.

Usage

```
data(onto.DO)
```

Value

an object of S4 class [Onto](#). It has slots for "nodeInfo" and "adjMatrix"

- nodeInfo: an object of S4 class [InfoDataFrame](#), describing information on nodes/terms including: "term_id" (i.e. Term ID), "term_name" (i.e. Term Name), "term_namespace" (i.e. Term Namespace), and "term_distance" (i.e. Term Distance: the distance to the root; always 0 for the root itself)
- adjMatrix: an object of S4 class [AdjData](#), containing adjacency data matrix, with rows for parent (arrow-outbound) and columns for children (arrow-inbound)

References

Schriml et al. (2012) Disease Ontology: a backbone for disease semantic integration. *Nucleic Acids Res*, 40:D940-946.
 Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

Examples

```
data(onto.DO)
onto.DO
```

`onto.GOBP`*Gene Ontology Biological Process (GOBP).*

Description

An R object that contains information on Gene Ontology Biological Process terms. These terms are organised as a direct acyclic graph (DAG), which is further stored as an object of the class 'igraph' (see <http://igraph.org/r/doc/aaa-igraph-package.html>). This data is prepared based on http://www.geneontology.org/ontology/obo_format_1_2/gene_ontology.1_2.obo. Only the edges with the relation (either 'is_a' or 'part_of') are retained.

Usage

```
data(onto.GOBP)
```

Value

an object of S4 class `Onto`. It has slots for "nodeInfo" and "adjMatrix"

- `nodeInfo`: an object of S4 class `InfoDataFrame`, describing information on nodes/terms including: "term_id" (i.e. Term ID), "term_name" (i.e. Term Name), "term_namespace" (i.e. Term Namespace), and "term_distance" (i.e. Term Distance: the distance to the root; always 0 for the root itself)
- `adjMatrix`: an object of S4 class `AdjData`, containing adjacency data matrix, with rows for parent (arrow-outbound) and columns for children (arrow-inbound)

References

Ashburner et al. (2000) Gene ontology: tool for the unification of biology. *Nat Genet*, 25:25-29.
Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

Examples

```
data(onto.GOBP)  
onto.GOBP
```

`onto.GOCC`*Gene Ontology Cellular Component (GOCC).*

Description

An R object that contains information on Gene Ontology Cellular Component terms. These terms are organised as a direct acyclic graph (DAG), which is further stored as an object of the class 'igraph' (see <http://igraph.org/r/doc/aaa-igraph-package.html>). This data is prepared based on http://www.geneontology.org/ontology/obo_format_1_2/gene_ontology.1_2.obo. Only the edges with the relation (either 'is_a' or 'part_of') are retained.

Usage

```
data(onto.GOCC)
```

Value

an object of S4 class [Onto](#). It has slots for "nodeInfo" and "adjMatrix"

- nodeInfo: an object of S4 class [InfoDataFrame](#), describing information on nodes/terms including: "term_id" (i.e. Term ID), "term_name" (i.e. Term Name), "term_namespace" (i.e. Term Namespace), and "term_distance" (i.e. Term Distance: the distance to the root; always 0 for the root itself)
- adjMatrix: an object of S4 class [AdjData](#), containing adjacency data matrix, with rows for parent (arrow-outbound) and columns for children (arrow-inbound)

References

Ashburner et al. (2000) Gene ontology: tool for the unification of biology. *Nat Genet*, 25:25-29.
Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

Examples

```
data(onto.GOCC)  
onto.GOCC
```

onto.GOMF

Gene Ontology Molecular Function (GOMF).

Description

An R object that contains information on Gene Ontology Molecular Function terms. These terms are organised as a direct acyclic graph (DAG), which is further stored as an object of the class 'igraph' (see <http://igraph.org/r/doc/aaa-igraph-package.html>). This data is prepared based on http://www.geneontology.org/ontology/obo_format_1_2/gene_ontology.1_2.obo. Only the edges with the relation (either 'is_a' or 'part_of') are retained.

Usage

```
data(onto.GOMF)
```

Value

an object of S4 class [Onto](#). It has slots for "nodeInfo" and "adjMatrix"

- nodeInfo: an object of S4 class [InfoDataFrame](#), describing information on nodes/terms including: "term_id" (i.e. Term ID), "term_name" (i.e. Term Name), "term_namespace" (i.e. Term Namespace), and "term_distance" (i.e. Term Distance: the distance to the root; always 0 for the root itself)

- `adjMatrix`: an object of S4 class `AdjData`, containing adjacency data matrix, with rows for parent (arrow-outbound) and columns for children (arrow-inbound)

References

Ashburner et al. (2000) Gene ontology: tool for the unification of biology. *Nat Genet*, 25:25-29.
 Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

Examples

```
data(onto.GOMF)
onto.GOMF
```

onto.HPMI

Human Phenotype Mode of Inheritance (HPMI).

Description

An R object that contains information on Human Phenotype Mode of Inheritance terms. These terms are organised as a direct acyclic graph (DAG), which is further stored as an object of the class 'igraph' (see <http://igraph.org/r/doc/aaa-igraph-package.html>). This data is prepared based on <http://compbio.charite.de/svn/hpo/trunk/src/ontology/human-phenotype-ontology.obo>.

Usage

```
data(onto.HPMI)
```

Value

an object of S4 class `Onto`. It has slots for "nodeInfo" and "adjMatrix"

- `nodeInfo`: an object of S4 class `InfoDataFrame`, describing information on nodes/terms including: "term_id" (i.e. Term ID), "term_name" (i.e. Term Name), "term_namespace" (i.e. Term Namespace), and "term_distance" (i.e. Term Distance: the distance to the root; always 0 for the root itself)
- `adjMatrix`: an object of S4 class `AdjData`, containing adjacency data matrix, with rows for parent (arrow-outbound) and columns for children (arrow-inbound)

References

Robinson et al. (2012) The Human Phenotype Ontology: a tool for annotating and analyzing human hereditary disease. *Am J Hum Genet*, 83:610-615.
 Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

Examples

```
data(onto.HPMI)
onto.HPMI
```

onto.HPON

Human Phenotype ONset and clinical course (HPON).

Description

An R object that contains information on Human Phenotype ONset and clinical course terms. These terms are organised as a direct acyclic graph (DAG), which is further stored as an object of the class 'igraph' (see <http://igraph.org/r/doc/aaa-igraph-package.html>). This data is prepared based on <http://compbio.charite.de/svn/hpo/trunk/src/ontology/human-phenotype-ontology.obo>.

