

Package ‘multiWGCNA’

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Description An R package for deeping mining gene co-expression networks in multi-trait expression data. Provides functions for analyzing, comparing, and visualizing WGCNA networks across conditions. multiWGCNA was designed to handle the common case where there are multiple biologically meaningful sample traits, such as disease vs wildtype across development or anatomical region.

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multiWGCNA-package	<i>multiWGCNA: multiWGCNA</i>
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Description

An R package for deeping mining gene co-expression networks in multi-trait expression data. Provides functions for analyzing, comparing, and visualizing WGCNA networks across conditions. multiWGCNA was designed to handle the common case where there are multiple biologically meaningful sample traits, such as disease vs wildtype across development or anatomical region.

Author(s)

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- Brent Fogel [contributor]

bidirectionalBestMatches	<i>Best matching modules</i>
--------------------------	------------------------------

Description

Find all the modules from dataset1 that have a best match to a module in dataset2 if that module in dataset2 is also a best match to the module in dataset1

Usage

```
bidirectionalBestMatches(comparisonList, plot = TRUE)
```

Arguments

comparisonList	a list with an elemnt "overlap", which is a data.frame resulting from a call to computeOverlapsFromWGCNA
plot	whether to generate a heatmap; default is TRUE

Value

A ggplot object

Author(s)

Dario Tommasini

Examples

```
library(ExperimentHub)
eh = ExperimentHub()
eh_query = query(eh, c("multiWGCNAdata"))
astrocyte_networks = eh_query[["EH8222"]]
comparisonList = list()
comparisonList$overlaps = computeOverlapsFromWGCNA(astrocyte_networks$EAE, astrocyte_networks$WT)
bidirectionalBestMatches(comparisonList)
```

BuildTOMFlowDF

BuildTOMFlowDF

Description

Preprocess for plotting a sankey flow diagram showing the movement of genes from one WGCNA to another WGCNA. Uses the flashClust framework.

Usage

```
BuildTOMFlowDF(WGCNAlist, networks, toms, genes_to_label, method = "average")
```

Arguments

WGCNAlist	list of WGCNA objects
networks	list of network names of length 2
toms	a list of TOM distance objects of length 2
genes_to_label	genes to label across two networks
method	linkage method to pass to flashClust for clustering, default is average linkage

Value

a data.frame

Author(s)

Dario Tommasini, Xinye Li

cleanDatExpr	<i>cleanDatExpr</i>
--------------	---------------------

Description

A function that converts a data.frame where row 1 is gene symbols to a numeric matrix where columns are genes and rows are samples for compatibility with most WGCNA functions.

Usage

```
cleanDatExpr(datExpr, checkGenesSamples = FALSE)
```

Arguments

datExpr	a data.frame where columns are samples and rows are samples and the gene symbols are in the first row
checkGenesSamples	call the WGCNA function checkGenesSamples?

Value

Returns a datExpr with rows as samples and columns as genes

Author(s)

Dario Tommasini

Examples

```
library(ExperimentHub)
eh = ExperimentHub()
eh_query = query(eh, c("multiWGCNAdata"))
astrocyte_se = eh_query[["EH8223"]]
datExpr = data.frame(X = rownames(assays(astrocyte_se)[[1]]), assays(astrocyte_se)[[1]])
cleanDatExpr(datExpr)
```

coexpressionLineGraph	<i>Coexpression Line Graph</i>
-----------------------	--------------------------------

Description

Plots a line graph showing the co-expression of selected genes across samples

Usage

```
coexpressionLineGraph(datExpr, splitBy = 1, fontSize = 2.15, colors = NULL)
```

Arguments

<code>datExpr</code>	a data.frame with genes as rows and samples as columns
<code>splitBy</code>	how much to split genes by on line graph
<code>fontSize</code>	the font size of the gene labels
<code>colors</code>	a vector of colors; default is random colors generated by colors function

Value

a ggplot object

Author(s)

Dario Tommasini

Examples

```
library(ExperimentHub)
eh = ExperimentHub()
eh_query = query(eh, c("multiWGCNAdata"))
astrocyte_networks = eh_query[["EH8222"]]
datExpr = GetDatExpr(astrocyte_networks[[1]],
  genes = topNGenes(astrocyte_networks$EAE, "EAE_015", 20))
coexpressionLineGraph(datExpr) +
  geom_vline(xintercept = 20.5, linetype='dashed')
```

`computeOverlapsFromWGCNA`

computeOverlapsFromWGCNA

Description

Computes overlap between the modules of two objects of class WGCNA

Usage

```
computeOverlapsFromWGCNA(dataset1, dataset2)
```

Arguments

<code>dataset1</code>	an object of class WGCNA to compare with dataset2
<code>dataset2</code>	an object of class WGCNA to compare with dataset1

