# Package 'CoDiNA'

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Type Package

Title Co-Expression Differential Network Analysis

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<b>Description</b> Categorize links and nodes from multiple networks in 3 categories: Common links (alpha) specific links (gamma), and different links (beta). Also categorizes the links into subcategories and groups. The package includes a visualization tool for the networks. More information about the methodology can be found at: Gysi et. al., 2018 <arxiv:1802.00828>.</arxiv:1802.00828>
License GPL-2
LazyData TRUE
<b>Depends</b> R (>= 3.1)
Imports data.table, igraph, magrittr, plyr, visNetwork, reshape2
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.Random.seed

.Random.seed

## **Description**

Random numbers generated by set.seed(123)

as.igraph

as.igraph

## Description

Converts the CoDiNA.plot into an igraph object.

#### Usage

```
as.igraph(x)
```

## **Arguments**

Х

the output from the function plot.

## Value

the CoDiNA plot as an igraph object.

## Author(s)

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

```
suppressWarnings(RNGversion("3.5.0"))
Nodes = LETTERS[1:10]
Net1 = data.frame(Node.1 = sample(Nodes) , Node.2 = sample(Nodes), wTO = runif(10,-1,1))
Net2 = data.frame(Node.1 = sample(Nodes) , Node.2 = sample(Nodes), wTO = runif(10,-1,1))
Net3 = data.frame(Node.1 = sample(Nodes) , Node.2 = sample(Nodes), wTO = runif(10,-1,1))
DiffNet = MakeDiffNet (Data = list(Net1,Net2,Net3), Code = c('Net1', 'Net2', 'Net3') )
Graph = plot(x = DiffNet,
    layout = NULL, smooth.edges = TRUE,
    path = NULL, MakeGroups = FALSE, Cluster = FALSE,
    legend = TRUE, manipulation = FALSE, sort.by.Phi = FALSE)
```

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```
x = as.igraph(Graph)
plot(x)
```

AST

AST

## Description

This data.table contains node and the weighted topological overlap (wTO) of Transcription Factors (TFs), from GSE4290 (Sun, 2006) for 50 brain samples with ASTgodendrogliomas. The wTO was calculated using the package wTO.

## Usage

```
data("AST")
```

#### **Format**

A data frame with 3488761 observations on the following 3 variables.

```
Node.1 a factor with levels. TF names

Node.2 a factor with levels. TF names

cor a numeric vector. wTO values calculated using only the TFs
```

## Source

https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=gse4290

## References

Sun L, Hui AM, Su Q, Vortmeyer A et al. Neuronal and glioma-derived stem cell factor induces angiogenesis within the brain. Cancer Cell 2006 Apr;9(4):287-300. PMID: 16616334

```
data(AST)
str(AST)
```

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ClusterNodes

ClusterNodes

## **Description**

Categorize the Nodes into Phi and Phi tilde.

#### Usage

```
ClusterNodes(DiffNet, cutoff.external = 0.8, cutoff.internal = 0.5)
```

## **Arguments**

DiffNet The Differential network from MakeDiffNet cutoff.external

The cut-off between the clusters (delta from the center to the edge coordinates), the closer to 1, the better.

cutoff.internal

The cut-off inside the clusters (delta from the theoretical cluster to the edge coordinates), the closer to zero, the better.

## **Examples**

```
DiffNet = MakeDiffNet (Data = list(CTR, AST), Code = c('CTR', 'AST') )
Genes_Phi = ClusterNodes(DiffNet, cutoff.external = 0.5, cutoff.internal = 0.25)
table(Genes_Phi$Phi_tilde)
```

CTR

CTR

## **Description**

This data.table contains node and the weighted topological overlap (wTO) of Transcription Factors (TFs), from GSE4290 (Sun, 2006) for 23 brain samples with Controls. The wTO was calculated using the package wTO.

## Usage

```
data("CTR")
```

#### **Format**

A data frame with 3488761 observations on the following 3 variables.