Usage

```
data(onto.HPON)
```

Value

an object of S4 class `Onto`. It has slots for "nodeInfo" and "adjMatrix"

- `nodeInfo`: an object of S4 class `InfoDataFrame`, describing information on nodes/terms including: "term_id" (i.e. Term ID), "term_name" (i.e. Term Name), "term_namespace" (i.e. Term Namespace), and "term_distance" (i.e. Term Distance: the distance to the root; always 0 for the root itself)
- `adjMatrix`: an object of S4 class `AdjData`, containing adjacency data matrix, with rows for parent (arrow-outbound) and columns for children (arrow-inbound)

References

Robinson et al. (2012) The Human Phenotype Ontology: a tool for annotating and analyzing human hereditary disease. *Am J Hum Genet*, 83:610-615.

Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

Examples

```
data(onto.HPON)
onto.HPON
```

`onto.HPPA`*Human Phenotype Phenotypic Abnormality (HPPA).*

Description

An R object that contains information on Human Phenotype Phenotypic Abnormality terms. These terms are organised as a direct acyclic graph (DAG), which is further stored as an object of the class 'igraph' (see <http://igraph.org/r/doc/aaa-igraph-package.html>). This data is prepared based on <http://compbio.charite.de/svn/hpo/trunk/src/ontology/human-phenotype-ontology.obo>.

Usage

```
data(onto.HPPA)
```

Value

an object of S4 class `Onto`. It has slots for "nodeInfo" and "adjMatrix"

- `nodeInfo`: an object of S4 class `InfoDataFrame`, describing information on nodes/terms including: "term_id" (i.e. Term ID), "term_name" (i.e. Term Name), "term_namespace" (i.e. Term Namespace), and "term_distance" (i.e. Term Distance: the distance to the root; always 0 for the root itself)
- `adjMatrix`: an object of S4 class `AdjData`, containing adjacency data matrix, with rows for parent (arrow-outbound) and columns for children (arrow-inbound)

References

Robinson et al. (2012) The Human Phenotype Ontology: a tool for annotating and analyzing human hereditary disease. *Am J Hum Genet*, 83:610-615.
Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

Examples

```
data(onto.HPPA)  
onto.HPPA
```

| | |
|---------|----------------------------------|
| onto.MP | <i>Mammalian Phenotype (MP).</i> |
|---------|----------------------------------|

Description

An R object that contains information on Mammalian Phenotype terms. These terms are organised as a direct acyclic graph (DAG), which is further stored as an object of the class 'igraph' (see <http://igraph.org/r/doc/aaa-igraph-package.html>). This data is prepared based on <http://sourceforge.net/p/diseaseontology/code/HEAD/tree/trunk/HumanMP.obo>.

Usage

```
data(onto.MP)
```

Value

an object of S4 class [Onto](#). It has slots for "nodeInfo" and "adjMatrix"

- nodeInfo: an object of S4 class [InfoDataFrame](#), describing information on nodes/terms including: "term_id" (i.e. Term ID), "term_name" (i.e. Term Name), "term_namespace" (i.e. Term Namespace), and "term_distance" (i.e. Term Distance: the distance to the root; always 0 for the root itself)
- adjMatrix: an object of S4 class [AdjData](#), containing adjacency data matrix, with rows for parent (arrow-outbound) and columns for children (arrow-inbound)

References

Smith et al. (2009) The Mammalian Phenotype Ontology: enabling robust annotation and comparative analysis. *Wiley Interdiscip Rev Syst Biol Med*, 1:390-399.
Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

Examples

```
data(onto.MP)  
onto.MP
```

| | |
|------|-----------------------------|
| Pfam | <i>Pfam domains (Pfam).</i> |
|------|-----------------------------|

Description

An object of class "InfoDataFrame" that contains information on Pfam domains (Pfam). This data is prepared based on ftp://ftp.sanger.ac.uk/pub/databases/Pfam/current_release/database_files/pfamA.txt.gz.

Usage

```
data(Pfam)
```

Value

an object of class `InfoDataFrame`. It has slots for data and dimLabels:

- data: a data.frame containing information about 14831 domains (in rows), with 3 columns ("id" for Pfam accession ID, and "level" always equals "Pfam", "description" for Pfam description)
- dimLabels: a character describing labels for rows and columns in data

References

Finn et al. (2014) The Pfam protein families database. *Nucleic Acids Res*, 42(Database issue):D222-D230.

See Also

[InfoDataFrame-class](#)

Examples

```
# load data
data(Pfam)
Pfam
# retrieve the dimension
dim(Pfam)
# retrieve names of columns
colNames(Pfam)
# retrieve the first 5 rows of data
Data(Pfam)[1:5,]
```

Pfam2GOBP

Annotations of Pfam domains by Gene Ontology Biological Process (GOBP).

Description

An object of class "Anno" that contains associations between Gene Ontology Biological Process terms and Pfam domains. This data is prepared based on the Pfam database (see <http://pfam.xfam.org>) and <ftp://ftp.geneontology.org/pub/go/external2go/pfam2go>.

Usage

```
data(Pfam2GOBP)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for Pfam accession ID, and "level" always equals "Pfam", "description" for Pfam description

References

Finn et al. (2014) The Pfam protein families database. *Nucleic Acids Res*, 42(Database issue):D222-D230.

See Also

[Anno-class](#)

Examples

```
# load data
data(Pfam2GOBP)
Pfam2GOBP
# retrieve info on ontology terms
termData(Pfam2GOBP)
# retrieve info on Pfam domains
domainData(Pfam2GOBP)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(Pfam2GOBP)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

Pfam2GOCC

Annotations of Pfam domains by Gene Ontology Cellular Component (GOCC).

Description

An object of class "Anno" that contains associations between Gene Ontology Cellular Component terms and Pfam domains. This data is prepared based on the Pfam database (see <http://pfam.xfam.org>) and <ftp://ftp.geneontology.org/pub/go/external2go/pfam2go>.

Usage

```
data(Pfam2GOCC)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for Pfam accession ID, and "level" always equals "Pfam", "description" for Pfam description

References

Finn et al. (2014) The Pfam protein families database. *Nucleic Acids Res*, 42(Database issue):D222-D230.

See Also

[Anno-class](#)

Examples

```
# load data
data(Pfam2GOCC)
Pfam2GOCC
# retrieve info on ontology terms
termData(Pfam2GOCC)
# retrieve info on Pfam domains
domainData(Pfam2GOCC)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(Pfam2GOCC)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

Pfam2GOMF

Annotations of Pfam domains by Gene Ontology Molecular Function (GOMF).

Description

An object of class "Anno" that contains associations between Gene Ontology Molecular Function terms and Pfam domains. This data is prepared based on the Pfam database (see <http://pfam.xfam.org>) and <ftp://ftp.geneontology.org/pub/go/external2go/pfam2go>.

Usage

```
data(Pfam2GOMF)
```


Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for Pfam accession ID, and "level" always equals "Pfam", "description" for Pfam description

References

Finn et al. (2014) The Pfam protein families database. *Nucleic Acids Res*, 42(Database issue):D222-D230.

