

Package ‘ProCoNA’

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Type Package

Title Protein co-expression network analysis (ProCoNA).

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Description Protein co-expression network construction using peptide level data, with statistical analysis. (Journal of Clinical Bioinformatics 2013, 3:11 doi:10.1186/2043-9113-3-11)

License GPL (>= 2)

LazyLoad yes

Depends R (>= 2.10), methods, WGCNA, MSnbase

Imports BiocGenerics, GOstats

Suggests RUnit

biocViews GraphAndNetwork, Software, Proteomics

R topics documented:

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procona-package *Peptide co-expression network construction.*

Description

Peptide co-expression network construction, analysis, and visualization.

Details

Package:	procona
Type:	Package
Title:	Peptide co-expression network construction.
Version:	0.13
Date:	2011-08-10
Author:	David L Gibbs
Maintainer:	David Gibbs <gibbsd@ohsu.edu>
License:	GPLv3
LazyLoad:	yes
Depends:	WGCNA, GOstats, multicore

Author(s)

David L Gibbs

Accessors

Accessors for the proconaNet S4 class

Description

Accessor functions allow access to the object data.

Methods

TOM: The topological overlap matrix or TOM. "matrix"

adj: The adjacency matrix. "matrix"

networkName: A name describing the data or experiment used to build the network. "character"

samples: The names of samples used in building the network. "character"

peptides: The names of peptides used in the network, also the node names. "character"

pepTree: The network dendrogram. "hclust"

dynamicColors: The module labels on each node (or peptide). "numeric"

MEs: The module eigenvectors (or eigen-peptides). "data.frame"

mergedMEs: The module eigenvectors after merging similar modules. "data.frame"

mergedColors: The module labels after merging similar modules. "numeric"

colorOrder: Modules are ordered by size, these labels correspond to that order. "character"

power: The soft thresholding power used in scaling the adjacency matrix. "numeric"

networkType: Either a signed or unsigned network regarding the method used in computing the initial correlations between nodes. "character"

permtest: The results of the permutation test on significance of topological overlap within modules. "matrix"

proconaVersion: Returns the version number of the software that built the object. "character"

Author(s)

David L Gibbs

Examples

```
data(ProCoNA_Data)
tomMatrix <- TOM(net1)
```

 bootstrapProconaNetwork

bootstrapProconaNetwork

Description

This function returns a peptide co-expression network object based on a bootstrapped correlation matrix.

Usage

```
bootstrapProconaNetwork(networkName = "bootstrap procona", pepdat = NULL,
  pow = NULL, powMax = 20, networkType = "signed", scaleFreeThreshold = 0.8,
  deepSplit = 2, minModuleSize = 30, mergeThreshold = 0.1,
  clusterType = "average", pamRespectsDendro = T, performT0Permtest = TRUE,
  toPermTestPermutates = 100, bootstrapThreshold = 1e-04)
```

Arguments

networkName	Name of this network
pepdat	This variable is the data set with rows as samples and cols as peptides
pow	The scaling power, NULL if unknown
powMax	The maximum power to be searched.
networkType	Whether the sign is considered in constructing adjacency and TOM
scaleFreeThreshold	The threshold for fitting to scale-free topology.. will use closest power.
deepSplit	Course grain control of module size
minModuleSize	The minimum module size allowed
mergeThreshold	Below this threshold, modules are merged.
clusterType	Clustering option
pamRespectsDendro	When cutting the dendrogram, pay attention to branch membership.
performT0Permtest	Performs permutation testing on modules
toPermTestPermutates	Number of permutations to do.
bootstrapThreshold	When to stop resampling...

Value

returns the procona network object

Author(s)

David L Gibbs

Examples

```
data(ProCoNA_Data)
net <- bootstrapProconaNetwork("peptide network", peptideData,
performT0Permtest=FALSE, bootstrapThreshold=0.1)
```

buildProconaNetwork *buildProconaNetwork*

Description

This function returns a peptide co-expression network object.

