# Package 'c3net'

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Title C3NET
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<b>Description</b> This package allows inferring regulatory networks with direct physical interactions from expression data using C3NET.
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Depends igraph
R topics documented:
c3
c3net
checknet
copula
expdata
makemim
netplot
sigtestMTC
sigtestp
truenet
Index 12

2 c3

C3NET second step: Selecting maximum valued elements

с3

## **Description**

c3 takes the mutual information matrix as input and implements the second step of C3NET - see details.

## Usage

```
c3 ( mim )
```

## **Arguments**

mim

A symmetric square mutual information matrix, where the elements (i,j) correspond to the mutual information I(i,j) or I(j,i) between variables i and j.

#### **Details**

The C3NET algorithm consists of two main steps. The first step is the same as for relevance networks (RELNET), where all the non-significant mutual information values in the matrix is eliminated statistically if not significant. The second step of C3NET keeps all maximum valued mutual information values for each row in the matrix and sets the rest of the elements in the matrix zero (the diagonal of the matrix is ignored).

## Value

c3 returns a symmetric mutual information matrix, which is obtained after implementing the second step of C3NET algorithm. Specifically, the non-zero elements in the returned matrix represents undirected links between variables if it is statistically significant (the first step of C3NET).

## References

G. Altay, F. Emmert-Streib, "Inferring the conservative causal core of gene regulatory networks", BMC Systems Biology (2010) 4:132.

#### See Also

```
makemim, copula, sigtestp, sigtestMTC, netplot, c3net
```

```
data(expdata)
expdata <- copula(expdata)
mim <- makemim(expdata)

Ic <- mean(mim[upper.tri(mim)]) #Example cut-off for the first step of C3NET
# Ic <- 2 can be set for the example.
mim[mim < Ic] <-0 #nonsignificant values eliminated wrt C3NET step 1.</pre>
```

c3net 3

c3net

All in one function to infer network with C3NET

## **Description**

c3net takes a data set as input and computes the inferred network using C3NET. - see details.

#### Usage

#### **Arguments**

dataset Data set where rows are variables (e.g. genes) and columns are samples.

alpha Statistical significance threshold

itnum Number of iterations to resample data to get sampling distribution.

methodstep1 If it is "cutoff" than the input cutoffMI is used as threshold for elimination in step

1. If it is "MTC" then MTCmethod input is used as the specific multiple testing correction method to employ. If it is "justp" then only significance threshold

alpha is used to compute significance threshold MI.

MTCmethod It can be any of the options available in the R function p.adjust (e.g. "holm",

"hochberg", "hommel", "bonferroni", "BH", "BY")

cutoffMI If methodstep1 is set to "cutoff" then this value is used for MI threshold. If this

is set to 0 (or by default) it uses mean MI for threshold.

network If TRUE, the network is plotted in the end.

## **Details**

For Step 1 of C3NET, there are three options that can be used. If methodstep1="cutoff" then cutoffMI input is taken as the significance threshold for step 1. In case cutoffMI is set to 0 then by default mean MI is taken as cutoffMI. If methodstep1 is set to "MTC" then the method from input MTCmethod, itnum for iteration number to get null distribution and alpha for statistical significance is taken for computations. If methodstep1 is set to "justp" or anything else then just alpha and itnum is used for computations.

#### Value

c3net returns a symmetric mutual information matrix, which is obtained after implementing C3NET. Specifically, non-zero elements in the returned matrix represents undirected link between variables. The inferred network may also be plotted if the argument network is set TRUE.

## References

G. Altay, F. Emmert-Streib, "Inferring the conservative causal core of gene regulatory networks", BMC Systems Biology (2010) 4:132.

4 checknet

#### See Also

```
makemim, copula, c3, sigtestp, sigtestMTC
```

## **Examples**

```
data(expdata)
data(truenet)

net <- c3net(expdata, network=TRUE)
scores <- checknet(net,truenet)</pre>
```

checknet

Validating the inferred network

## **Description**

checknet takes the inferred mutual information matrix and the true network for validation. - see details.

## Usage

```
checknet( finalrelationmatrix, realrelationmatrix)
```

## **Arguments**

finalrelationmatrix

The inferred symmetric mutual information matrix, where i,j th element is the mutual information I(i,j) or I(j,i) between two variables i and j. The diagonal is set to zero.

realrelationmatrix

A symmetric reference connection matrix, where a 1 at i,j th element defines the connection between variables i and j and non-connection is represented by 0. The diagonal is all zero.

## Value

checknet returns a vector with 6 elements, that contains the parameters as output <- c(precision, Fscore, recall, TP, FP, FN).

## References

G. Altay, F. Emmert-Streib, "Inferring the conservative causal core of gene regulatory networks", BMC Systems Biology, (2010) 4:132.

