# Package 'dcanr'

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Title Differential co-expression/association network analysis

**Version** 1.21.0

**Description** This package implements methods and an evaluation framework to infer differential co-expression/association networks. Various methods are implemented and can be evaluated using simulated datasets. Inference of differential co-expression networks can allow identification of networks that are altered between two conditions (e.g., health and disease).

**biocViews** NetworkInference, GraphAndNetwork, DifferentialExpression, Network

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**Encoding UTF-8** 

LazyData false

**Imports** igraph, foreach, plyr, stringr, reshape2, methods, Matrix, graphics, stats, RColorBrewer, circlize, doRNG

**Suggests** EBcoexpress, testthat, EBarrays, GeneNet, mclust, minqa, SummarizedExperiment, Biobase, knitr, rmarkdown, BiocStyle, edgeR

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VignetteBuilder knitr

Enhances parallel, doSNOW, doParallel

URL https://davislaboratory.github.io/dcanr/,
 https://github.com/DavisLaboratory/dcanr

BugReports https://github.com/DavisLaboratory/dcanr/issues

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2 dcanr-package

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# **Contents**

dcan	r-package dcanr: Differential co-expression/association network analysis	
Index		20
	sim102	19
	plotSimNetwork	
	performanceMeasure	
	perfMethods	
	mi.ap	
	getSimData	14
	dcZscore	13
	dcTest	12
	dcScore	10
	dcPipeline	8
	dcNetwork	7
	dcMethods	6
	dcEvaluate	5
	dcAdjust	4
	cor.pairs	4
	dcanr-package	2

# Description

Methods and an evaluation framework for the inference of differential co-expression/association networks.

## **Details**

There are three categories of functions available

- 1. Differential co-expression methods (DC) These functions are used to perform a differential co-expression analysis on experimental data with binary conditions.
- 2. Functions to evaluate DC methods These functions are used to evaluate methods implemented in the package and novel methods on simulated data. Expression data is simulated for 2 conditions, wild-type and knock-down of given genes.
- 3. By-products of implementations

dcanr-package 3

#### Differential co-expression methods (DC)

- dcMethods
- dcScore
- dcTest
- dcAdjust
- dcNetwork

#### **Functions to evaluate DC methods**

Accessors of simulated data:

- getConditionNames
- getSimData
- getTrueNetwork
- plotSimNetwork

Functions for evaluating inference methods

- dcPipeline
- dcEvaluate

## By-products of implementations

These are functions used in the package but have further uses in general:

- cor.pairs a faster implementation of pairwise correlation computation
- mi.ap pairwise computation of mutual information MI with data discretisation performed using adaptive partitioning
- perfMethods available performance metrics
- performanceMeasure performance measures of prediction algorithms. Predictions have to be binary

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#### See Also

Useful links:

- https://davislaboratory.github.io/dcanr/
- https://github.com/DavisLaboratory/dcanr
- Report bugs at https://github.com/DavisLaboratory/dcanr/issues

4 dcAdjust

cor.pairs

Fast pairwise correlation estimation

# Description

Fast estimation of pairwise correlation coefficients.

## Usage

```
cor.pairs(emat, cor.method = c("pearson", "spearman"))
```

# **Arguments**

emat a numeric matrix

cor.method a character, specifying the method to use for estimation. Possible values are

'pearson' (default) and 'spearman'

## Value

a numeric matrix with estimated correlation coefficients

# **Examples**

```
x <- matrix(rnorm(200), 100, 2)
cor.pairs(x)
cor.pairs(x, cor.method = 'spearman')</pre>
```

dcAdjust

Adjust for multiple testing in differential association analysis

# Description

Adjust for multiple hypothesis testing after performing statistical tests using dcTest. This can be performed using a method provided by the users. p.adjust is used by default.

# Usage

```
dcAdjust(dcpvals, f = stats::p.adjust, ...)
```

#### Arguments

dcpvals	a matrix, the result of the dcTest function. The results should be passed as
	produced by the function and not modified in intermediate steps
f	a function, the function to be used for adjustment. p.adjust from the stats package is the default with the specific adjustment method 'fdr' used. The range
	of available methods can be accessed using p.adjust.methods. Custom functions should accept a numeric vector of p-values as the first argument
	additional parameters to the adjustment function such as method

dcEvaluate 5

#### **Details**

Ensure that the p-value matrix passed to this function is the one produced by dcTest. Any modification to the result matrix will result in failure of the function.