Usage

```
buildProconaNetwork(networkName = "ProCoNA", pepdat, pow=1,
powMax = 20, networkType = "signed", pearson = FALSE, scaleFreeThreshold = 0.8,
deepSplit = 2, minModuleSize = 30, mergeThreshold = 0.1,
clusterType = "average", pamRespectsDendro = TRUE, performT0Permtest = TRUE,
toPermTestPermutates = 100)
```

Arguments

networkName	Name of this network
pepdat	This variable is the data set with rows as samples and cols as peptides
pow	The scaling power, NULL if unknown
powMax	The maximum power to be searched.
networkType	Should the sign be considered in constructing adjacency and TOM ("signed" or "unsigned")
pearson	use Pearson's cor or the robust bi-weight correlation
scaleFreeThreshold	The threshold for fitting to scale-free topology.. will use closest power.
deepSplit	Course grain control of module size
minModuleSize	The minimum module size allowed
mergeThreshold	Below this threshold, modules are merged.
clusterType	Clustering option
pamRespectsDendro	When cutting the dendrogram, pay attention to branch membership.
performT0Permtest	Performs permutation testing on modules
toPermTestPermutates	Number of permutations to do.

Details

The procona network object contains a number of slots which store information relevant to the construction of the network. Accessor functions provide direct access to the slots. See `getSlots("proconaNet")` for a complete list.

Value

returns the procona network object

Author(s)

David L Gibbs

Examples

```
data(ProCoNA_Data)
net <- buildProconaNetwork("peptide network", peptideData)
```

c2i

c2i

Description

coordinates to index

Usage

```
c2i(nrows, x, y)
```

Arguments

nrows	number of rows in the matrix
x	the row coordinate
y	the col coordinate

Value

the index into the matrix

Author(s)

David L Gibbs

```
compareNetworksWithFishersExactTest  
    compareNetworksWithFishersExactTest
```

Description

Fisher's exact test is used pairwise on modules to compare two networks. The arguments to Fisher's exact test are given below.

n == number of entities in the network

m == number of entities in intersection of two modules

d1 == number of entities in module A but not in module B

d2 == number of entities in module B but not in module A

2x2 matrix for the test is then: m d1 d2 n-d1-d2-m

Usage

```
compareNetworksWithFishersExactTest(peps1, peps2, colors1, colors2,  
    title = "", net1label = "", net2label = "")
```

Arguments

peps1	Nodes in network 1, character vector
peps2	Nodes in network 2, character vector
colors1	modules for net 1
colors2	modules for net 2
title	Plot title
net1label	xlabel
net2label	ylabel

Value

Returns fishers exact test -log pvalues and overlap matrix showing the number of shared members for each pair of modules.

Author(s)

David L Gibbs

Examples

```
## Not run:
data(ProCoNA_Data)
#net1 <- buildProconaNetwork("peptide network", peptideData, pow=12)
#net2 <- buildProconaNetwork("peptide network", peptideData, pow=6)
compareNetworksWithFishersExactTest(peptides(net1), peptides(net2),
mergedColors(net1), mergedColors(net2), "network comparison", "net1", "net2")

## End(Not run)
```

```
compareNetworksWithFishersExactTestProcona
      compareNetworksWithFishersExactTestProcona
```

Description

Convenience function for calling the `compareNetworksWithFishersExactTest` using only two procona objects.

Usage

```
compareNetworksWithFishersExactTestProcona(net1, net2,
      title)
```

Arguments

<code>net1</code>	procona object for network 1
<code>net2</code>	procona object for network 2
<code>title</code>	plot title

Value

Returns a list of fisher -log pvalues, and overlaps between modules.