Value

Returns a data.frame showing the overlap results for modules from dataset1 with dataset2

Author(s)

Dario Tommasini

Examples

```
library(ExperimentHub)
eh = ExperimentHub()
eh_query = query(eh, c("multiWGCNAdata"))
astrocyte_networks = eh_query[["EH8222"]]
computeOverlapsFromWGCNA(astrocyte_networks$EAE, astrocyte_networks$WT)
```

constructNetworks	<i>constructNetworks: Construct all the weighted gene correlation networks</i>
-------------------	--

Description

A high level function that returns all networks possible for a given experimental design

Usage

```
constructNetworks(
  datExpr,
  sampleTable,
  conditions1,
  conditions2,
  write = FALSE,
  alphaLevel = 0.05,
  plot = FALSE,
  detectNumbers = TRUE,
  ...
)
```

Arguments

datExpr	either a SummarizedExperiment object or data.frame with genes are rows and samples as columns
sampleTable	data.frame with sample names in first column and sample traits in the second and third column. First column should be called "Sample"
conditions1	first design conditions, ie healthy/disease
conditions2	second design conditions, ie frontal lobe/temporal lobe
write	write results out to files?
alphaLevel	significance value passed to findBestTrait function, default is 0.05
plot	plot modules? Default is false
detectNumbers	passed to makeTraitTable2; if you have any numeric traits, make sure this is TRUE so that these get detected.
...	Arguments to pass to blockwiseModules function

Value

A list of WGCNA objects, ie level one, two, and three networks.

Author(s)

Dario Tommasini

Examples

```
library(ExperimentHub)
eh = ExperimentHub()
eh_query = query(eh, c("multiWGCNAdata"))
autism_se = eh_query[["EH8219"]]
set.seed(1)
autism_se = autism_se[sample(rownames(autism_se), 500),]
sampleTable = colData(autism_se)
conditions1 = unique(sampleTable[,2])
conditions2 = unique(sampleTable[,3])
autism_networks = constructNetworks(autism_se, sampleTable, conditions1[[1]], conditions2[[1]],
  networkType = "signed", TOMType = "unsigned",
  power = 10, minModuleSize = 100, maxBlockSize = 25000,
  reassignThreshold = 0, minKMEtoStay = 0, mergeCutHeight = 0,
  numericLabels = TRUE, pamRespectsDendro = FALSE,
  deepSplit = 4, verbose = 3)
autism_networks[["combined"]]
```

diffCoexpression

Differential co-expresison analysis

Description

Performs a differential co-expression ananlysis given an expression data.frame and a conditions vector

Usage

```
diffCoexpression(
  datExpr,
  conditions,
  geneList = NULL,
  plot = FALSE,
  method = c("pearson", "spearman"),
  removeFreeNodes = TRUE,
  labelSize = 0.5,
  labelDist = 0,
  shape = "circle",
  degreeForSize = FALSE,
```

```

    label = FALSE,
    onlyPositive = FALSE,
    z.threshold = NULL,
    FDR.threshold = 0.05,
    nodeSize = 3
  )

```

Arguments

<code>datExpr</code>	a data.frame containing expression values
<code>conditions</code>	a vector containing conditions for the samples
<code>geneList</code>	vector of genes, will use all genes if NULL (default)
<code>plot</code>	plot a network?
<code>method</code>	either "pearson" or "spearman"
<code>removeFreeNodes</code>	remove free nodes from network?
<code>labelSize</code>	label size
<code>labelDist</code>	distance from labels to nodes
<code>shape</code>	shape of nodes
<code>degreeForSize</code>	should node size correspond to degree?
<code>label</code>	label nodes?
<code>onlyPositive</code>	only draw positive correlations?
<code>z.threshold</code>	z-score threshold
<code>FDR.threshold</code>	FDR threshold
<code>nodeSize</code>	size of node

Value

A list including a matrix of z-scores, a matrix of raw p-values, a matrix of adjusted p-values, and a summary data.frame

Author(s)

Dario Tommasini

Examples

```

library(ExperimentHub)
eh = ExperimentHub()
eh_query = query(eh, c("multiWGCNadata"))
astrocyte_se = eh_query[["EH8223"]]
datExpr = assays(astrocyte_se)[[1]]
diffCoexpression(datExpr, c(rep(1,20), rep(2,16)),
  geneList = c("Gfap", "Vim", "Aspg", "Serpina3n", "Cp", "Osmr", "Cd44",
    "Cxcl10", "Hspb1", "Timp1", "Slpr3", "Steap4", "Lcn2"))

```

diffModuleExpression *Differential module expression*

Description

Runs (and plots) the differential module expression analysis

Usage

```
diffModuleExpression(
  WGCNAobject,
  geneList,
  design,
  plotTitle = NULL,
  mode = c("PC1", "Zscore"),
  testColumn = 2,
  refColumn = 3,
  test = c("ANOVA", "PERMANOVA"),
  plot = TRUE
)
```

Arguments

WGCNAobject	WGCNA object
geneList	vector of genes in WGCNAobject
design	the sampleTable
plotTitle	title for the plot
mode	either PC1 or Zscore, default is PC1
testColumn	the column of the sampleTable to be resolved
refColumn	the column of the sampleTable to be used as biological variation
test	statistical test to perform, either "ANOVA" or "PERMANOVA"
plot	generate a plot?