This method applies the adjustment method only to one triangle of the matrix to ensure adjustment is not performed for duplicated tests (symmetric matrix). As results from the DiffCoEx and EBcoexpress do not produce p-values, this method does not change anything thereby returning the original matrix.

#### Value

a matrix, of adjusted p-values (or scores in the case of DiffCoEx and EBcoexpress) representing significance of differential associations.

#### See Also

```
dcTest p.adjust
```

## **Examples**

```
x <- matrix(rnorm(60), 2, 30)
cond <- rep(1:2, 15)
zscores <- dcScore(x, cond)
pvals <- dcTest(zscores, emat = x, condition = cond)
dcAdjust(pvals, p.adjust, method = 'fdr')</pre>
```

dcEvaluate

Evaluate performance of DC methods on simulations

# **Description**

Quantify the performance of a differential co-expression pipeline on simulated data.

```
dcEvaluate(
   simulation,
   dclist,
   truth.type = c("association", "influence", "direct"),
   perf.method = "f.measure",
   combine = TRUE,
   ...
)
```

6 dcMethods

## **Arguments**

simulation a list, storing data and results generated from simulations

dclist a list of igraphs, produced using dcPipeline

truth.type a character, specifying which level of the true network to retrieve: 'association'

(default), 'influence' or 'direct'

perf.method a character, specifying the method to use. Available methods can be accessed

using perfMethods

combine a logical, indicating whether differential networks from independent knock-outs

should be treated as a single inference or independent inferences (defaults to

TRUE)

.. additional parameters to be passed on to the performance metric method (see

performanceMeasure)

#### Value

a numeric, representing the performance metric. A single value if combine = TRUE and a named vector otherwise.

#### See Also

dcPipeline, performanceMeasure, perfMethods

#### **Examples**

```
data(sim102)
#run a standard pipeline
resStd <- dcPipeline(sim102, dc.func = 'zscore')
dcEvaluate(sim102, resStd)
dcEvaluate(sim102, resStd, combine = FALSE)</pre>
```

dcMethods

Get names of differential co-expression methods

#### **Description**

Returns a list of differential co-expression methods

#### Usage

dcMethods()

# Value

names of methods implemented

dcNetwork 7

## **Examples**

```
dcMethods()
```

dcNetwork

Generate a differential network from a DC analysis

# Description

Threshold the results from a differential co-expression analysis and create a differential network.

# Usage

```
dcNetwork(dcscores, dcpvals = NULL, thresh = NULL, ...)
```

# Arguments

dcscores	a matrix, the result of the dcScore function. The results should be passed as produced by the function and not modified in intermediate steps
dcpvals	a matrix or NULL, raw or adjusted p-values resulting from dcTest or dcAdjust respectively. Should be left NULL only if method is EBcoexpress or DiffCoEx
thresh	a numeric, threshold to apply. If NULL, defaults to $0.1$ for methods that generate a p-value, $0.9$ for posterior probabilities from EBcoexpress and $0.1$ on the absolute score from DiffCoEx
	see details

# **Details**

No extra arguments required for this function. The ellipsis are used to allow flexibility in pipelines.

## Value

an igraph object, representing the differential network. Scores are added as edge attributes with the name 'score'

#### See Also

```
dcScore, dcTest, dcAdjust
```

```
#create data
set.seed(360)
x <- matrix(rnorm(120), 4, 30)
cond <- rep(1:2, 15)

#perform analysis - z-score
zscores <- dcScore(x, cond)</pre>
```

8 dcPipeline

```
pvals <- dcTest(zscores, emat = x, condition = cond)
pvals <- dcAdjust(pvals, p.adjust, method = 'fdr')
ig <- dcNetwork(zscores, pvals, 0.1)

#perform analysis - DiffCoEx
dcscores <- dcScore(x, cond, dc.method = 'diffcoex')
ig <- dcNetwork(dcscores, thresh = 0.001)

#plot the resulting differential co-expression network
igraph::plot.igraph(ig)</pre>
```

dcPipeline

Run a DC pipeline on a simulation

## **Description**

Run a differential co-expression pipeline on data from a simulation experiment. A default pipeline can be used which consists of methods in the package or custom pipelines can be provided.