Author(s)

David L Gibbs

Examples

```
## Not run:
data(ProCoNA_Data)
#net1 <- buildProconaNetwork("peptide network", peptideData)
#net2 <- buildProconaNetwork("peptide network", peptideData)
compareNetworksWithFishersExactTestProcona(net1, net2, "new comparison")

## End(Not run)
```

`corBootstrap`*corBootstrap*

Description

Boostraps a correlation matrix. In order to bootstrap a large correlation matrix, several thousand samplings may be necessary. To avoid storing thousands of matrices, a running mean is kept for each pairwise correlation. In addition, a running standard deviation is computed so that for each pairwise correlation, we can estimate the distribution of values across resamplings. After each resampling, a new correlation matrix is computed. A difference is taken between this new matrix and the running mean. If all differences are less than the specified threshold, then the bootstrapped matrix has converged to a final state.

Usage

```
corBootstrap(dataMatrix, networkType = "signed", threshold = 1e-04,  
             tmpSaveFile = TRUE)
```

Arguments

<code>dataMatrix</code>	Matrix with samples in rows and peptides (or other data type) in columns.
<code>networkType</code>	Whether the sign is considered in constructing adjacency and TOM
<code>threshold</code>	Maximum difference allowed between running mean bootstrap correlation matrix, and new resampled cor matrix. Defines how soon we consider the bootstrap to have converged.
<code>tmpSaveFile</code>	Should temporary saves be done?

Value

Returns a list of the bootstrapped matrix, standard deviation matrix, and the number of resamplings done.

Author(s)

David L Gibbs

Examples

```
data(ProCoNA_Data)  
x <- peptideData[,1:10]  
y <- corBootstrap(dataMatrix=x, networkType="unsigned", threshold=0.1, tmpSaveFile=FALSE)
```

correlationWithPhenotypesHeatMap
correlationWithPhenotypesHeatMap

Description

Plots a heatmap showing the Pearson correlation of modules with phenotypes.

Usage

```
correlationWithPhenotypesHeatMap(net, phenotypes, modules,  
  plotName, title, textSize)
```

Arguments

net	The ProCoNA network object.
phenotypes	Matrix of phenotypic traits, can include character strings (converted to factors).
modules	Vector of modules to plot. Default is all modules.
plotName	Name of the saved plot, NULL to show on screen.
title	Plot title.
textSize	The font size of the correlations shown in each module-phenotype pair.

Value

the module eigenvector correlations

Author(s)

David L Gibbs

Examples

```
data(ProCoNA_Data)  
#net1 <- buildProconaNetwork("pepnet", peptideData, pow=12)  
n <- length(samples(net1))  
phenotypes <- matrix(rnorm(10*n), nrow=60)  
moduleCors <- correlationWithPhenotypesHeatMap(net1, phenotypes, modules = 1:7,  
  plotName = "Phenotype Associations", title = "Module-trait relationships", textSize = 0.5)
```

getFisherMatrix	<i>getFisherMatrix</i>
-----------------	------------------------

Description

Fisher's exact test pairwise on modules.

Usage

```
getFisherMatrix(peps1, peps2, colors1, colors2)
```

Arguments

peps1	Names of entities in the network (nodes of network 1)
peps2	Names of entities in the network (nodes of network 2)
colors1	the module assignments for network 1
colors2	the module assignments for network 2

Value

Returns the fisher test pvalues and count of overlapping peptides.

Author(s)

David L Gibbs

Examples

```
data(ProCoNA_Data)
#net1 <- buildProconaNetwork("peptide network", peptideData, pow=12)
#net2 <- buildProconaNetwork("peptide network", peptideData + 0.3*rnorm(length(peptideData)), pow=12)
getFisherMatrix(peptides(net1), peptides(net2), mergedColors(net1), mergedColors(net2))
```

getPeptideNAs	<i>getPeptideNAs</i>
---------------	----------------------

Description

This function returns the number of NAs for each peptide.

Usage

```
getPeptideNAs(pepdat)
```

Arguments

pepdat the peptide data.

Value

returns a list of counts of NAs for each peptide.