Value

a data.frame with the resulting p-values

Examples

```
library(ExperimentHub)
eh = ExperimentHub()
eh_query = query(eh, c("multiWGCNAdata"))
astrocyte_se = eh_query[["EH8223"]]
sampleTable = colData(astrocyte_se)
astrocyte_networks = eh_query[["EH8222"]]
```

```
diffModuleExpression(astrocyte_networks[["combined"]],
  topNGenes(astrocyte_networks$combined, "combined_013"),
  sampleTable,
  test = "ANOVA",
  plotTitle = "combined_013",
  plot = TRUE)
```

drawMultiWGCNAnetwork *Draw multiWGCNA network*

Description

Draw a network where nodes are modules and edges represent significant gene overlap. Modules are sorted by levels 1, 2, and 3.

Usage

```
drawMultiWGCNAnetwork(
  WGCNAlist,
  comparisonList,
  moduleOfInterest,
  design,
  overlapCutoff = 0,
  padjCutoff = 1,
  removeOutliers = TRUE,
  alpha = 1e-50,
  layout = NULL,
  hjust = 0.4,
  vjust = 0.3,
  width = 0.5,
  colors = NULL
)
```

Arguments

WGCNAlist	list of WGCNA objects
comparisonList	the list of overlap comparisons ie from <code>iterate(myNetworks, overlapComparisons, ...)</code>
moduleOfInterest	module of interest, ie "combined_001"
design	the sampleTable design matrix
overlapCutoff	cutoff to remove module correspondences with less than this number of genes
padjCutoff	cutoff to remove module correspondences above this significance value
removeOutliers	remove outlier modules?
alpha	alpha level of significance

layout	layout of network to be passed to plot function of igraph object, defaults to multiWGCNA custom layout
hjust	horizontal justification of labels
vjust	vertical justification of labels
width	width of labels
colors	colors to use for modules, should be the same length as the number of WGCNA objects in the WGCNAlist. Defaults to random colors for each condition.

Value

an igraph plot

Author(s)

Dario Tommasini

Examples

```
library(ExperimentHub)
eh = ExperimentHub()
eh_query = query(eh, c("multiWGCNAdata"))
astrocyte_se = eh_query[["EH8223"]]
sampleTable = colData(astrocyte_se)
astrocyte_networks = eh_query[["EH8222"]]
results = list()
results$overlaps = iterate(astrocyte_networks, overlapComparisons, plot=FALSE)
drawMultiWGCNAnetwork(astrocyte_networks,
  results$overlaps,
  "combined_013",
  sampleTable)
```

GetDatExpr	<i>Get expression data</i>
------------	----------------------------

Description

Returns the expression data frame a WGCNA object as a data.frame

Usage

```
GetDatExpr(object, genes = NULL)
```

Arguments

object	An object of class WGCNA
genes	a list of genes to subset to; default is NULL

Value

a data.frame

Author(s)

Dario Tommasini

Examples

```
library(ExperimentHub)
eh = ExperimentHub()
eh_query = query(eh, c("multiWGCNAdata"))
astrocyte_networks = eh_query[["EH8222"]]
datExpr = GetDatExpr(astrocyte_networks[[1]],
  genes = topNGenes(astrocyte_networks$EAE, "EAE_015", 20))
coexpressionLineGraph(datExpr) +
  geom_vline(xintercept = 20.5, linetype='dashed')
```

getModule

name: Name of WGCNAobject

Description

Returns the module name for a given gene

Usage

```
getModule(WGCNAobject, gene, use = "dynamicLabels")
```

Arguments

WGCNAobject	an object of class WGCNA
gene	a gene symbol, should be present in WGCNA object
use	which module label to use e.g. dynamicLabels or dynamicColors

Value

A module name

Examples

```
library(ExperimentHub)
eh = ExperimentHub()
eh_query = query(eh, c("multiWGCNAdata"))
astrocyte_networks = eh_query[["EH8222"]]
getModule(astrocyte_networks[[1]], 'Srgn')
```

getPreservation	<i>getPreservation</i>
-----------------	------------------------

Description

Performs a network preservation analysis

Usage

```
getPreservation(reference, test, nPermutations = 100, write = FALSE)
```

Arguments

reference	reference network of class WGCNA
test	test network of class WGCNA
nPermutations	number of permutations to perform; at least 50 permutations
write	write to file?