```
Node.1 a factor with levels. TF names

Node.2 a factor with levels. TF names

cor a numeric vector. wTO values calculated using only the TFs
```

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#### Source

https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=gse4290

#### References

Sun L, Hui AM, Su Q, Vortmeyer A et al. Neuronal and glioma-derived stem cell factor induces angiogenesis within the brain. Cancer Cell 2006 Apr;9(4):287-300. PMID: 16616334

## **Examples**

```
data(CTR)
str(CTR)
```

GLI

GLI

## Description

This data.table contains node and the weighted topological overlap (wTO) of Transcription Factors (TFs), from GSE4290 (Sun, 2006) for 81 brain samples with glioblastomas. The wTO was calculated using the package wTO.

## Usage

```
data("GLI")
```

## **Format**

A data frame with 3488761 observations on the following 3 variables.

```
Node.1 a factor with levels. TF names

Node.2 a factor with levels. TF names

cor a numeric vector. wTO values calculated using only the TFs
```

#### Source

https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=gse4290

#### References

Sun L, Hui AM, Su Q, Vortmeyer A et al. Neuronal and glioma-derived stem cell factor induces angiogenesis within the brain. Cancer Cell 2006 Apr;9(4):287-300. PMID: 16616334

```
data(GLI)
str(GLI)
```

6 MakeDiffNet

fNet MakeDiffNet	

#### **Description**

Categorize links into Phi categories, calculate the distance to the center and also normlize the distance into some categories: Phi and Phi tilda, group and all.

#### Usage

```
MakeDiffNet(Data, Code, cutoff = 0.33, stretch = TRUE)
```

## **Arguments**

Data	List of data.frames containig Node.1, Node.2 and the correlation value
Code	Name of each one of the networks.
cutoff	By default, the cutoff is $0.33$ . If the user wants to use another value, it has to be cited on the description of the used methodology that the cutoff was changed.
stretch	Should the input data be normalized? Default to TRUE.

#### Value

Returns a data.table contating: Nodes names, correlation value for each network (the input values), the k means cluster that link belongs, the Phi groups (Phi and Phi tilda), the signed group that link belongs to, the unsigned group. The distance to the center, and the distance normalized by: Phi\_tilda, Phi, signed group or all data.

#### Author(s)

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

```
suppressWarnings(RNGversion("3.5.0"))
Nodes = LETTERS[1:20]
Net1 = data.frame(Node.1 = sample(Nodes) , Node.2 = sample(Nodes), wTO = runif(10,-1,1))
Net2 = data.frame(Node.1 = sample(Nodes) , Node.2 = sample(Nodes), wTO = runif(10,-1,1))
Net3 = data.frame(Node.1 = sample(Nodes) , Node.2 = sample(Nodes), wTO = runif(10,-1,1))
DiffNet = MakeDiffNet (Data = list(Net1,Net2,Net3), Code = c('Net1', 'Net2', 'Net3') )
print(DiffNet)
```

normalize 7

normalize

normalize

#### **Description**

Normalize a given variable.

#### Usage

```
normalize(m)
```

## **Arguments**

m

variable to be normalized in the interval [0,1]

## Author(s)

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

## **Examples**

```
Z = runif(10,-10,10)
normalize(Z)
```

OLI

OLI

## Description

This data.table contains node and the weighted topological overlap (wTO) of Transcription Factors (TFs), from GSE4290 (Sun, 2006) for 50 brain samples with oligodendrogliomas. The wTO was calculated using the package wTO.

#### Usage

```
data("OLI")
```

#### **Format**

A data frame with 3488761 observations on the following 3 variables.

```
Node.1 a factor with levels. TF names

Node.2 a factor with levels. TF names

cor a numeric vector. wTO values calculated using only the TFs
```

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#### **Source**

https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=gse4290

## References

Sun L, Hui AM, Su Q, Vortmeyer A et al. Neuronal and glioma-derived stem cell factor induces angiogenesis within the brain. Cancer Cell 2006 Apr;9(4):287-300. PMID: 16616334

## **Examples**

```
data(OLI)
str(OLI)
```

OrderNames

OrderNames

## Description

Sorts each link's Nodes by the smallest value. Removes links that both nodes are the same.

## Usage

```
OrderNames(M)
```

## **Arguments**

Μ

data.frame to have the names ordered. Node.1, Node.2 and correlation value.

#### Value

a data.table whith Node.1 and Node.2, sorted by the smallest value between both.

## Author(s)

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