Author(s)

David L Gibbs

goStatTest

goStatTest

Description

Wrapper function to run the hyperGTest from package GOstats, after mapping each peptide to an entrez ID.

Usage

```
goStatTest(pnet, module, pepinfo, pepColName, protColName, universe,
           onto, annot, pvalue, cond)
```

Arguments

pnet	Procona network object.
module	Module of interest (numeric)
pepinfo	The mass tag info, mapping peptides to proteins.
pepColName	Column name in mass tag info for peptides
protColName	Column name in mass tag info for proteins
universe	Table mapping protein IDs to entrez IDs
onto	The ontology category (bp etc)..
annot	The annotation database to use
pvalue	pvalue cutoff
cond	conditional parameter, see GOstats.

Value

Returns the results of the hyper geometric test.

Author(s)

David L Gibbs

Examples

```
## Not run:
data(ProCoNA_Data)
#net1 <- buildProconaNetwork("peptide network", peptideData, pow=12)
goStatTest(net1, 1, masstagdb, "Mass_Tag_ID", "Reference", universe, "BP", "org.Mm.eg.db", 0.005, FALSE)

## End(Not run)
```

hclust-class	Class "hclust"
--------------	----------------

Description

From the OneHandClapping package. Thanks! Dummy class to permit object of S3 class hclust in S4 class definition of Screening

Objects from the Class

Objects can be created by calls of the form `new("hclust", ...)`.

Slots

.Data: Object of class "list" ~~

Extends

Class "list", from data part. Class "vector", by class "list", distance 2.

Methods

No methods defined with class "hclust" in the signature.

Warning

This class is just defined as a dummy class. No objects should be instantiated.

Note

This class is just defined as a dummy class. No objects should be instantiated.

Examples

```
showClass("hclust")
```

*i2c**i2c*

Description

Index to coordinates

Usage

`i2c(nrows, i)`

Arguments

<code>nrows</code>	number of rows in the matrix
<code>i</code>	the index into the matrix

Value

the row col coordinates into the matrix

Author(s)

David L Gibbs

*i2col**i2col*

Description

index to column

Usage

`i2col(nrows, i)`

Arguments

<code>nrows</code>	number of rows in matrix
<code>i</code>	the index

Value

returns the column of the matrix

Author(s)

David L Gibbs

MMvsPS

Module members vs Peptide Significance

Description

Plots the module membership (correlation to eigenvector) against the peptide significance (correlation to phenotype) for a given trait and module

Usage

```
MMvsPS(pnet, pepdat, phenoVec, mod)
```

Arguments

pnet	The procona network
pepdat	the peptide data, with rows as samples and columns as peptides
phenoVec	the phenotypic trait, vector
mod	the module of interest

Value

returns a list of module memberships and peptide significances.

Author(s)

David L Gibbs

Examples

```
data(ProCoNA_Data)
#net1 <- buildProconaNetwork("peptide network", peptideData, pow=13)
MMvsPS(net1, peptideData, phenotypes[,5], 1)
```

MMvsPSallModules

MMvsPSallModules

Description

Call MMvsPS, producing plots for all modules.

Usage

```
MMvsPSallModules(net, peptable, phenoVec, prefixName)
```

Arguments

net	The procona network object
peptable	The peptide data
phenoVec	The phenotypic trait, as a numeric vector
prefixName	The plot files prefix name. Writes pdfs.

Value

nothing returned

Author(s)

David L Gibbs

Examples

```
## Not run:  
# This function outputs a set of pdfs.  
data(ProCoNA_Data)  
#net1 <- buildProconaNetwork("peptide network", peptideData, pow=13)  
MMvsPSallModules(net1, peptideData, phenotypes[,5], 1)  
  
## End(Not run)
```

moduleMemberCorrelations

moduleMemberCorrelations

Description

Computes the relation between peptides and eigenvector summaries and also peptides and phenotypes.