Value

a data.frame summarizing results of preservation analysis

Author(s)

Dario Tommasini

GetSignificantOverlap	<i>Get significant overlap</i>
-----------------------	--------------------------------

Description

A function that subsets an overlap comparisons data.frame.

Usage

```
GetSignificantOverlap(df, p.adj.threshold = 0.05, overlap.threshold = 10)
```

Arguments

df	a data.frame resulting from a call to computeOverlapsFromWGCNA
p.adj.threshold	an object of class WGCNA to compare with dataset2
overlap.threshold	an object of class WGCNA to compare with dataset1

Value

Returns a data.frame with significant overlaps

Author(s)

Dario Tommasini

iterate	<i>iterate: Iterate function across networks</i>
---------	--

Description

A high level function that iterates functions across a list of WGCNA objects

Usage

```
iterate(WGCNAlist, FUN, ...)
```

Arguments

WGCNAlist	a vector of objects of type WGCNAobject
FUN	function to iterate, either overlapComparisons or preservationComparisons
...	arguments to be passed on to overlapComparisons or preservationComparisons

Value

a comparison list from overlapComparisons or preservationComparisons

Author(s)

Dario Tommasini

Examples

```
library(ExperimentHub)
eh = ExperimentHub()
eh_query = query(eh, c("multiWGCNAdata"))
astrocyte_networks = eh_query[["EH8222"]]
results = list()
iterate(astrocyte_networks, overlapComparisons, plot=FALSE)
```

makeTraitTable	<i>Generate a trait table from a sample table</i>
----------------	---

Description

Generates a WGCNA-compatible trait table from a sampleTable dataframe. This function is deprecated. Use makeTraitTable2 instead.

Usage

```
makeTraitTable(inputTable, column, detectNumbers = FALSE)
```

Arguments

inputTable	the sampleTable data.frame
column	the column from the sampleTable to use as traits
detectNumbers	whether to consider traits with numbers as numerical rather than categorical variables

Value

a data.frame with integer values denoting the categorical sample traits

Examples

```
sampleTable = data.frame(Sample = c(paste0("EAE", 1:10), paste0("WT", 1:10)),
                          Disease = c(rep("EAE", 10), rep("WT", 10)),
                          Region = c(rep(c("Cbl", "Sc"), 5)))
makeTraitTable(sampleTable, 2)
```

makeTraitTable2	<i>Generate a trait table from a sample table (version 2)</i>
-----------------	---

Description

Generates a WGCNA-compatible trait table from a sampleTable dataframe.

Usage

```
makeTraitTable2(inputTable, column, detectNumbers = TRUE)
```

Arguments

inputTable	the sampleTable data.frame
column	the column from the sampleTable to use as traits
detectNumbers	whether to consider traits with numbers as numerical rather than categorical variables

Value

a data.frame with integer values denoting the categorical sample traits

Examples

```
sampleTable = data.frame(Sample = c(paste0("EAE", 1:10), paste0("WT", 1:10)),  
                          Disease = c(rep("EAE", 10), rep("WT", 10)),  
                          Region = c(rep(c("Cbl", "Sc"), 5)))  
makeTraitTable2(sampleTable, 2)
```

moduleComparisonPlot	<i>Module comparison plot</i>
----------------------	-------------------------------

Description

A plotting function that returns a heatmap and barplot for a module

Usage

```
moduleComparisonPlot(overlapDf, dataset1, dataset2)
```

Arguments

overlapDf	a data.frame resulting from a call to computeOverlapsFromWGCNA
dataset1	an object of class WGCNA to compare with dataset2
dataset2	an object of class WGCNA to compare with dataset1

Value

Returns a ggplot object (flowplot and heatmap) showing the module correspondence between two objects of class WGCNA

Author(s)

Dario Tommasini

Examples

```
library(ExperimentHub)
eh = ExperimentHub()
eh_query = query(eh, c("multiWGCNAdata"))
astrocyte_networks = eh_query[["EH8222"]]
overlapDf = computeOverlapsFromWGCNA(astrocyte_networks$EAE, astrocyte_networks$WT)
moduleComparisonPlot(overlapDf, astrocyte_networks$EAE, astrocyte_networks$WT)
```

moduleExpressionPlot *Plots an expression profile for a module*

Description

A plotting function that returns a heatmap and barplot for a module

Usage

```
moduleExpressionPlot(
  WGCNAobject,
  geneList,
  mode = c("PC1", "averageZscore"),
  legend = FALSE,
  title = NULL,
  clusterGenes = FALSE
)
```

Arguments

WGCNAobject	an object of class WGCNAobject
geneList	a vector of gene names to be extracted from WGCNAobject
mode	use first principal component or averageZscore?
legend	plot legend?
title	title of the plot
clusterGenes	cluster heatmap genes by hierarchical clustering?