#### See Also

```
makemim, copula, c3, c3net, c3, makemim, copula, sigtestp, sigtestMTC
```

copula 5

#### **Examples**

```
data(expdata)
data(truenet)
expdata <- copula(expdata)
mim <- makemim(expdata)

Ic <- 2  #Example cut-off value for mutual information for the first step of C3NET

mim[mim < Ic] <-0  #nonsignificant values eliminated wrt C3NET step 1.

net <- c3(mim)  # regulatory network inferred (non zero elements stand for links of # the predicted network)

scores <- checknet(net,truenet)</pre>
```

copula

Copula transformation of data set

## **Description**

copula takes a data set as input and applies a copula transformation - see details.

## Usage

```
copula ( expdata )
```

## **Arguments**

expdata

Data set where rows are variables (e.g. genes) and columns are samples.

#### **Details**

The data set can be copula transformed for more stable estimations of the mutual information matrix. Each row of the data set (gene samples of each gene) is copula transformed and obtained rdata. Here, the formula for conversion is copuladata =(rdata- 0.5)/numberofsamples.

## Value

copula returns a copula transformed data set.

#### References

G. Altay, F. Emmert-Streib, "Inferring the conservative causal core of gene regulatory networks", BMC Systems Biology (2010) 4:132.

## See Also

```
c3, c3net, makemim
```

6 expdata

#### **Examples**

```
data(expdata)
expdata <- copula(expdata)</pre>
```

expdata

Example data set

## **Description**

expdata This data set is the subset generated by SynTRen. The source network is taken from the within SynTren package program and it is real E.coli network with file name "EColi\_full\_HongWu\_Ma\_NAR2004.sif" of Ma HW, et. al. Nuc. Ac. Res. 2004.

## Usage

```
data ( expdata )
```

## **Arguments**

expdata

Example data set where rows are variables (e.g. genes) and columns are samples.

#### **Details**

The example is also given to provide a sample data set so that one can take it as reference while formatting the new data set as entry to c3net. Data sets must be complete and no empty or non-numerical character allowed.

## Value

expdata returns an example data set where rows are variables (e.g. genes) and columns are samples. It has 400 genes, with 898 edges, and 800 steady-state samples.

#### References

Ma HW, Kumar B, Ditges U, Gunzer F, Buer J, Zeng AP: An extended transcriptional regulatory network of Escherichia coli and analysis of its hierarchical structure and network motifs. Nucleic Acids Res 2004, 32:6643-6649

Bulcke, T.V., Leemput, K.V., Naudts, B., et. al. (2006) Syntren: a generator of synthetic gene expression data for design and analysis of structure learning algorithms. BMC Bioinformatics, 7(1), 43.

G. Altay, F. Emmert-Streib, "Inferring the conservative causal core of gene regulatory networks", BMC Systems Biology (2010) 4:132.

#### See Also

```
c3, c3net, makemim
```

```
data(expdata)
```

makemim 7

makemim

Constructing mutual information matrix

## **Description**

makemim takes the data set as input and computes mutual information values for each pair of variables (e.g. gene). - see details.

## Usage

```
makemim( expdata )
```

## **Arguments**

expdata

Data matrix where rows correspond to variables (e.g. genes) and columns are samples.

#### **Details**

A mutual information matrix is generated from a data set using an empirical Gaussian estimator.

## Value

makemim returns a symmetric mutual information matrix using empirical Gaussian estimator.

#### References

- G. A. Darbellay, I. Vajda, "Estimation of the information by an adaptive partitioning of the observation space", IEEE Trans. Inf. Theory vol.45 (1999)
- G. Altay, F. Emmert-Streib, "Inferring the conservative causal core of gene regulatory networks", BMC Systems Biology, (2010) 4:132.

#### See Also

```
c3,c3net, copula, checknet
```

```
data(expdata)
expdata <- copula(expdata)
mim <- makemim(expdata)</pre>
```

8 netplot

netplot

Plotting the inferred network

## **Description**

netplot takes the inferred mutual information matrix and plots a network wrt nonzero elements of the matrix - see details.

#### Usage

```
netplot(gnet)
```

## **Arguments**

gnet

A symmetric squared mutual information matrix that is inferred as final connection matrix, where i,j th element is the mutual information I(i,j) or I(j,i) between variables i and j. The nonzero elements may also be 1. Diagonal is all zero.

#### **Details**

A labelled undirected network is plotted wrt the row (column) names of the input matrix. When plotted, select "Layout" and then "Fruchterman-Reingold" and "OK" to get a better view. To remove labels, select "View" and deselect "Labels". To change the colors or size of nodes and edges, from "Select", click on "all edges" or "vertices" and on the network right click on nodes and change color or size.

## Value

netplot returns an undirected network.

## References

G. Altay, F. Emmert-Streib, "Inferring the conservative causal core of gene regulatory networks", BMC Systems Biology (2010) 4:132.