See Also

[Anno-class](#)

Examples

```
# load data
data(Pfam2GOMF)
Pfam2GOMF
# retrieve info on ontology terms
termData(Pfam2GOMF)
# retrieve info on Pfam domains
domainData(Pfam2GOMF)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(Pfam2GOMF)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

Rfam

RNA families (Rfam).

Description

An object of class "InfoDataFrame" that contains information on RNA families (Rfam). This data is prepared based on ftp://anonymous@ftp.sanger.ac.uk/pub/databases/Rfam/11.0/database_files/rfam.txt.gz.

Usage

```
data(Rfam)
```

Value

an object of class `InfoDataFrame`. It has slots for data and dimLabels:

- data: a data.frame containing information about 2208 RNA families (in rows), with 3 columns ("id" for Rfam accession ID, and "level" always equals "Rfam", "description" for Rfam description)
- dimLabels: a character describing labels for rows and columns in data

References

Gardner et al. (2011) Rfam: Wikipedia, clans and the "decimal" release. *Nucleic Acids Res*, 39(Database issue):D141-D145.

See Also

[InfoDataFrame-class](#)

Examples

```
# load data
data(Rfam)
Rfam
# retrieve the dimension
dim(Rfam)
# retrieve names of columns
colNames(Rfam)
# retrieve the first 5 rows of data
Data(Rfam)[1:5,]
```

Rfam2GOBP

Annotations of Rfam RNA families by Gene Ontology Biological Process (GOBP).

Description

An object of class "Anno" that contains associations between Gene Ontology Biological Process terms and Rfam RNA families. This data is prepared based on the Rfam database (see <http://rfam.xfam.org>) and <http://geneontology.org/external2go/rfam2go>.

Usage

```
data(Rfam2GOBP)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of RNAs X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing RNAs (i.e. rows in annoData), including: "id" for Rfam accession ID, and "level" always equals "Rfam", "description" for Rfam description

References

Gardner et al. (2011) Rfam: Wikipedia, clans and the "decimal" release. *Nucleic Acids Res*, 39(Database issue):D141-D145.

See Also

[Anno-class](#)

Examples

```
# load data
data(Rfam2GOBP)
Rfam2GOBP
# retrieve info on ontology terms
termData(Rfam2GOBP)
# retrieve info on Rfam RNAs
domainData(Rfam2GOBP)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(Rfam2GOBP)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

Rfam2GOCC

Annotations of Rfam RNA families by Gene Ontology Cellular Component (GOCC).

Description

An object of class "Anno" that contains associations between Gene Ontology Cellular Component terms and Rfam RNA families. This data is prepared based on the Rfam database (see <http://rfam.xfam.org>) and <http://geneontology.org/external2go/rfam2go>.

Usage

```
data(Rfam2GOCC)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of RNAs X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing RNAs (i.e. rows in annoData), including: "id" for Rfam accession ID, and "level" always equals "Rfam", "description" for Rfam description

References

Gardner et al. (2011) Rfam: Wikipedia, clans and the "decimal" release. *Nucleic Acids Res*, 39(Database issue):D141-D145.

See Also

[Anno-class](#)

Examples

```
# load data
data(Rfam2GOCC)
Rfam2GOCC
# retrieve info on ontology terms
termData(Rfam2GOCC)
# retrieve info on Rfam RNAs
domainData(Rfam2GOCC)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(Rfam2GOCC)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

Rfam2GOMF

Annotations of Rfam RNA families by Gene Ontology Molecular Function (GOMF).

Description

An object of class "Anno" that contains associations between Gene Ontology Molecular Function terms and Rfam RNA families. This data is prepared based on the Rfam database (see <http://rfam.xfam.org>) and <http://geneontology.org/external2go/rfam2go>.

Usage

```
data(Rfam2GOMF)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of RNAs X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing RNAs (i.e. rows in annoData), including: "id" for Rfam accession ID, and "level" always equals "Rfam", "description" for Rfam description

References

Gardner et al. (2011) Rfam: Wikipedia, clans and the "decimal" release. *Nucleic Acids Res*, 39(Database issue):D141-D145.

See Also

[Anno-class](#)

Examples

```
# load data
data(Rfam2GOMF)
Rfam2GOMF
# retrieve info on ontology terms
termData(Rfam2GOMF)
# retrieve info on Rfam RNAs
domainData(Rfam2GOMF)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(Rfam2GOMF)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

SCOP.fa

SCOP domain families (fa).

Description

An object of class "InfoDataFrame" that contains information on SCOP domain families (fa).

Usage

```
data(SCOP.fa)
```

Value

an object of class `InfoDataFrame`. It has slots for data and dimLabels:

- data: a data.frame containing information about 2223 domains (in rows), with 3 columns ("id" for SCOP sunid, and "level" for SCOP level, "description" for SCOP description)
- dimLabels: a character describing labels for rows and columns in data

References

Morais et al. (2011) SUPERFAMILY 1.75 including a domain-centric gene ontology method. *Nucleic Acids Res*, 39(Database issue):D427-34.
Andreeva et al. (2008) Data growth and its impact on the SCOP database: new developments. *Nucleic Acids Res*, 36(Database issue):D419-425

See Also

[InfoDataFrame-class](#)

Examples

```
# load data
data(SCOP.fa)
SCOP.fa
# retrieve the dimension
dim(SCOP.fa)
# retrieve names of columns
colNames(SCOP.fa)
# retrieve the first 5 rows of data
Data(SCOP.fa)[1:5,]
```

SCOP.fa2DO

Annotations of SCOP domain families (fa) by Disease Ontology (DO).

Description

An object of class "Anno" that contains associations between Disease Ontology terms and SCOP domain families (fa). This data is prepared based on the dcGO database (see <http://supfam.org/SUPERFAMILY/dcGO/>).

Usage

```
data(SCOP.fa2DO)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for SCOP sunid, and "level" for SCOP level, "description" for SCOP description

References

Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

See Also

[Anno-class](#)

Examples

```
# load data
data(SCOP.fa2D0)
SCOP.fa2D0
# retrieve info on ontology terms
termData(SCOP.fa2D0)
# retrieve info on SCOP domains
domainData(SCOP.fa2D0)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(SCOP.fa2D0)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

SCOP.fa2GOBP

Annotations of SCOP domain families (fa) by Gene Ontology Biological Process (GOBP).

Description

An object of class "Anno" that contains associations between Gene Ontology Biological Process terms and SCOP domain families (fa). This data is prepared based on the dcGO database (see <http://supfam.org/SUPERFAMILY/dcGO/>).

Usage