Value

a patchworked ggplot object

Author(s)

Dario Tommasini

Examples

```
library(ExperimentHub)
eh = ExperimentHub()
eh_query = query(eh, c("multiWGCNAdat"))
astrocyte_networks = eh_query[["EH8222"]]
moduleExpressionPlot(astrocyte_networks[["combined"]],
  geneList = topNGenes(astrocyte_networks$combined, "combined_013"))
```

ModuleFlowPlot	<i>Module sankey diagram</i>
----------------	------------------------------

Description

A plotting function that returns a plot

Usage

```
ModuleFlowPlot(
  WGCNAlist,
  comparisonList,
  networks,
  labels = NULL,
  alpha = 1,
  x.scale = 2,
  y.scale = 1,
  width = 0.3,
  height = 0.2,
  color.by = c("trait", "network", "none"),
  color.low = "cyan",
  color.high = "magenta",
  col = NULL,
  my_layout = NULL,
  use.padj = FALSE,
  only.contiguous = TRUE,
  only.signif = TRUE,
  show.legend = TRUE,
  scale.by.size = TRUE,
  spacer = 10,
  label.y = 50,
  label.size = 4,
  base_family = "Helvetica",
  p.adj.threshold = 0.05,
  overlap.threshold = 10
)
```

Arguments

<code>WGCNAlist</code>	a data.frame resulting from a call to <code>computeOverlapsFromWGCNA</code>
<code>comparisonList</code>	an object of class <code>WGCNA</code> to compare with <code>dataset2</code>
<code>networks</code>	an object of class <code>WGCNA</code> to compare with <code>dataset1</code>
<code>labels</code>	labels to show under each networks
<code>alpha</code>	alpha value for the node tiles, default is 1
<code>x.scale</code>	x spacing parameter
<code>y.scale</code>	y spacing parameter
<code>width</code>	width of the block
<code>height</code>	height of the block
<code>color.by</code>	color fill for nodes, either 'network', 'trait', or NULL
<code>color.low</code>	default is cyan
<code>color.high</code>	default is magenta
<code>col</code>	color palette for node tiles, default is internal colors function
<code>my_layout</code>	pass a custom layout for nodes? Default is null
<code>use.padj</code>	use p.adj for edge width, default is false (use gene overlap value)
<code>only.contiguous</code>	only show continuous overlaps? Default is TRUE
<code>only.signif</code>	plot only significant overlap?
<code>show.legend</code>	show the legend?
<code>scale.by.size</code>	scale the node height by the size of the module? Default is TRUE.
<code>spacer</code>	space between the nodes
<code>label.y</code>	adjust the y coordinate for the network labels, default is 1
<code>label.size</code>	size of the node labels, default is 4
<code>base_family</code>	font family for ggraph
<code>p.adj.threshold</code>	adjusted p-value significance threshold for overlap
<code>overlap.threshold</code>	threshold for number of genes overlapping

Value

Returns a ggalluvial diagram comparing two networks

Author(s)

Dario Tommasini

Examples

```
library(ExperimentHub)
eh = ExperimentHub()
eh_query = query(eh, c("multiWGCNAdata"))
astrocyte_networks = eh_query[["EH8222"]]
results = list()
results$overlaps = iterate(astrocyte_networks, overlapComparisons, plot=FALSE)
ModuleFlowPlot(astrocyte_networks, results$overlaps, c('combined', 'EAE', 'WT'))
```

ModuleFlowPlot2Way *Module sankey diagram*

Description

A plotting function that returns a sankey plot comparing two networks. This is a deprecated function, please use ModuleFlowPlot instead.

Usage

```
ModuleFlowPlot2Way(
  overlapDf,
  dataset1,
  dataset2,
  only.signif = TRUE,
  show.legend = TRUE,
  col = NULL,
  ...
)
```

Arguments

overlapDf	a data.frame resulting from a call to computeOverlapsFromWGCNA
dataset1	an object of class WGCNA to compare with dataset2
dataset2	an object of class WGCNA to compare with dataset1
only.signif	plot only significant overlap?
show.legend	show the legend?
col	color palette, default is internal colors function
...	params to GetSignificantOverlap function for thresholds

Value

Returns a ggalluvial diagram comparing two networks

Author(s)

Dario Tommasini

Examples

```
library(ExperimentHub)
eh = ExperimentHub()
eh_query = query(eh, c("multiWGCNAdata"))
astrocyte_networks = eh_query[["EH8222"]]
overlapDf = computeOverlapsFromWGCNA(astrocyte_networks$EAE, astrocyte_networks$WT)
ModuleFlowPlot2Way(overlapDf, astrocyte_networks$EAE, astrocyte_networks$WT)
```

moduleToModuleHeatmap *Module to module heatmap*

Description

Returns a heatmap where color corresponds to FDR-adjusted overlap (hypergeometric test) and the label corresponds to the number of overlapping genes