## Usage

```
dcPipeline(
   simulation,
   dc.func = "zscore",
   precomputed = FALSE,
   continuous = FALSE,
   cond.args = list(),
   ...
)
```

## **Arguments**

simulation	a list, storing data and results generated from simulations
dc.func	a function or character. Character represents one of the method names from dcMethods which is run with the default settings. A function can be used to provide custom processing pipelines (see details)
precomputed	a logical, indicating whether the precomputed inference should be used or a new one computed (default FALSE)
continuous	a logical, indicating whether binary or continuous conditions should be used (default FALSE). No methods implemented currently use continuous conditions. This is to allow custom methods that require continuous conditions
cond.args	a list, containing condition-specific arguments for the DC inference pipeline. See details
	additional parameters to dc.func

dcPipeline 9

#### **Details**

If dc. func is a character, the existing methods in the package will be run with their default parameters. The pipeline is as such: dcScore -> dcTest -> dcAdjust -> dcNetwork, resulting in a igraph object. Parameters to the independent processing steps can also be provided to this function as shown in the examples.

If precomputed is TRUE while dc.func is a character, pre-computed results will be used. These can then be evaluated using dcEvaluate.

Custom pipelines need to be coded into a function which can then be provided instead of a character. Functions must have the following structure:

```
function(emat, condition, ...)
```

They must return either an igraph object or an adjacency matrix stored in a base R 'matrix' or the S4 'Matrix' class, containing all genes in the expression matrix 'emat'. See examples for how the in-built functions are combined into a pipeline.

If the pipeline (in-built or custom) requires condition-specific parameters to run, cond.args can be used to pass these. For instance, LDGM requires lambda OR the number of edges in the target network to be specified for each inference/condition. For the latter case and with 3 different conditions, this can be done by setting cond.args = list('ldgm.ntarget' = c(100, 140, 200)). Non-specific arguments should be passed directly to the dcPipeline function call.

#### Value

a list of igraphs, representing the differential network for each independent condition (knock-out).

#### See Also

```
plot.igraph, dcScore, dcTest, dcAdjust, dcNetwork, dcMethods
```

```
#run a standard pipeline
resStd <- dcPipeline(sim102, dc.func = 'zscore')

#run a standard pipeline and specify params
resParam <- dcPipeline(sim102, dc.func = 'zscore', cor.method = 'pearson')

#run a standard pipeline and specify condition-specific params
resParam <- dcPipeline(
    sim102,
    dc.func = 'diffcoex',
    #arguments for the conditions ADR1 knockdown and UME6 knockdown resp.
    cond.args = list(diffcoex.beta = c(6, 20))
)

#retrieve pre-computed results
resPrecomputed <- dcPipeline(sim102, dc.func = 'zscore', precomputed = TRUE)
#run a custom pipeline</pre>
```

10 dcScore

```
analysisInbuilt <- function(emat, condition, dc.method = 'zscore', ...) {
  #compute scores
  score = dcScore(emat, condition, dc.method, ...)
  #perform statistical test
  pvals = dcTest(score, emat, condition, ...)
  #adjust tests for multiple testing
  adjp = dcAdjust(pvals, ...)
  #threshold and generate network
  dcnet = dcNetwork(score, adjp, ...)
  return(dcnet)
}
resCustom <- dcPipeline(sim102, dc.func = analysisInbuilt)

plot(resCustom[[1]])</pre>
```

dcScore

Compute scores from differential association analysis

## **Description**

Implementations and wrappers for existing implementations for methods inferring differential associations/co-expression. This method requires a matrix of expression and a binary condition to compute the differential association scores for all pairs of features (genes). Applications are not limited to analysis of gene expression data and may be used for differential associations in general.

```
dcScore(emat, condition, dc.method, ...)
## S4 method for signature 'matrix'
dcScore(emat, condition, dc.method = "zscore", ...)
## S4 method for signature 'Matrix'
dcScore(emat, condition, dc.method = "zscore", ...)
## S4 method for signature 'data.frame'
dcScore(emat, condition, dc.method = "zscore", ...)
## S4 method for signature 'ExpressionSet'
dcScore(emat, condition, dc.method = "zscore", ...)
## S4 method for signature 'SummarizedExperiment'
dcScore(emat, condition, dc.method = "zscore", ...)
## S4 method for signature 'DGEList'
dcScore(emat, condition, dc.method = "zscore", ...)
```

dcScore 11

### Arguments

emat	a matrix, Matrix, data.frame, ExpressionSet, SummarizedExperiment or DGE-List
condition	a numeric, (with 1's and 2's representing a binary condition), a factor with 2 levels or a character representing 2 conditions
dc.method	a character, representing the method to use. Use $dcMethods()$ to get a list of methods
	possible arguments are cor.method, diffcoex.beta, ebcoexpress.useBWMC, ebcoexpress.plot, ldgm.lambda, ldgm.ntarget and ldgm.iter. See details