Usage

```
moduleMemberCorrelations(pnet, pepdat, phenotypes)
```

Arguments

pnet	The peptide net object
pepdat	The peptide data matrix
phenotypes	The matrix of traits

Value

Matrix of Pearson correlations with peptides in rows.

Author(s)

David L Gibbs

Examples

```

data(ProCoNA_Data)
#net1 <- buildProconaNetwork("peptide network", peptideData)
n <- length(samples(net1))
phenotypes <- matrix(rnorm(10*n), nrow=60)
pepcor <- moduleMemberCorrelations(net1, peptideData, phenotypes)

# To plot the heatmap:
# moduleCors <- correlationWithPhenotypesHeatMap(net1, phenotypes, modules = 1:5,
#   plot = NULL, title = "Module-trait relationships", textSize = 0.5)

#####
# quick function to write out the tables for specific modules.
#moduleData <- function(pepnet, pepcors, module, pepinfo, fileprefix) {
#   moduleX <- pepnet@peptides[which(pepnet@mergedColors==module)]
#   moduleInfo <- pepinfo[which(pepinfo$Mass_Tag_ID %in% moduleX),]
#   moduleCors <- pepcors[which(pepcors$Module==module),]
#   corname <- paste(fileprefix, "_correlations.csv", sep="")
#   write.table(moduleCors, file=corname, sep=",", row.names=F)
#   infoname <- paste(fileprefix, "_peptide_info.csv", sep="")
#   write.table(moduleInfo, file=infoname, sep=",", row.names=F)
#}
#####

# WRITE OUT A TABLE WITH THE BELOW FUNCTION CALL :)#
# moduleData(peptideNetwork, pepcor, 1, masstagdb, "Module_1")

```

modulePhenotypeCorrelations

modulePhenotypeCorrelations

Description

Computes the relation between the modules and the phenotypes.

Usage

```
modulePhenotypeCorrelations(pnet, phenotypes)
```

Arguments

pnet	The peptide net object
phenotypes	The matrix of traits

Value

returns a matrix of correlations between modules and phenotypes.

Author(s)

David L Gibbs

Examples

```
data(ProCoNA_Data)
#net1 <- buildProconaNetwork("peptide network", peptideData, pow=13)
n <- length(samples(net1))
phenotypes <- matrix(rnorm(10*n), nrow=60)
m <- modulePhenotypeCorrelations(net1, phenotypes)

# To plot the heatmap:
# moduleCors <- correlationWithPhenotypesHeatMap(net1, phenotypes, modules = 1:5,
#   plot = NULL, title = "Module-trait relationships", textSize = 0.5)
```

orderMatrixIndex

orderMatrixIndex

Description

Order the the matrix by upper diag in a greedy fashion

Usage

```
orderMatrixIndex(mat)
```

Arguments

mat A matrix

Value

returns a matrix in order of greatest in upper diagonal direction.

Author(s)

David L Gibbs

```
peptideConnectivityTest  
    peptideConnectivityTest
```

Description

This function will compare the connectivity between peptides mapped to a given protein, against a randomly drawn, similarly sized, selection of peptides. The hypothesis is that peptides from a given protein should be more connected than random.

Usage

```
peptideConnectivityTest(pnet, pepInfo, pepCol, protCol, repsPerProt)
```

Arguments

pnet	The peptide net object
pepInfo	The peptide information table, mapping peptides to proteins
pepCol	The string identifying the column in the pepInfo table with peptide ID
protCol	String identifying column in pepInfo with Protein ID.
repsPerProt	number of repetitions for the null

Value

Returns a list of the connected peptides and the random samples.

Author(s)

David L Gibbs

Examples

```
data(ProCoNA_Data)  
#net1 <- buildProconaNetwork("peptide network", peptideData, pow=12)  
p <- peptideConnectivityTest(net1, masstagdb, "Mass_Tag_ID", "Reference", 200)
```

peptideCorrelationTest

peptideCorrelationTest

Description

Take the data, and a mapping of peptides to proteins, and compute the mean correlation between peptides linked to a given protein. Compare a similar number of random correlations.