```
data(SCOP.fa2GOBP)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for SCOP sunid, and "level" for SCOP level, "description" for SCOP description

References

Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

See Also

[Anno-class](#)

Examples

```
# load data
data(SCOP.fa2GOBP)
SCOP.fa2GOBP
# retrieve info on ontology terms
termData(SCOP.fa2GOBP)
# retrieve info on SCOP domains
domainData(SCOP.fa2GOBP)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(SCOP.fa2GOBP)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

SCOP.fa2GOCC

Annotations of SCOP domain families (fa) by Gene Ontology Cellular Component (GOCC).

Description

An object of class "Anno" that contains associations between Gene Ontology Cellular Component terms and SCOP domain families (fa). This data is prepared based on the dcGO database (see <http://supfam.org/SUPERFAMILY/dcGO/>).

Usage

```
data(SCOP.fa2GOCC)
```


Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for SCOP sunid, and "level" for SCOP level, "description" for SCOP description

References

Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

See Also

[Anno-class](#)

Examples

```
# load data
data(SCOP.fa2GOCC)
SCOP.fa2GOCC
# retrieve info on ontology terms
termData(SCOP.fa2GOCC)
# retrieve info on SCOP domains
domainData(SCOP.fa2GOCC)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(SCOP.fa2GOCC)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

SCOP.fa2GOMF

Annotations of SCOP domain families (fa) by Gene Ontology Molecular Function (GOMF).

Description

An object of class "Anno" that contains associations between Gene Ontology Molecular Function terms and SCOP domain families (fa). This data is prepared based on the dcGO database (see <http://supfam.org/SUPERFAMILY/dcGO/>).

Usage

```
data(SCOP.fa2GOMF)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for SCOP sunid, and "level" for SCOP level, "description" for SCOP description

References

Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

See Also

[Anno-class](#)

Examples

```
# load data
data(SCOP.fa2GOMF)
SCOP.fa2GOMF
# retrieve info on ontology terms
termData(SCOP.fa2GOMF)
# retrieve info on SCOP domains
domainData(SCOP.fa2GOMF)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(SCOP.fa2GOMF)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

SCOP.fa2HPMI

Annotations of SCOP domain families (fa) by Human Phenotype Mode of Inheritance (HPMI).

Description

An object of class "Anno" that contains associations between HPMI terms and SCOP domain families (fa). This data is prepared based on the dcGO database (see <http://supfam.org/SUPERFAMILY/dcGO/>).

Usage

```
data(SCOP.fa2HPMI)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for SCOP sunid, and "level" for SCOP level, "description" for SCOP description

References

Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

See Also

[Anno-class](#)

Examples

```
# load data
data(SCOP.fa2HPMI)
SCOP.fa2HPMI
# retrieve info on ontology terms
termData(SCOP.fa2HPMI)
# retrieve info on SCOP domains
domainData(SCOP.fa2HPMI)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(SCOP.fa2HPMI)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

SCOP.fa2HPON

Annotations of SCOP domain families (fa) by Human Phenotype ON-set and clinical course (HPON).

Description

An object of class "Anno" that contains associations between HPON terms and SCOP domain families (fa). This data is prepared based on the dcGO database (see <http://supfam.org/SUPERFAMILY/dcGO/>).

Usage

```
data(SCOP.fa2HPON)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for SCOP sunid, and "level" for SCOP level, "description" for SCOP description

References

Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

See Also

[Anno-class](#)

Examples

```
# load data
data(SCOP.fa2HPON)
SCOP.fa2HPON
# retrieve info on ontology terms
termData(SCOP.fa2HPON)
# retrieve info on SCOP domains
domainData(SCOP.fa2HPON)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(SCOP.fa2HPON)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

SCOP . fa2HPPA

Annotations of SCOP domain families (fa) by Human Phenotype Phenotypic Abnormality (HPPA).

Description

An object of class "Anno" that contains associations between HPPA terms and SCOP domain families (fa). This data is prepared based on the dcGO database (see <http://supfam.org/SUPERFAMILY/dcGO/>).

Usage

```
data(SCOP . fa2HPPA)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for SCOP sunid, and "level" for SCOP level, "description" for SCOP description

References

Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

See Also

[Anno-class](#)

Examples

```
# load data
data(SCOP.fa2HPPA)
SCOP.fa2HPPA
# retrieve info on ontology terms
termData(SCOP.fa2HPPA)
# retrieve info on SCOP domains
domainData(SCOP.fa2HPPA)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(SCOP.fa2HPPA)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

SCOP.fa2MP

Annotations of SCOP domain families (fa) by Mammalian Phenotype (MP).

Description

An object of class "Anno" that contains associations between Mammalian Phenotype terms and SCOP domain families (fa). This data is prepared based on the dcGO database (see <http://supfam.org/SUPERFAMILY/dcGO/>).

Usage

```
data(SCOP.fa2MP)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for SCOP sunid, and "level" for SCOP level, "description" for SCOP description

References

Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

See Also

[Anno-class](#)

Examples

```
# load data
data(SCOP.fa2MP)
SCOP.fa2MP
# retrieve info on ontology terms
termData(SCOP.fa2MP)
# retrieve info on SCOP domains
domainData(SCOP.fa2MP)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(SCOP.fa2MP)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

SCOP.sf

SCOP domain superfamilies (sf).

Description

An object of class "InfoDataFrame" that contains information on SCOP domain superfamilies (sf).

Usage

```
data(SCOP.sf)
```

Value

an object of class `InfoDataFrame`. It has slots for data and dimLabels:

- data: a data.frame containing information about 2223 domains (in rows), with 3 columns ("id" for SCOP sunid, and "level" for SCOP level, "description" for SCOP description)
- dimLabels: a character describing labels for rows and columns in data

References

Morais et al. (2011) SUPERFAMILY 1.75 including a domain-centric gene ontology method. *Nucleic Acids Res*, 39(Database issue):D427-34.
Andreeva et al. (2008) Data growth and its impact on the SCOP database: new developments. *Nucleic Acids Res*, 36(Database issue):D419-425

See Also

[InfoDataFrame-class](#)

Examples

```
# load data
data(SCOP.sf)
SCOP.sf
# retrieve the dimension
dim(SCOP.sf)
# retrieve names of columns
colNames(SCOP.sf)
# retrieve the first 5 rows of data
Data(SCOP.sf)[1:5,]
```

SCOP.sf2DO

Annotations of SCOP domain superfamilies (sf) by Disease Ontology (DO).

Description

An object of class "Anno" that contains associations between Disease Ontology terms and SCOP domain superfamilies (sf). This data is prepared based on the dcGO database (see <http://supfam.org/SUPERFAMILY/dcGO/>).

