Package 'wTO'

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Туре	Package				
	e Computing Weighted Topological Overlaps (wTO) & Consensus wTO Network				
Versi	on 2.1				
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Descr	ription Computes the Weighted Topological Overlap with positive and negative signs (wTO) networks given a data frame containing the mRNA count/ expression/ abundance per sample, and a vector containing the interested nodes of interaction (a subset of the elements of the full data frame). It also computes the cut-off threshold or p-value based on the individuals bootstrap or the values reshuffle per individual. It also allows the construction of a consensus network, based on multiple wTO networks. The package includes a visualization tool for the networks. More about the methodology can be found at <doi:10.1186 s12859-018-2351-7="">.</doi:10.1186>				
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	CorrelationOverlap				

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Description

This function computes the correlation between Nodes and the Overlapping Nodes of interest.

Usage

CorrelationOverlap(Data, Overlap, method)

Arguments

Data data.frame containing the expression data. Nodes on the Rows, Individuals on

the Columns. Don't forget to give the names to the Nodes and to the Individuals.

Nodes must have the row.names() with the Node Name.

Overlap A vector containg the names of the Nodes of interest.

method Spearman ("s", "spearman") or Pearson ("p", "pearson") correlation

Author(s)

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Example GRF Example GRF

Description

ExampleGRF data.frame containing data.frame containing names of GRFs.

Usage

data(ExampleGRF)

Format

data.frame 184 lines, 1 column.

metagenomics_abundance

metagenomics_abundance

Description

metagenomics_abundance

Usage

```
data('metagenomics_abundance')
```

Format

data.frame from The USC Microbial Observatory. The data is public available at https://www.ebi.ac.uk/metagenomics/proje

Microarray_Expression1

Microarray_Expression1

Description

Microarray_Expression1 data.frame containing expression data for 1000 genes and 18 individuals.

Usage

Microarray_Expression1

Format

data.frame 1000 lines, 18 columns.

Microarray_Expression2

Microarray_Expression2

Description

Microarray_Expression2 data.frame containing expression data for 1000 genes and 18 individuals.

Usage

Microarray_Expression2

Format

data.frame 1000 lines, 18 columns.

4 NetVis

Description

Given a set of Nodes and the weight of the edges, a cutoff for the edges, it draws the networks. Returns a list with the nodes and edges attributes. And plots the network.

Usage

```
NetVis(
 Node.1,
 Node.2,
 wTO,
  pval = NULL,
 MakeGroups = FALSE,
  padj = NULL,
  cutoff = list(kind = "Threshold", value = 0.5),
  layout = NULL,
  smooth.edges = T,
  path = NULL,
  Cluster = F,
  legend = T,
  shape = list(shape = "triangle", names = NULL),
 manipulation = F
)
```

Arguments

Node.1	Names of the Nodes.1 that are connected to the Nodes.2. It's the output from wTO.Complete or Consensus.
Node.2	Names of the Nodes.2 that are connected to the Nodes.1. It's the output from wTO.Complete or Consensus.
wTO	weight of the links, the wTO output from wTO.Complete or wTO.Consensus.
pval	p-values for the wTO value. By default it is NULL.
MakeGroups	algorithm to find clusters. One of the followings: walktrap, optimal, spinglass, edge.betweenness, fast_greedy, infomap, louvain, label_prop, leading_eigen. Default to FALSE.
padj	Adjusted p-values for the wTO value. By default it is NULL.
cutoff	It's a list containing the kind of cutoff to be used (pval, Threshold or pval.adj)and it's value. Example: cutoff= list(kind = "Threshold", value = 0.5)
layout	a layout from the igraph package.
smooth.edges	If the edges should be smoothed or not.
path	If the graph should be saved specify the name of the file.

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Cluster TRUE or FALSE if the nodes should be clustered (double click to uncluster).

legend TRUE or FALSE if the legend should appear.

shape a list shape=list(shape = "triangle", names = NULL), with the shape and the

IDs that should have a different shape, shape can be: diamond, star, triangle,

triangleDown or square.

manipulation TRUE or FALSE if the graph should be editable.

Author(s)

Deisy Morselli Gysi <deisy at bioinf.uni-leipzig.de>

Examples

```
## Not run:
X = wTO.Complete( k =1, n = 5, Data = Microarray_Expression1,
Overlap = ExampleGRF$x[1:10], method = "p", plot = FALSE)
# Plot with the default aguments.
NetVis(Node.1 = X$wTO$Node.1, Node.2 = X$wTO$Node.2,
wTO = X$wTO$wTO_sign, cutoff = list(kind =
"Threshold", value = 0.50))
# Plotting just the edges with p-value < 0.05, with straight edges, nodes clustered,
# no legend and mapipulation of the graph enabled.
 NetVis(Node.1 = X$wTO$Node.1, Node.2 = X$wTO$Node.2,
 wTO = X$wTO$wTO_sign, pval = X$wTO$pval_sign,
 padj = X$wTO$pval_sign,
 cutoff= list(kind = "pval", value = 0.05),
 smooth.edges = FALSE,
Cluster = TRUE, legend = FALSE, manipulation = TRUE)
\# Plotting just the edges with wTO > 0.50, no legend and the nodes:
# "ZNF738", "ZNF677" with triagle shape,
# no legend and mapipulation of the graph enabled.
NetVis(Node.1 = X$wTO$Node.1, Node.2 = X$wTO$Node.2,
wTO = X$wTO$wTO_sign, pval = X$wTO$pval_sign,
 padj = X$wTO$pval_sign, cutoff= list(kind = "Threshold", value = 0.5),legend = FALSE,
 shape = list(shape = "triangle", names = c("ZNF738", "ZNF677")))
## End(Not run)
```