Usage

```
peptideCorrelationTest(dat, pepinfo, pepCol, protCol)
```

Arguments

dat	The data with samples as rows and peptides as columns
pepinfo	The mapping of peptides to proteins as a data frame
pepCol	The column name of peptide info table containing peptide IDs
protCol	The column name of pepinfo info table containing protein IDs

Value

return a t-test comparing protein correlations to random correlations.

Author(s)

David L Gibbs

Examples

```
data(ProCoNA_Data)
net1 <- buildProconaNetwork("peptide network", peptideData, pow=12)
peptideCorrelationTest(peptideData, masstagdb, "Mass_Tag_ID", "Reference")
```

plotNet

plotNet

Description

Plots the dendrogram and module colors. See ?plotDendroAndColors

Usage

```
plotNet(object)
```

Arguments

object The procona network object.

Value

None.

Author(s)

David L Gibbs

Examples

```
data(ProCoNA_Data)
#net1 <- buildProconaNetwork("peptide network", peptideData)
plotNet(net1)
```

ppiPermTest

ppiPermTest

Description

Performs a permutation test for enrichment of PPI edges given a database. Peptides are selected from each module and mapped to potential protein parents in the mass tag database. We check if these proteins are found in the PPI network, and record any edges between them. This is compared to edges found using randomly selected proteins (taken from the mass tag database). A p-value is computed as the number of times the randomly sampled proteins incurred more edges than the observed proteins, divided by the number of iterations.

Usage

```
ppiPermTest(pnet, pepdat, pepinfo, pepColName, pi_colName, pi_edges,
            threshold, iterations)
```

Arguments

pnet	procona network object
pepdat	the data matrix with peptides as columns.
pepinfo	Maps peptides to proteins ... same format as in ppiTable
pepColName	The column in pepinfo with peptide IDs... as in pepdat (the peptide data matrix)
pi_colName	The column in pepinfo that maps peptides to unit found in pi_edges
pi_edges	Must be two columns A-B ... sort out evidence levels (in vivo or in vitro) in advance
threshold	Minimum peptide correlation with module eigenvector.
iterations	Number of repetitions

Value

returns list of test results.

Author(s)

David L Gibbs

Examples

```
data(ProCoNA_Data)
#net1 <- buildProconaNetwork("peptide network", peptideData, pow=12)
ppis <- data.frame(A=sample(masstagdb$Reference, 50), B=sample(masstagdb$Reference, 50))
ppiPermTest(net1, peptideData, masstagdb, "Mass_Tag_ID", "Reference", ppis, 0.33, 100)
```

printNet

printNet

Description

Prints general information about the network object.

Usage

```
printNet(object)
```

Arguments

object The procona network object.

Value

None.

Author(s)

David L Gibbs

Examples

```
## Not run:
data(ProCoNA_Data)
net1 <- buildProconaNetwork("peptide network", peptideData)
printNet(net1)

## End(Not run)
```

 ProCoNA-Data

A simulated mass tag data base

Description

The mass tag database, which would be used to identify peptides, simply maps peptide IDs to peptide sequences and protein matches.

This simulated peptide dataset was generated using OpenMS's MSSimulator. A set of proteins was randomly sampled, and used to generate a likely set of observed peptides. Then data for a co-expression network was simulated with WGCNA's simulation function, and columns were named with simulated peptides.

The mass tag database, which would be used to identify peptides, simply maps peptide IDs to peptide sequences and protein matches. This represents a mapping to Entrez IDs.

The matrix annotates the biological samples according to ... phenotypic observations!

The two network objects are included to avoid rebuilding them in the other man page examples.

 proconaNet-class

proconaNet S4 class

Description

The main ProCoNA object - holder of data.