Usage

```
moduleToModuleHeatmap(
  comparisonDf,
  dataset1 = NULL,
  dataset2 = NULL,
  trait1 = NULL,
  trait2 = NULL,
  list1 = NULL,
  list2 = NULL,
  filterByTrait = FALSE,
  alphaLevel = 0.05
)
```

Arguments

comparisonDf	the data.frame output of computeOverlapFromWGCNA
dataset1	optional; WGCNA object for dataset 1
dataset2	optional; WGCNA object for dataset 2
trait1	optional; subset to modules correlated to this trait for dataset 1
trait2	optional; subset to modules correlated to this trait for dataset 2
list1	subset to this list of modules for dataset 1
list2	subset to this list of modules for dataset 2
filterByTrait	only plot for modules that correlate with some trait?
alphaLevel	the alpha level of significance for module-trait correlation, defaults to 0.05

Value

A ggplot object

Examples

```
library(ExperimentHub)
eh = ExperimentHub()
eh_query = query(eh, c("multiWGCNAdata"))
astrocyte_networks = eh_query[["EH8222"]]
overlapDf = computeOverlapsFromWGCNA(astrocyte_networks$EAE, astrocyte_networks$WT)
moduleToModuleHeatmap(overlapDf)
```

name	<i>name: Name of WGCNAobject</i>
------	----------------------------------

Description

Returns the name of a WGCNAobject.

Usage

```
name(WGCNAobject)
```

Arguments

WGCNAobject an object of class WGCNA

Value

Returns the name of the WGCNA object, ie "EAE" for astrocyte_networks\$EAE.

Examples

```
library(ExperimentHub)
eh = ExperimentHub()
eh_query = query(eh, c("multiWGCNAdata"))
astrocyte_networks = eh_query[["EH8222"]]
name(astrocyte_networks$EAE)
```

overlapComparisons	<i>Overlap comparisons</i>
--------------------	----------------------------

Description

Compares modules between two objects of type WGCNAobjects within a WGCNAobject list given the indices. Recommended to be used in conjunction with the iterate function.

Usage

```
overlapComparisons(  
  comparisonList,  
  WGCNAlist,  
  first,  
  second,  
  element,  
  plot = TRUE,  
  write = FALSE  
)
```

Arguments

comparisonList	a list passed by the iterate function
WGCNAlist	list of objects of class WGCNA
first	index of first WGCNA object
second	index of second WGCNA object
element	element position in the comparison list (passed by iterate function)
plot	generate plots?
write	write results to file?

Value

A list, in which the first element is a data.frame showing the overlap results and the second element is a data.frame showing the best matching modules between the two WGCNA objects.

Author(s)

Dario Tommasini

Examples

```
library(ExperimentHub)  
eh = ExperimentHub()  
eh_query = query(eh, c("multiWGCNAdata"))  
astrocyte_networks = eh_query[["EH8222"]]  
results = list()  
results$overlaps = iterate(astrocyte_networks, overlapComparisons, plot=FALSE)
```

performANOVA	<i>Perform ANOVA</i>
--------------	----------------------

Description

Test association between module expression to traits using ANOVA

Usage

```
performANOVA(datExpr, design, testCondition, refCondition, alphaLevel = 0.05)
```

Arguments

datExpr	expression data.frame
design	the sampleTable
testCondition	test column in sampleTable
refCondition	reference column in sampleTable
alphaLevel	the significance level

Value

a data.frame with p-values for each association

permutationTestResults	<i>Permutation test results</i>
------------------------	---------------------------------

Description

The results of running the PreservationPermutationTest in the astrocyte vignette. This is provided since this function is quite slow. Please see the astrocyte vignette for more details.

Usage

```
data(permutationTestResults)
```

Format

A list of data.frames containing preservation results for each permutation

PlotMultiNodesTOMflow	<i>PlotMultiNodesTOMflow</i>
-----------------------	------------------------------

Description

Plots a sankey flow diagram showing the movement of genes from one WGCNA to multi-WGCNA networks. Uses the ggalluvial framework.