Usage

```
data(SCOP.sf2DO)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for SCOP sunid, and "level" for SCOP level, "description" for SCOP description

References

Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

See Also

[Anno-class](#)

Examples

```
# load data
data(SCOP.sf2D0)
SCOP.sf2D0
# retrieve info on ontology terms
termData(SCOP.sf2D0)
# retrieve info on SCOP domains
domainData(SCOP.sf2D0)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(SCOP.sf2D0)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

SCOP.sf2GOBP

Annotations of SCOP domain superfamilies (sf) by Gene Ontology Biological Process (GOBP).

Description

An object of class "Anno" that contains associations between Gene Ontology Biological Process terms and SCOP domain superfamilies (sf). This data is prepared based on the dcGO database (see <http://supfam.org/SUPERFAMILY/dcGO/>).

Usage

```
data(SCOP.sf2GOBP)
```


Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for SCOP sunid, and "level" for SCOP level, "description" for SCOP description

References

Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

See Also

[Anno-class](#)

Examples

```
# load data
data(SCOP.sf2GOBP)
SCOP.sf2GOBP
# retrieve info on ontology terms
termData(SCOP.sf2GOBP)
# retrieve info on SCOP domains
domainData(SCOP.sf2GOBP)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(SCOP.sf2GOBP)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

SCOP.sf2GOCC

Annotations of SCOP domain superfamilies (sf) by Gene Ontology Cellular Component (GOCC).

Description

An object of class "Anno" that contains associations between Gene Ontology Cellular Component terms and SCOP domain superfamilies (sf). This data is prepared based on the dcGO database (see <http://supfam.org/SUPERFAMILY/dcGO/>).

Usage

```
data(SCOP.sf2GOCC)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for SCOP sunid, and "level" for SCOP level, "description" for SCOP description

References

Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

See Also

[Anno-class](#)

Examples

```
# load data
data(SCOP.sf2GOCC)
SCOP.sf2GOCC
# retrieve info on ontology terms
termData(SCOP.sf2GOCC)
# retrieve info on SCOP domains
domainData(SCOP.sf2GOCC)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(SCOP.sf2GOCC)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

SCOP.sf2GOMF

Annotations of SCOP domain superfamilies (sf) by Gene Ontology Molecular Function (GOMF).

Description

An object of class "Anno" that contains associations between Gene Ontology Molecular Function terms and SCOP domain superfamilies (sf). This data is prepared based on the dcGO database (see <http://supfam.org/SUPERFAMILY/dcGO/>).

Usage

```
data(SCOP.sf2GOMF)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for SCOP sunid, and "level" for SCOP level, "description" for SCOP description

References

Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

See Also

[Anno-class](#)

Examples

```
# load data
data(SCOP.sf2GOMF)
SCOP.sf2GOMF
# retrieve info on ontology terms
termData(SCOP.sf2GOMF)
# retrieve info on SCOP domains
domainData(SCOP.sf2GOMF)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(SCOP.sf2GOMF)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

SCOP.sf2HPMI

Annotations of SCOP domain superfamilies (sf) by Human Phenotype Mode of Inheritance (HPMI).

Description

An object of class "Anno" that contains associations between HPMI terms and SCOP domain superfamilies (sf). This data is prepared based on the dcGO database (see <http://supfam.org/SUPERFAMILY/dcGO/>).

Usage

```
data(SCOP.sf2HPMI)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for SCOP sunid, and "level" for SCOP level, "description" for SCOP description

References

Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

See Also

[Anno-class](#)

Examples

```
# load data
data(SCOP.sf2HPMI)
SCOP.sf2HPMI
# retrieve info on ontology terms
termData(SCOP.sf2HPMI)
# retrieve info on SCOP domains
domainData(SCOP.sf2HPMI)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(SCOP.sf2HPMI)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

SCOP.sf2HPON

Annotations of SCOP domain superfamilies (sf) by Human Phenotype ONset and clinical course (HPON).

Description

An object of class "Anno" that contains associations between HPON terms and SCOP domain superfamilies (sf). This data is prepared based on the dcGO database (see <http://supfam.org/SUPERFAMILY/dcGO/>).

Usage

```
data(SCOP.sf2HPON)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for SCOP sunid, and "level" for SCOP level, "description" for SCOP description

References

Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

See Also

[Anno-class](#)

Examples

```
# load data
data(SCOP.sf2HPON)
SCOP.sf2HPON
# retrieve info on ontology terms
termData(SCOP.sf2HPON)
# retrieve info on SCOP domains
domainData(SCOP.sf2HPON)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(SCOP.sf2HPON)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

SCOP.sf2HPPA

Annotations of SCOP domain superfamilies (sf) by Human Phenotype Phenotypic Abnormality (HPPA).

Description

An object of class "Anno" that contains associations between HPPA terms and SCOP domain superfamilies (sf). This data is prepared based on the dcGO database (see <http://supfam.org/SUPERFAMILY/dcGO/>).

Usage

```
data(SCOP.sf2HPPA)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for SCOP sunid, and "level" for SCOP level, "description" for SCOP description

References

Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

See Also

[Anno-class](#)

Examples

```
# load data
data(SCOP.sf2HPPA)
SCOP.sf2HPPA
# retrieve info on ontology terms
termData(SCOP.sf2HPPA)
# retrieve info on SCOP domains
domainData(SCOP.sf2HPPA)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(SCOP.sf2HPPA)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

SCOP.sf2MP

Annotations of SCOP domain superfamilies (sf) by Mammalian Phenotype (MP).

Description

An object of class "Anno" that contains associations between Mammalian Phenotype terms and SCOP domain superfamilies (sf). This data is prepared based on the dcGO database (see <http://supfam.org/SUPERFAMILY/dcGO/>).

Usage

```
data(SCOP.sf2MP)
```

Value

an object of class `Anno`. It has slots for "annoData", "termData" and "domainData":

- annoData: a sparse matrix of domains X terms
- termData: variables describing ontology terms (i.e. columns in annoData), including: "ID" (i.e. term ID), "Name" (i.e. term Names), "Namespace" (i.e. term Namespace), and "Distance" (i.e. term Distance to the ontology root)
- domainData: variables describing domains (i.e. rows in annoData), including: "id" for SCOP sunid, and "level" for SCOP level, "description" for SCOP description

References

Fang H and Gough J. (2013) dcGO: database of domain-centric ontologies on functions, phenotypes, diseases and more. *Nucleic Acids Res*, 41(Database issue):D536-44.

See Also

[Anno-class](#)

Examples