```
Nodes = LETTERS[1:10]
Z = data.frame(Node.1 = sample(Nodes) ,
Node.2 = sample(Nodes), cor = runif(10,-1,1))
OrderNames(Z)
```

plot.CoDiNA 9

plot.CoDiNA plot.CoDiNA

#### **Description**

Categorize the Nodes into Phi and groups categories. Also, creates an interactive view of the CoD-iNA network.

#### **Usage**

```
## S3 method for class 'CoDiNA'
plot(
    x,
    cutoff.external = 0,
    cutoff.internal = 1,
    cutoff.ratio = 1,
    layout = NULL,
    smooth.edges = TRUE,
    path = NULL,
    MakeGroups = FALSE,
    Cluster = FALSE,
    legend = TRUE,
    manipulation = FALSE,
    sort.by.Phi = FALSE,
    ...
)
```

#### Arguments

x Output from MakeDiffNet

cutoff.external

The cut-off between the clusters (delta from the center to the edge coordinates), the closer to 1, the better.

cutoff.internal

The cut-off inside the clusters (delta from the theoretical cluster to the edge

coordinates), the closer to zero, the better.

cutoff.ratio The cut-off for the ratio of both scores. Default is set to 1. The greater, the

better.

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.

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.

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

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```
legend TRUE or FALSE if the legend should appear.

manipulation TRUE or FALSE if the graph should be editable.

sort.by.Phi if the graph should be plotted in the Phi order

Additional plotting parameters.
```

#### Value

Returns a list contatining: The nodes description, the Edges description and the network graph.

#### Author(s)

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

#### **Examples**

```
suppressWarnings(RNGversion("3.5.0"))

Nodes = LETTERS[1:10]
Net1 = data.frame(Node.1 = sample(Nodes) , Node.2 = sample(Nodes), wTO = runif(10,-1,1))
Net2 = data.frame(Node.1 = sample(Nodes) , Node.2 = sample(Nodes), wTO = runif(10,-1,1))
Net3 = data.frame(Node.1 = sample(Nodes) , Node.2 = sample(Nodes), wTO = runif(10,-1,1))
DiffNet = MakeDiffNet (Data = list(Net1,Net2,Net3), Code = c('Net1', 'Net2', 'Net3') )
Graph = plot(x = DiffNet,
    layout = NULL, smooth.edges = TRUE,
    path = NULL, MakeGroups = FALSE, Cluster = FALSE,
    legend = TRUE, manipulation = FALSE, sort.by.Phi = FALSE)
Graph
```

print.CoDiNA

print.CoDiNA

## Description

Print on the screen the number of nodes and edges. To see the data.frame, call: data.frame().

#### Usage

```
## S3 method for class 'CoDiNA'
print(x, ...)
```

## **Arguments**

x Output from MakeDiffNet... Additional plotting parameters.

#### Value

Print on the screen the number of nodes and edges.

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#### Author(s)

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

#### **Examples**

```
Nodes = LETTERS[1:10]
Net1 = data.frame(Node.1 = sample(Nodes) , Node.2 = sample(Nodes), wTO = runif(10,-1,1))
Net2 = data.frame(Node.1 = sample(Nodes) , Node.2 = sample(Nodes), wTO = runif(10,-1,1))
Net3 = data.frame(Node.1 = sample(Nodes) , Node.2 = sample(Nodes), wTO = runif(10,-1,1))
DiffNet = MakeDiffNet (Data = list(Net1,Net2,Net3), Code = c('Net1', 'Net2', 'Net3') )
print(DiffNet)
```

summary.CoDiNA

summary.CoDiNA

#### Description

summary of the CoDiNA network.

## Usage

```
## S3 method for class 'CoDiNA'
summary(object, ...)
```

#### **Arguments**

object Output from MakeDiffNet
... Additional plotting parameters.

#### Value

Returns a summary describing the network.

#### Author(s)

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

```
Nodes = LETTERS[1:10]
Net1 = data.frame(Node.1 = sample(Nodes) , Node.2 = sample(Nodes), wTO = runif(10,-1,1))
Net2 = data.frame(Node.1 = sample(Nodes) , Node.2 = sample(Nodes), wTO = runif(10,-1,1))
Net3 = data.frame(Node.1 = sample(Nodes) , Node.2 = sample(Nodes), wTO = runif(10,-1,1))
DiffNet = MakeDiffNet (Data = list(Net1,Net2,Net3), Code = c('Net1', 'Net2', 'Net3') )
summary(DiffNet)
```

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