Usage

```
PlotMultiNodesTOMflow(  
  TOMDF,  
  labels = NULL,  
  alpha = 0.1,  
  width = 0.05,  
  color = "black"  
)
```

Arguments

TOMDF	created by BuildTOMFlowDF
labels	labels to use for the networks, default is Network1, Network2, etc.
alpha	alpha of flows
width	width of the strata
color	color of flows

Value

a ggplot object

Author(s)

Dario Tommasini, Xinye Li

preservationComparisonPlot	<i>Preservation Comparison Scatterplot</i>
----------------------------	--

Description

A plotting function that draws a scatterplot of preservation scores between two WGCNA objects

Usage

```
preservationComparisonPlot(  
  preservationList,  
  dataset1,  
  dataset2,  
  alphaLevel = 0.05,  
  outliers = FALSE  
)
```

Arguments

preservationList	a list resulting from a call to <code>preservationComparisons</code>
dataset1	an object of class <code>WGCNAobject</code> to compare with dataset2
dataset2	an object of class <code>WGCNAobject</code> to compare with dataset1
alphaLevel	alpha level of significance, default is 0.05
outliers	leave outlier modules? By default these are removed

Value

a ggplot object

Author(s)

Dario Tommasini

Examples

```
library(ExperimentHub)  
eh = ExperimentHub()  
eh_query = query(eh, c("multiWGCNAdata"))  
astrocyte_networks = eh_query[["EH8222"]]  
results = list()  
results$preservation=iterate(astrocyte_networks[c("EAE", "WT")],  
  preservationComparisons,  
  write=FALSE,  
  plot=FALSE,  
  nPermutations=2)  
preservationComparisonPlot(results$preservation$EAE_vs_WT,  
  astrocyte_networks$EAE,  
  astrocyte_networks$WT)
```

`preservationComparisons`*Preservation comparisons*

Description

A high level function that performs a perservation comparison between two WGCNAobjects in a WGCNAlist, usually supplied by iterate function

Usage

```
preservationComparisons(  
  comparisonList,  
  WGCNAlist,  
  first,  
  second,  
  element,  
  plot = FALSE,  
  write = FALSE,  
  alphaLevel = 0.05,  
  nPermutations = 100  
)
```

Arguments

<code>comparisonList</code>	a list passed by the iterate function
<code>WGCNAlist</code>	list of objects of type WGCNAobject
<code>first</code>	index of first WGCNAobject
<code>second</code>	index of second WGCNAobject
<code>element</code>	element position in the comparison list (passed by iterate function)
<code>plot</code>	generate plots?
<code>write</code>	write results to file?
<code>alphaLevel</code>	alpha level of significance for module-trait correlation
<code>nPermutations</code>	number of permutations, defaults to 100

Value

a list of preservation comparisons results across levels 1, 2, 3

Author(s)

Dario Tommasini

Examples

```
library(ExperimentHub)
eh = ExperimentHub()
eh_query = query(eh, c("multiWGCNAdat"))
astrocyte_networks = eh_query[["EH8222"]]
results = list()
iterate(astrocyte_networks[c("EAE", "WT")],
  preservationComparisons,
  write=FALSE,
  plot=FALSE,
  nPermutations=2)
```

PreservationPermutationTest

PreservationPermutationTest

Description

Performs a permutation test to determine if a null distribution of expected preservation scores for modules in this dataset if the labels were randomly assigned. Please look at the astrocyte vignette for more info.

Usage

```
PreservationPermutationTest(
  referenceDatExpr,
  design,
  constructNetworksIn,
  testPreservationIn,
  nPermutations = 100,
  nPresPermutations = 100,
  ...
)
```

Arguments

referenceDatExpr	
	the combined datExpr
design	the sampleTable
constructNetworksIn	
	the condition to use for network construction, e.g. for the astrocyte data, this is "EAE"
testPreservationIn	
	the condition to use for testing preservation, e.g. for the astrocyte data, this was "WT"
nPermutations	the number of permutations to perform for permutation test

```

nPresPermutations      the number of permutations to perform in modulePreservation function
...                    arguments to pass to blockwiseModules function for network construction (should
                        be the same as used for constructing the original network)

```

Value

A list of data.frames with preservation results for each permutation

Author(s)

Dario Tommasini

Examples

```

## Not run:
library(ExperimentHub)
eh = ExperimentHub()
eh_query = query(eh, c("multiWGCNAdata"))
astrocyte_networks = eh_query[["EH8222"]]
astrocyte_se = eh_query[["EH8223"]]
sampleTable = colData(astrocyte_se)
results = list()
results$permutation.test = PreservationPermutationTest(
  astrocyte_networks$combined@datExpr[sample(17000,3000),],
  sampleTable,
  constructNetworksIn = "EAE", # Construct networks using EAE samples
  testPreservationIn = "WT", # Test preservation of disease samples in WT
  nPermutations = 10, # Number of permutations for permutation test
  nPresPermutations = 10, # Number of permutations for modulePreservation
  networkType = "signed", TOMType = "unsigned",
  power = 12, minModuleSize = 100, maxBlockSize = 25000,
  reassignThreshold = 0, minKMEtoStay = 0, mergeCutHeight = 0,
  numericLabels = TRUE, pamRespectsDendro = FALSE,
  deepSplit = 4, verbose = 3
)

## End(Not run)

```

PreservationScoreDistribution

PreservationScoreDistribution

Description

Extracts the preservation score distribution from the results of PreservationPermutationTest.