Objects from the Class

Objects can be created by calls of the form `new(proconaNet ...)`

Slots

networkName: A name describing the data or experiment used to build the network. "character"

samples: The names of samples used in building the network. "character"

adj: The adjacency matrix. "matrix"

TOM: The topological overlap matrix or TOM. "matrix"

peptides: The names of peptides used in the network, also the node names. "character"

pepTree: The network dendrogram. "hclust"

dynamicColors: The module labels on each node (or peptide). "numeric"

MEs: The module eigenvectors (or eigen-peptides). "data.frame"

mergedMEs: The module eigenvectors after merging similar modules. "data.frame"

mergedColors: The module labels after merging similar modules. "numeric"

colorOrder: Modules are ordered by size, these labels correspond to that order. "character"

power: The soft thresholding power used in scaling the adjacency matrix. "numeric"

networkType: Either a signed or unsigned network regarding the method used in computing the initial correlations between nodes. "character"

permtest: The results of the permutation test on significance of topological overlap within modules. "matrix"

proconaVersion: Returns the version number of the software that built the object. "character"

Methods

show signature(x = "proconaNet"): Shows info about the network.

print signature(x = "proconaNet"): Prints info about the network.

Author(s)

David L Gibbs

proconaVersionFun *Procona Software Version*

Description

Returns the current version of the software.

Usage

```
proconaVersionFun()
```

Value

returns the version

Author(s)

David L Gibbs

runningStats	<i>runningStats</i>
--------------	---------------------

Description

Computing the running mean and variance

Usage

```
runningStats(newMat, runningMean, Mk1, Sk1, k)
```

Arguments

newMat	The matrix from resampled data
runningMean	The running mean matrix
Mk1	Matrix used in calculation of mean
Sk1	Matrix used in calculation of sd
k	Current resampling iteration

Value

returns the list of runningMean, runningSD, Mk, Sk

Author(s)

David L Gibbs

subsetModCors	<i>subsetModCors</i>
---------------	----------------------

Description

subsets the module-phenotype correlation matrix which has funny rownames

Usage

```
subsetModCors(modCors, modules)
```

Arguments

modCors	The matrix of module-phenotype correlations
modules	Which modules are desired.

Author(s)

David L Gibbs

subsetPeptideData	<i>subsetPeptideData</i>
-------------------	--------------------------

Description

Given a matrix of peptide data, omit columns with excess missing data, specified by NAs.

Usage

```
subsetPeptideData(pepdat, numNAsAllowed = NULL, percentageNAsAllowed = 0.05)
```

Arguments

pepdat	The peptide matrix, with peptides in columns and samples in rows.
numNAsAllowed	The maximum count of missing values for each peptide (counts NAs).
percentageNAsAllowed	The percentage of missing data allowed for each peptide over samples.

Value

Returns a matrix.

Author(s)

David L Gibbs

Examples

```
data(ProCoNA_Data)
subsetPeptideData(peptideData, percentageNAsAllowed=0.2)
```

toPermTest	<i>toPermTest</i>
------------	-------------------

Description

Uses the procona network object, the data with peptides as columns, samples in rows. And the power that the net was built at the number of permutations to do... Modules are permuted and mean topological overlap is recorded, constructing the null. The number of random permutations with mean TO greater than observed provides the p-value.

Usage

```
toPermTest(pnet, numPermutates)
```

Arguments

`pnet` ProCoNA network object.
`numPermutates` The number of permutations to perform

Value

returns the network obj with the perm test

Author(s)

David L Gibbs

Examples

```
data(ProCoNA_Data)
#net1 <- buildProconaNetwork("peptide network", peptideData, pow=12)
toPermTest(net1, 100)
```

<code>utri</code>	<i>utri</i>
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Description

The upper triangle of a matrix

Usage

```
utri(mat)
```

Arguments

`mat` A matrix

Value

Returns a vector

Author(s)

David L Gibbs

Examples

```
m <- matrix(rnorm(9), nrow=3, ncol=3)
utri(m)
```

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