```
# load data
data(SCOP.sf2MP)
SCOP.sf2MP
# retrieve info on ontology terms
termData(SCOP.sf2MP)
# retrieve info on SCOP domains
domainData(SCOP.sf2MP)
# retrieve the first 5 rows and columns of annotation data
x <- annoData(SCOP.sf2MP)[1:5,1:5]
x
# convert the above retrieval to the full matrix
as.matrix(x)
```

visEnrichment

Function to visualise enrichment analysis outputs in the context of the ontology hierarchy

Description

visEnrichment is supposed to visualise enrichment analysis outputs (represented as an 'Eoutput' object) in the context of the ontology hierarchy (direct acyclic graph; DAG). Only part of DAG induced by those nodes/terms specified in query nodes (and the mode defining the paths to the root of DAG) will be visualised. Nodes in query are framed in black (by default), and all nodes (in query plus induced) will be color-coded according to a given data.type ('zscore'; otherwise taking the form of 10-based negative logarithm for 'adjp' or 'pvalue'). If no nodes in query, the top 5 significant terms (in terms of adjusted p-value) will be used for visualisation

Usage

```
visEnrichment(e, nodes_query = NULL, num_top_nodes = 5,
  path.mode = c("all_shortest_paths", "shortest_paths", "all_paths"),
  data.type = c("adjp", "pvalue", "zscore"), height = 7, width = 7,
  margin = rep(0.1, 4), colormap = c("yr", "bwr", "jet", "gbr", "wyr",
  "br",
  "rainbow", "wb", "lightyellow-orange"), ncolors = 40, zlim = NULL,
  colorbar = T, colorbar.fraction = 0.1, newpage = T,
  layout.orientation = c("left_right", "top_bottom", "bottom_top",
  "right_left"), node.info = c("both", "none", "term_id", "term_name",
  "full_term_name"), graph.node.attrs = NULL, graph.edge.attrs = NULL,
  node.attrs = NULL)
```

Arguments

| | |
|---------------|--|
| e | an object of S4 class Eoutput |
| nodes_query | a vector containing a list of nodes/terms in query. These nodes are used to produce a subgraph of the ontology DAG induced by them. If NULL, the top significant terms (in terms of p-value) will be determined by the next 'num_top_nodes' |
| num_top_nodes | a numeric value specifying the number of the top significant terms (in terms of p-value) will be used. This parameter does not work if the previous 'nodes_query' has been specified |
| path.mode | the mode of paths induced by nodes in query. It can be "all_paths" for all possible paths to the root, "shortest_paths" for only one path to the root (for each node in query), "all_shortest_paths" for all shortest paths to the root (i.e. for each node, find all shortest paths with the equal lengths) |
| data.type | a character telling which data type for nodes in query is used to color-code nodes. It can be one of 'adjp' for adjusted p-values (by default), 'pvalue' for p-values and 'zscore' for z-scores. When 'adjp' or 'pvalue' is used, 10-based negative logarithm is taken. For the style of how to color-code, please see the next arguments: colormap, ncolors, zlim and colorbar |
| height | a numeric value specifying the height of device |
| width | a numeric value specifying the width of device |
| margin | margins as units of length 4 or 1 |
| colormap | short name for the colormap. It can be one of "yr" (yellow-red colormap; by default), "jet" (jet colormap), "bwr" (blue-white-red colormap), "gbr" (green-black-red colormap), "wyr" (white-yellow-red colormap), "br" (black-red colormap), "wb" (white-black colormap), and "rainbow" (rainbow colormap, that is, red-yellow-green-cyan-blue-magenta). Alternatively, any hyphen-separated HTML color names, e.g. "lightyellow-orange" (by default), "blue-black-yellow", "royalblue-white-sandybrown", "darkgreen-white-darkviolet". A list of standard color names can be found in http://html-color-codes.info/color-names |
| ncolors | the number of colors specified over the colormap |
| zlim | the minimum and maximum z/data values for which colors should be plotted, defaulting to the range of the finite values of z. Each of the given colors will be |

| | |
|--------------------|---|
| | used to color an equispaced interval of this range. The midpoints of the intervals cover the range, so that values just outside the range will be plotted |
| colorbar | logical to indicate whether to append a colorbar. If data is null, it always sets to false |
| colorbar.fraction | the relative fraction of colorbar block against the device size |
| newpage | logical to indicate whether to open a new page. By default, it sets to true for opening a new page |
| layout.orientation | the orientation of the DAG layout. It can be one of "left_right" for the left-right layout (viewed from the DAG root point; by default), "top_bottom" for the top-bottom layout, "bottom_top" for the bottom-top layout, and "right_left" for the right-left layout |
| node.info | tells the ontology term information used to label nodes. It can be one of "both" for using both of Term ID and Name (the first 15 characters; by default), "none" for no node labeling, "term_id" for using Term ID, "term_name" for using Term Name (the first 15 characters), and "full_term_name" for using the full Term Name |
| graph.node.attrs | a list of global node attributes. These node attributes will be changed globally. See 'Note' below for details on the attributes |
| graph.edge.attrs | a list of global edge attributes. These edge attributes will be changed globally. See 'Note' below for details on the attributes |
| node.attrs | a list of local edge attributes. These node attributes will be changed locally; as such, for each attribute, the input value must be a named vector (i.e. using Term ID as names). See 'Note' below for details on the attributes |

Value

An object of class 'Ragraph'

Note

A list of global node attributes used in "graph.node.attrs":

- "shape": the shape of the node: "circle", "rectangle", "rect", "box" and "ellipse"
- "fixedsize": the logical to use only width and height attributes. By default, it sets to true for not expanding for the width of the label
- "fillcolor": the background color of the node
- "color": the color for the node, corresponding to the outside edge of the node
- "fontcolor": the color for the node text/labelings
- "fontsize": the font size for the node text/labelings
- "height": the height (in inches) of the node: 0.5 by default
- "width": the width (in inches) of the node: 0.75 by default

- "style": the line style for the node: "solid", "dashed", "dotted", "invis" and "bold"

A list of global edge attributes used in "graph.edge.attrs":

- "color": the color of the edge: gray by default
- "weight": the weight of the edge: 1 by default
- "style": the line style for the edge: "solid", "dashed", "dotted", "invis" and "bold"

A list of local node attributes used in "node.attrs" (only those named Term IDs will be changed locally!):

- "label": a named vector specifying the node text/labelings
- "shape": a named vector specifying the shape of the node: "circle", "rectangle", "rect", "box" and "ellipse"
- "fixedsize": a named vector specifying whether it sets to true for not expanding for the width of the label
- "fillcolor": a named vector specifying the background color of the node
- "color": a named vector specifying the color for the node, corresponding to the outside edge of the node
- "fontcolor": a named vector specifying the color for the node text/labelings
- "fontsize": a named vector specifying the font size for the node text/labelings
- "height": a named vector specifying the height (in inches) of the node: 0.5 by default
- "width": a named vector specifying the width (in inches) of the node: 0.75 by default
- "style": a named vector specifying the line style for the node: "solid", "dashed", "dotted", "invis" and "bold"

See Also

[dcEnrichment](#), [dcRDataLoader](#), [dcConverter](#)

Examples