Usage

```
PreservationScoreDistribution(preservationData, moduleOfInterestSize)
```

Arguments

```

preservationData
    the results from PreservationPermutationTest
moduleOfInterestSize
    the number of genes in your module of interest

```

Value

A data.frame with Z-summary preservation scores of the module from each permutation and the corresponding module size

Author(s)

Dario Tommasini

Examples

```

# Remove outlier modules
permutationTestResultsFiltered = lapply(permutationTestResults, function(x)
x[!x$is.outlier.module,])

# Find preservation score distribution for a given module size
scores.summary = PreservationScoreDistribution(
permutationTestResultsFiltered,
moduleOfInterestSize = 303)

```

runDME

Run differential module expression

Description

A wrapper to run diffModuleExpression on all the modules in a network

Usage

```

runDME(
  WGCNAobject,
  design,
  alphaLevel = 0.05,
  testCondition = NULL,
  refCondition = NULL,
  p.adjust = "fdr",
  plot = FALSE,
  test = c("ANOVA", "PERMANOVA"),
  write = FALSE,
  out = NULL
)

```

Arguments

WGCNAobject	object of class WGCNA with the modules to run DME on
design	the sampleTable
alphaLevel	level of significance
testCondition	the column of the sampleTable to be resolved
refCondition	the column of the sampleTable to be used as biological variation
p.adjust	adjust for multiple comparisons, argument to pass to p.adjust function
plot	generate a plot?
test	statistical test to perform, either "ANOVA" or "PERMANOVA"
write	write results to a file?
out	file name for DME plots, only used if write is TRUE

Value

a data.frame summarizing the results of the analysis

Author(s)

Dario Tommasini

Examples

```
library(ExperimentHub)
eh = ExperimentHub()
eh_query = query(eh, c("multiWGCNAdata"))
astrocyte_se = eh_query[["EH8223"]]
sampleTable = colData(astrocyte_se)
astrocyte_networks = eh_query[["EH8222"]]
runDME(astrocyte_networks[["combined"]],
  design = sampleTable,
  p.adjust = "fdr",
  refCondition = "Region",
  testCondition = "Disease")
```

summarizeResults

summarizeResults: Summarize results from a results list object

Description

Prints (or writes) a summary of the results from a results list object

Usage

```
summarizeResults(  
  myNetworks,  
  results,  
  alphaLevel = 0.05,  
  write = FALSE,  
  outputFile = "results.txt"  
)
```

Arguments

- myNetworks a list of WGCNAobjects
- results results list
- alphaLevel alpha level of significance
- write write to file?
- outputFile name of output file, defaults to results.txt

Value

prints a summary of results from the multiWGCNA analysis

TOMFlowPlot	<i>TOMFlowPlot</i>
-------------	--------------------

Description

Plots a sankey flow diagram showing the movement of genes from one WGCNA to another WGCNA.
Uses the ggalluvial framework.

Usage

```
TOMFlowPlot(  
  WGCNAlist,  
  networks,  
  toms,  
  genes_to_label,  
  alpha = 0.1,  
  color = "black",  
  width = 0.05  
)
```

Arguments

WGCNAlist	list of WGCNA objects
networks	list of network names of length 2
toms	a list of TOM distance objects of length 2
genes_to_label	genes to label across two networks
alpha	alpha of flows
color	color of flows
width	width of the strata

Value

a ggplot object

Author(s)

Dario Tommasini

topNGenes	<i>topNGenes: Top N genes of a module</i>
-----------	---

Description

Returns the top N number of genes of a module. All genes returned if no number is specified. Genes are in order of intramodular connectivity.

Usage

```
topNGenes(WGCNAobject, module, nGenes = NULL)
```

Arguments

WGCNAobject	an object of class WGCNA
module	the name of the module in WGCNAobject
nGenes	an integer from 1 to module size; returns all genes if left NULL

Value

a character vector of the genes/probes in the module

Examples

```
library(ExperimentHub)
eh = ExperimentHub()
eh_query = query(eh, c("multiWGCNAdata"))
astrocyte_networks = eh_query[["EH8222"]]
topNGenes(astrocyte_networks$EAE, "EAE_015", nGenes = 10)
```

WGCNA-class	<i>The WGCNA Class</i>
-------------	------------------------

Description

The WGCNA class is the main class used in multiWGCNA to store results from a weighted gene co-expression network analysis. These include the original unaltered expression data used as input, connectivity metrics, module assignment, input sample conditions, trait

Value

NA

Slots

- datExpr The expression data, connectivity data, and module assignment
- conditions A data.frame with integer conditions for WGCNA
- trait A data.frame showing pearson correlation values to traits
- moduleEigengenes A data.frame of module eigengenes for each module across samples
- outlierModules A vector of modules classified by our algorithm as being driven by sample outliers

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