wTO wTO

Description

Calculates the weighted topologycal overlap (wTO) between a set of Nodes and the Overlapping nodes. This function implements the method from Nowick (2009).

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Usage

```
wTO(A_TF, sign = c("abs", "sign"))
```

Arguments

A_TF Is the weighted adjency matrix (correlation matrix).

sign ("abs", "sign") if the user wants to use the absolute correlation or the signed

correlation.

Value

A matrix containing the wTO values.

Author(s)

Deisy Morselli Gysi <deisy at bioinf.uni-leipzig.de>

References

Katja Nowick, Tim Gernat, Eivind Almaas and Lisa Stubbs (2009) <doi:10.1073/pnas.0911376106>

wTO.Complete

wTO.Complete

Description

Compute the wTO and also the bootstraps. Proposed at: arXiv:1711.04702

Usage

```
wTO.Complete(
    k = 1,
    n = 100,
    Data,
    Overlap = row.names(Data),
    method = "p",
    method_resampling = "Bootstrap",
    pvalmethod = "BH",
    savecor = F,
    expected.diff = 0.2,
    lag = NULL,
    ID = NULL,
    normalize = F,
    plot = T
)
```

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Arguments

k Number of threads to be used for computing the weight Topological Overlap.

Default is set to 1.

n Number of resamplings, used to compute the empirical distribuitions of the

links. Default is set to 100.

Data data.frame containing the count / expression data for the correlation.

Overlap Set of nodes of interest, where the Overlapping weights will be computed.

method Type of the correlation that should be used. "s" / "spearman" will compute the

rank spearman correlation, "p" / "pearson" will compute the linear correlation.

If no value is given, the default is to use "p".

method_resampling

method of the resampling. Bootstrap, BlockBootstrap or Reshuffle. Bootstrap null hypothesis is that the wTO is random, and Reshuffle tests if the wTO is

equal to zero.

pvalmethod method to compute the multiple test correction for the pvalue. for more infor-

mation check the function p.adjust.

savecor T/F if need to save the correlation.

expected.diff Difference expected between the real wTO and resampled wTO By default, it is

set to 0.2.

lag time dependency, lag, if you are using the BlockedBootstrap.

ID of the samples for the blocked bootstrap (for repeated measures).

normalize T/F Should the data be normalized?

plot T/F Should the diagnosis plot be plotted?

Value

a list with results.

- wTO is a data.frame containing the Nodes, the wTO computed using the signed correlations, the pvalue and the adj.pvalue.
- abs.wTO is a data.frame containing the Nodes, the wTO computed using the absolute correlations, the pvalue and the adj.pvalue.
- Correlation is a data.frame containing the correlation between all the nodes.
- Empirical.Quantile quantile values for the empirical distribution.
- Quantile quantile values for the sample distribution.

Author(s)

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Examples

```
## Not run:
# Using spearman rank correlation and bonferroni correction for the pvalues.
wTO.Complete( k = 8, n = 1000, Data = Microarray_Expression1,
Overlap = ExampleGRF$x, method = "s", pvalmethod = "bonferroni")
 # Changing the resampling method to Reshuffle.
wTO.Complete( k =1, n = 1000, Data = Microarray_Expression1,
Overlap = ExampleGRF$x, method_resampling = "Reshuffle")
 # Changing the resampling method to BlockBootstrap, with a lag of 2.
 row.names(metagenomics_abundance) = metagenomics_abundance$OTU
 metagenomics_abundance = metagenomics_abundance[,-1]
wTO.Complete( k =1, n = 1000, Data = metagenomics_abundance, method = "s",
Overlap = row.names(metagenomics_abundance), method_resampling = "BlockBootstrap", lag = 2)
wTO.Complete( k =2, n = 1000, Data = Microarray_Expression1, method = "s",
Overlap = ExampleGRF$x, method_resampling = "BlockBootstrap", ID = rep(1:9,each = 2))
X = wTO.Complete( k =1, n = 1000, Data = Microarray_Expression1,
Overlap = ExampleGRF$x, method = "p", plot = FALSE)
## End(Not run)
```

wTO.Consensus

wTO.Consensus

Description

Consensus requires a list of data.frame containing the pair of nodes, and the wTO values for all networks that need to be joined. Reference: arXiv:1711.04702

Usage

```
wTO.Consensus(data)
```

Arguments

data

list of data.frame containing the "Node.1", "Node.2" and "wTO".