```
## Not run:
# 1) load SCOP.sf (as 'InfoDataFrame' object)
SCOP.sf <- dcRDataLoader('SCOP.sf')
# randomly select 20 domains
data <- sample(rowNames(SCOP.sf), 20)

# 2) perform enrichment analysis, producing an object of S4 class 'Eoutput'
eoutput <- dcEnrichment(data, domain="SCOP.sf", ontology="GOMF")
eoutput

# 3) visualise the top 10 significant terms
# color-coded according to 10-based negative logarithm of p-values
visEnrichment(eoutput)
# color-coded according to zscore
visEnrichment(eoutput, data.type='zscore')
```

```
# 4) visualise the top 5 significant terms in the ontology hierarchy
nodes_query <- names(sort(adjp(eoutput))[1:5])
visEnrichment(eoutput, nodes_query=nodes_query)
# change the frame color: highlight (framed in blue) nodes/terms in query
nodes.highlight <- rep("blue", length(nodes_query))
names(nodes.highlight) <- nodes_query
visEnrichment(eoutput, nodes_query=nodes_query,
node.attrs=list(color=nodes.highlight))

## End(Not run)
```

Index

*Topic **S4**

- AdjData-class, [3](#)
- Anno-class, [5](#)
- Anno-method, [7](#)
- AnnoData-class, [8](#)
- Cnetwork-class, [9](#)
- Cnetwork-method, [10](#)
- Coutput-class, [11](#)
- Coutput-method, [13](#)
- Dnetwork-class, [37](#)
- Dnetwork-method, [39](#)
- Eoutput-class, [40](#)
- Eoutput-method, [42](#)
- InfoDataFrame-class, [45](#)
- InfoDataFrame-method, [47](#)
- Onto-class, [52](#)
- Onto-method, [54](#)

*Topic **classes**

- AdjData-class, [3](#)
- Anno-class, [5](#)
- AnnoData-class, [8](#)
- Cnetwork-class, [9](#)
- Coutput-class, [11](#)
- Dnetwork-class, [37](#)
- Eoutput-class, [40](#)
- InfoDataFrame-class, [45](#)
- Onto-class, [52](#)

*Topic **datasets**

- Ancestral_domainome, [4](#)
- eTOL, [44](#)
- InterPro, [48](#)
- InterPro2GOBP, [49](#)
- InterPro2GOCC, [50](#)
- InterPro2GOMF, [51](#)
- onto.DO, [55](#)
- onto.GOBP, [56](#)
- onto.GOCC, [56](#)
- onto.GOMF, [57](#)
- onto.HPMI, [58](#)

- onto.HPON, [59](#)
- onto.HPPA, [60](#)
- onto.MP, [61](#)
- Pfam, [61](#)
- Pfam2GOBP, [62](#)
- Pfam2GOCC, [63](#)
- Pfam2GOMF, [64](#)
- Rfam, [65](#)
- Rfam2GOBP, [66](#)
- Rfam2GOCC, [67](#)
- Rfam2GOMF, [68](#)
- SCOP.fa, [69](#)
- SCOP.fa2DO, [70](#)
- SCOP.fa2GOBP, [71](#)
- SCOP.fa2GOCC, [72](#)
- SCOP.fa2GOMF, [73](#)
- SCOP.fa2HPMI, [74](#)
- SCOP.fa2HPON, [75](#)
- SCOP.fa2HPPA, [76](#)
- SCOP.fa2MP, [77](#)
- SCOP.sf, [78](#)
- SCOP.sf2DO, [79](#)
- SCOP.sf2GOBP, [80](#)
- SCOP.sf2GOCC, [81](#)
- SCOP.sf2GOMF, [82](#)
- SCOP.sf2HPMI, [83](#)
- SCOP.sf2HPON, [84](#)
- SCOP.sf2HPPA, [85](#)
- SCOP.sf2MP, [86](#)

*Topic **methods**

- Anno-method, [7](#)
- Cnetwork-method, [10](#)
- Coutput-method, [13](#)
- Dnetwork-method, [39](#)
- Eoutput-method, [42](#)
- InfoDataFrame-method, [47](#)
- Onto-method, [54](#)
- [,Anno,ANY,ANY,ANY-method
(Anno-method), [7](#)

- [,Anno-method (Anno-method), 7
- [,Cnetwork, ANY, ANY, ANY-method
(Cnetwork-method), 10
- [,Cnetwork-method (Cnetwork-method), 10
- [,Dnetwork, ANY, ANY, ANY-method
(Dnetwork-method), 39
- [,Dnetwork-method (Dnetwork-method), 39
- [,InfoDataFrame, ANY, ANY, ANY-method
(InfoDataFrame-method), 47
- [,InfoDataFrame-method
(InfoDataFrame-method), 47
- [,Onto, ANY, ANY, ANY-method
(Onto-method), 54
- [,Onto-method (Onto-method), 54
- AdjData, 9, 23, 37, 52, 55–61
- AdjData (AdjData-class), 3
- AdjData-class, 3
- adjMatrix (Onto-method), 54
- adjMatrix, Cnetwork-method
(Cnetwork-method), 10
- adjMatrix, Dnetwork-method
(Dnetwork-method), 39
- adjMatrix, Onto-method (Onto-method), 54
- adjp (Eoutput-method), 42
- adjp, Coutput-method (Coutput-method), 13
- adjp, Eoutput-method (Eoutput-method), 42
- Ancestral_domainome, 4
- Anno, 4, 15, 16, 20, 49–51, 63–65, 67–69,
71–78, 80–87
- Anno (Anno-class), 5
- Anno-class, 5
- Anno-method, 7
- AnnoData, 5
- AnnoData (AnnoData-class), 8
- annoData (Anno-method), 7
- annoData, Anno-method (Anno-method), 7
- AnnoData-class, 8
- Cnetwork, 12
- Cnetwork (Cnetwork-class), 9
- cnetwork (Coutput-method), 13
- cnetwork, Coutput-method
(Coutput-method), 13
- Cnetwork-class, 9
- Cnetwork-method, 10
- colNames (InfoDataFrame-method), 47
- colNames, InfoDataFrame-method
(InfoDataFrame-method), 47
- Coutput (Coutput-class), 11
- Coutput-class, 11
- Coutput-method, 13
- Data (InfoDataFrame-method), 47
- Data, InfoDataFrame-method
(InfoDataFrame-method), 47
- data.frame2InfoDataFrame
(InfoDataFrame-method), 47
- dcBuildAnno, 15
- dcBuildInfoDataFrame, 16
- dcBuildOnto, 17
- dcConverter, 18, 21, 23, 29, 36, 90
- dcDAGannotate, 20, 22, 23, 29, 36
- dcDAGdomainSim, 21, 21, 36, 37