CSARDI G, NEPUSZ T.: igraph-package for R, 2008

## See Also

```
makemim, copula, c3, sigtestp, sigtestMTC
```

```
data(expdata)
data(truenet)
expdata <- copula(expdata)
mim <- makemim(expdata)
Ic <- 2 #Example cut-off for the first step of C3NET</pre>
```

sigtestMTC 9

```
mim[mim < Ic] <-0 #nonsignificant values eliminated wrt C3NET step 1.

net <- c3(mim) # regulatory network inferred (non zero elements stand for links of # the predicted network)

netplot(net)</pre>
```

sigtestMTC

Elimination of nonsignificant edges by application of a procedure for multiple testing correction

## **Description**

sigtestMTC takes a data set as input and computes the mutual information matrix in which the nonsignificant elements are eliminated by a multiple testing correction - see details.

#### Usage

```
sigtestMTC( data, alpha, itnum, methodsig="BH")
```

## **Arguments**

data Data set where rows are variables (e.g. genes) and columns are samples.

alpha Significance level

itnum Number of iterations to resample data to estimate sampling distribution.

methodsig A procedure to perform a multiple testing correction, either controling the false

discovery rate (FDR) or the family-wise error.

#### **Details**

The data set is resampled for each iteration and at the end the sampling distribution is obtained.

## Value

sigtestMTC returns an environment res that contains the new MI matrix, res\$Inew, that is obtained after the elimination of nonsignificant elements with a MTC procedure. For advanced users, who are able to write code in R, we provided the variables obtained in the function so that one might want to make further analysis without running the function again. Please see the source code for the additional return values of the environment res.

#### References

G. Altay, F. Emmert-Streib, "Inferring the conservative causal core of gene regulatory networks", BMC Systems Biology (2010) 4:132.

#### See Also

```
makemim, copula, c3, sigtestp
```

10 sigtestp

#### **Examples**

```
data(expdata)
data(truenet)
alpha <- 0.001
itnum <-2
res <- sigtestMTC( expdata, alpha, itnum, methodsig="BH")
net <- c3(res$Inew) # regulatory network inferred (non zero elements stand for links of # the predicted network)
scores <- checknet(net,truenet)</pre>
```

sigtestp

Significance test for elimination of nonsignificant edges

## **Description**

sigtestp takes the data set as input and computes the mutual information matrix in which the nonsignificant elements are eliminated by using only significance threshold alpha. - see details.

## Usage

```
sigtestp( data, alpha, itnum)
```

#### **Arguments**

data	Data set where rows are variables (e.g. genes) and columns are samples.
alpha	Statistical significance threshold

itnum Number of iterations to resample data to get sampling distribution.

#### **Details**

The data set is resampled completely at each iteration and at the end the sampling distribution is obtained for using in significance test.

#### Value

sigtestp returns environment res that contains the obtained threshold value res\$10. For advanced users, who are able to write code in R, we provided the variables obtained in the function so that one might want to make further analysis without running the function again. Please see the source code for the additional return values of the environment res.

## References

G. Altay, F. Emmert-Streib, "Inferring the conservative causal core of gene regulatory networks", BMC Systems Biology (2010) 4:132.

truenet 11

#### See Also

```
makemim, copula, c3,c3, sigtestMTC
```

## **Examples**

```
data(expdata)
data(truenet)
alpha <- 0.001
itnum <- 2
res <- sigtestp( expdata, alpha, itnum)
net <- c3(res$Inew) # regulatory network inferred (non zero elements stand for links of # the predicted network)
scores <- checknet(net,truenet)</pre>
```

truenet

Reference true network of the example data set

## **Description**

truenet A symmetric matrix, where non-zero elements stand for true connections.

## Usage

```
data( truenet )
```

## **Arguments**

truenet

Reference true network of the example data set.

## Value

truenet Reference true network of the example data set.

## References

G. Altay, F. Emmert-Streib, "Inferring the conservative causal core of gene regulatory networks", BMC Systems Biology (2010) 4:132.

## See Also

```
c3, c3net, makemim
```

```
data(truenet)
```

## **Index**

```
*Topic misc
    c3, 1
    c3net, 2
    checknet, 4
    copula, 5
    expdata, 6
    {\tt makemim}, \textcolor{red}{7}
    netplot, 8
    sigtestMTC, 9
    sigtestp, 10
    truenet, 11
c3, 1, 3–9, 11
c3net, 2, 2, 4-7, 11
checknet, 4, 7
copula, 2-4, 5, 7-9, 11
expdata, 6
makemim, 2-6, 7, 8, 9, 11
netplot, 2, 8
sigtestMTC, 2-4, 8, 9, 11
sigtestp, 2-4, 8, 9, 10
truenet, 11
```