Author(s)

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Examples

wTO.export

```
wto_sig = EXAMPLE$wTO$wTO_sign,
pvalsig = EXAMPLE$wTO$pval_sig),
data.frame(Node.1 = EXAMPLE$wTO$Node.1,
Node.2 = EXAMPLE$wTO$Node.2,
wtoabs = EXAMPLE$wTO$wTO_abs,
pvalabs = EXAMPLE$wTO$pval_abs))
CONS = wTO.Consensus(data)
## End(Not run)
```

wTO.export

wTO.export

Description

Exports the significative interactions, the wTO weight and pvalues into a .txt file, tab separeted. This file can be imported in other visualization tools (Cytoscape for example).

Usage

```
wTO.export(DATA, path, sign = TRUE, pvalue = 0.05, padj = 0.05, prop.NA = 0.5)
```

Arguments

DATA	Output from the function wTO.Complete or wTO.Consensus.
path	Path and file name where the .txt file should be saved.
sign	Should the network contain the results for the signed network or unsigned? Only for data coming from wTO.Complete.
pvalue	cutoff p-value for the network. Only for data coming from wTO.Complete.
padj	cutoff adjusted p-value for the network. Only for data coming from wTO.Complete.
prop.NA	cutoff proportion of NAs for the network. Only for data coming from wTO.Consensus.

Examples

10 wTO.fast

```
CN = wTO.Consensus(data = list(Ex_k1_cor_p_boot_p005_sig,
Ex_k1_cor_p_boot_p005_abs))
wTO.export(CN, './CN.txt')
### You can store the result on the workspace.
y = wTO.export(CN, './CN.txt')
head(y)
## End(Not run)
```

wTO.fast

wTO.fast

Description

Compute the wTO and also the bootstraps. Proposed at arXiv:1711.04702. This is a quicker version of the wTO.Complete. It doesn't contain diagnose plots nor a parallel version.

Usage

```
wTO.fast(
  Data,
  Overlap = row.names(Data),
  method = "p",
  sign = "sign",
  delta = 0.2,
  n = 10,
  method_resampling = "Bootstrap",
  lag = NULL,
  ID = NULL
)
```

Arguments

Data	data.frame containing the count / expression data for the correlation.
Overlap	Set of nodes of interest, where the Overlapping weights will be computed.
method	Type of the correlation that should be used. "s" / "spearman" will compute the rank spearman correlation, "p" / "pearson" will compute the linear correlation. If no value is given, the default is to use "p".
sign	Should the wTO be signed?
delta	expected difference between the real wTO and the bootstraped.
n	Number of resamplings, used to compute the empirical distribuitions of the links. Default is set to 100.

wTO.in.line

method_resampling

method of the resampling. Bootstrap or BlockBootstrap.If the second is used,

please give the lag (time dependency among the data).

lag Time dependency for the blocked bootstrap (for time series).

ID of the samples for the blocked bootstrap (for repeated measures).

Author(s)

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Examples

```
wTO.fast(Data = Microarray_Expression1,
Overlap = ExampleGRF$x,
method = "p")

# For a time series with lag = 4
# wTO.fast(Data = Microarray_Expression1,
# Overlap = ExampleGRF$x,
# method = "p",
# method_resampling = 'BlockBootstrap',
# lag = 4)

# For a study where the individuals were measured multiple times.
# wTO.fast(Data = Microarray_Expression1,
# Overlap = ExampleGRF$x,
# method = "p",
# method_resampling = 'BlockBootstrap',
# ID = rep(1:9, each= 2))
```

wTO.in.line

wTO.in.line

Description

Transforms a correlation matrix into the line format.

Transforms a correlation matrix into the line format.

Usage

```
wTO.in.line(d)
```

Arguments

d

correlation matrix to be converted into the line format.

Value

```
the wTO matrix into a data.frame: Node1, Node2 and wTO. the wTO matrix into a data.frame: Node1, Node2 and wTO.
```

wTO.rep_measure

Author(s)

Deisy Morselli Gysi <deisy at bioinf.uni-leipzig.de>

wTO.rep_measure wTO.rep_measure

Description

Compute the wTO for a repeated measures experiment and also the bootstraps. Proposed at arXiv:1711.04702. This is a quicker version of the wTO.Complete. It doesn'T contain diagnose plots nor a parallel version.

Usage

```
wTO.rep_measure(
  Data,
  Overlap = row.names(Data),
  ID,
  sign = "sign",
  delta = 0.2,
  n = 10
)
```

Arguments

Data data.frame containing the count / expression data for the correlation.

Overlap Set of nodes of interest, where the Overlapping weights will be computed.

ID a vector with the individuals identification

sign Should the wTO be signed?

delta expected difference between the real wTO and the bootstraped.

Number of resamplings, used to compute the empirical distribuitions of the links. Default is set to 100.

Author(s)

Deisy Morselli Gysi <deisy at bioinf.uni-leipzig.de>

Examples

```
#wTO.rep_measure(Data = Microarray_Expression1, ID = rep(c(1:9),2),
#Overlap = ExampleGRF$x)
```

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