- dcEnrichment, 21, 26, 34, 40, 41, 43, 90
- dcRDataLoader, 19, 21, 23, 29, 32, 36, 90
- dcRWRpipeline, 9, 11, 14, 34
- dData (Anno-method), 7
- dData, Anno-method (Anno-method), 7
- description (Dnetwork-method), 39
- description, Dnetwork-method
(Dnetwork-method), 39
- dgCMatrix2Anno (Anno-method), 7
- dgCMatrix2Cnetwork (Cnetwork-method), 10
- dgCMatrix2Dnetwork (Dnetwork-method), 39
- dgCMatrix2Onto (Onto-method), 54
- dim, Anno-method (Anno-method), 7
- dim, Cnetwork-method (Cnetwork-method),
10
- dim, Dnetwork-method (Dnetwork-method),
39
- dim, InfoDataFrame-method
(InfoDataFrame-method), 47
- dim, Onto-method (Onto-method), 54
- dimLabels (InfoDataFrame-method), 47
- dimLabels, InfoDataFrame-method
(InfoDataFrame-method), 47
- Dnetwork, 23, 35
- Dnetwork (Dnetwork-class), 37
- Dnetwork-class, 37
- Dnetwork-method, 39
- domainData (Anno-method), 7
- domainData, Anno-method (Anno-method), 7
- domainNames (Anno-method), 7
- domainNames, Anno-method (Anno-method), 7
- drop, 8, 11, 40, 48, 55
- Eoutput, 28, 88

- Eoutput (Eoutput-class), 40
- Eoutput-class, 40
- Eoutput-method, 42
- eTOL, 44

- id (Dnetwork-method), 39
- id,Dnetwork-method (Dnetwork-method), 39
- InfoDataFrame, 5, 9, 16, 17, 37, 48, 52, 55–62, 66, 70, 79
- InfoDataFrame (InfoDataFrame-class), 45
- InfoDataFrame-class, 45
- InfoDataFrame-method, 47
- InterPro, 48
- InterPro2GOBP, 49
- InterPro2GOCC, 50
- InterPro2GOMF, 51

- level (Dnetwork-method), 39
- level,Dnetwork-method (Dnetwork-method), 39

- matrix2Anno (Anno-method), 7
- matrix2Cnetwork (Cnetwork-method), 10
- matrix2Dnetwork (Dnetwork-method), 39
- matrix2Onto (Onto-method), 54

- ncol (InfoDataFrame-method), 47
- ncol,InfoDataFrame-method (InfoDataFrame-method), 47
- nInfo (Onto-method), 54
- nInfo,Cnetwork-method (Cnetwork-method), 10
- nInfo,Dnetwork-method (Dnetwork-method), 39
- nInfo,Onto-method (Onto-method), 54
- nodeInfo (Onto-method), 54
- nodeInfo,Cnetwork-method (Cnetwork-method), 10
- nodeInfo,Dnetwork-method (Dnetwork-method), 39
- nodeInfo,Onto-method (Onto-method), 54
- nodeName (Onto-method), 54
- nodeName,Cnetwork-method (Cnetwork-method), 10
- nodeName,Dnetwork-method (Dnetwork-method), 39
- nodeName,Onto-method (Onto-method), 54
- nrow (InfoDataFrame-method), 47
- nrow,InfoDataFrame-method (InfoDataFrame-method), 47

- Onto, 17, 18, 20, 22, 55–61
- Onto (Onto-class), 52
- Onto-class, 52
- Onto-method, 54
- onto.DO, 55
- onto.GOBP, 56
- onto.GOCC, 56
- onto.GOMF, 57
- onto.HPMI, 58
- onto.HPON, 59
- onto.HPPA, 60
- onto.MP, 61

- Pfam, 61
- Pfam2GOBP, 62
- Pfam2GOCC, 63
- Pfam2GOMF, 64
- pvalue (Eoutput-method), 42
- pvalue,Coutput-method (Coutput-method), 13
- pvalue,Eoutput-method (Eoutput-method), 42

- ratio (Coutput-method), 13
- ratio,Coutput-method (Coutput-method), 13

- Rfam, 65
- Rfam2GOBP, 66
- Rfam2GOCC, 67
- Rfam2GOMF, 68
- rownames (InfoDataFrame-method), 47
- rownames,InfoDataFrame-method (InfoDataFrame-method), 47

- SCOP.fa, 69
- SCOP.fa2DO, 70
- SCOP.fa2GOBP, 71
- SCOP.fa2GOCC, 72
- SCOP.fa2GOMF, 73
- SCOP.fa2HPMI, 74
- SCOP.fa2HPON, 75
- SCOP.fa2HPPA, 76
- SCOP.fa2MP, 77
- SCOP.sf, 78
- SCOP.sf2DO, 79
- SCOP.sf2GOBP, 80
- SCOP.sf2GOCC, 81
- SCOP.sf2GOMF, 82
- SCOP.sf2HPMI, 83

SCOP.sf2HPON, [84](#)
SCOP.sf2HPPA, [85](#)
SCOP.sf2MP, [86](#)
show,Anno-method (Anno-method), [7](#)
show,Cnetwork-method (Cnetwork-method),
[10](#)
show,Coutput-method (Coutput-method), [13](#)
show,Dnetwork-method (Dnetwork-method),
[39](#)
show,Eoutput-method (Eoutput-method), [42](#)
show,InfoDataFrame-method
(InfoDataFrame-method), [47](#)
show,Onto-method (Onto-method), [54](#)

tData (Anno-method), [7](#)
tData,Anno-method (Anno-method), [7](#)
term_distance (Onto-method), [54](#)
term_distance,Onto-method
(Onto-method), [54](#)
term_id (Onto-method), [54](#)
term_id,Onto-method (Onto-method), [54](#)
term_name (Onto-method), [54](#)
term_name,Onto-method (Onto-method), [54](#)
term_namespace (Onto-method), [54](#)
term_namespace,Onto-method
(Onto-method), [54](#)
termData (Anno-method), [7](#)
termData,Anno-method (Anno-method), [7](#)
termNames (Anno-method), [7](#)
termNames,Anno-method (Anno-method), [7](#)

view (Eoutput-method), [42](#)
view,Eoutput-method (Eoutput-method), [42](#)
visEnrichment, [29](#), [87](#)

write (Eoutput-method), [42](#)
write,Coutput-method (Coutput-method),
[13](#)
write,Eoutput-method (Eoutput-method),
[42](#)

zscore (Eoutput-method), [42](#)
zscore,Coutput-method (Coutput-method),
[13](#)
zscore,Eoutput-method (Eoutput-method),
[42](#)