#### **Details**

When using data from sequencing experiments, make sure appropriate filtering for low counts and data transformation has been performed. Not doing so will affect estimation of correlation coefficients which most methods rely on.

Additional method specific parameters can be supplied to the function. cor.method can be set to either 'pearson' (default) or 'spearman' to determine the method to use for estimating correlations. These are the two measures currently supported in the package. We recommend using the 'spearman' correlation when dealing with sequencing data.

The beta parameter in the DiffCoEx method can be specified using diffcoex.beta (defaults to 6). This enable soft thresholding of correlations similar to WGCNA.

EBcoexpress specific parameters include ebcoexpress.useBWMC (defaults to TRUE) representing whether to use the bi-weight mid-correlation coefficient or not, and ebcoexpress.plot which plots the diagnostic plots if set to TRUE (defaults to FALSE).

LDGM specific parameters include ldgm.lambda, ldgm.ntarget and ldgm.iter. ldgm.lambda specifies the L1 regularisation parameter to use when fitting the model. This can be tuned and specified by the user. Alternatively, this can be tuned such that the resulting network has a specified number of edges. In this case, ldgm.ntarget should be specified instead. ldgm.iter is the maximum number of iterations to perform when tuning ldgm.lambda using ldgm.ntarget (defaults to 50).

EBcoexpress and GGM-based are implemented by providing interfaces to, or using functions from the EBcoexpress, GeneNet, and COSINE packages respectively. If using any of these methods, please **cite** the appropriate packages and the appropriate methodology articles.

#### Value

a matrix, of scores/statistics representing differential associations; p-values will be returned if FTGI is used and posterior probabilities if EBcoexpress is used.

#### See Also

dcMethods

12 dcTest

#### **Examples**

```
x <- matrix(rnorm(60), 2, 30)
cond <- rep(1:2, 15)
dcScore(x, cond) #defaults to zscore
dcScore(x, cond, dc.method = 'diffcoex')</pre>
```

dcTest

Statistical test for differential association analysis

## **Description**

Perform statistical tests for scores generated using dcScore. Selects appropriate tests for the different methods used in computing scores. The exact test is selected based on the scoring method used and cannot be manually specified. Available tests include the z-test and permutation tests. Parallel computation supported for the permutation test.

#### Usage

```
dcTest(dcscores, emat, condition, ...)
```

## **Arguments**

descores

a matrix, the result of the deScore function. The results should be passed as produced by the function and not modified in intermediate steps

emat

a matrix, data.frame, ExpressionSet, SummarizedExperiment or DGEList. This should be the one passed to deScore

condition

a numeric, (with 1's and 2's representing a binary condition), a factor with 2 levels or a character representing 2 conditions. This should be the one passed to deScore

... see details

#### **Details**

Ensure that the score matrix passed to this function is the one produced by dcScore. Any modification to the result matrix will cause this function to fail. This is intended as the test need to be performed on the entire score matrix, not subsets.

The appropriate test is chosen automatically based on the scoring method used. A z-test is performed for the z-score method while no tests are performed for DiffCoEx, EBcoexpress and FTGI. Permutation tests are performed for the remainder of methods by permutation sample labels. Statistics from a permutation are pooled such that statistics from all scores are used to evaluate a single observed score.

Additional method specific parameters can be supplied to the function when performing permutation tests. B specifies the number of permutations to be performed and defaults to 20.

If a cluster exists, computation in a permutation test will be performed in parallel (see examples).

dcZscore 13

## Value

a matrix, of p-values (or scores in the case of DiffCoEx and EBcoexpress) representing significance of differential associations. DiffCoEx will return scores as the publication specifies direct thresholding of scores and EBcoexpress returns posterior probabilities.

#### See Also

```
dcMethods, dcScore
```

#### **Examples**

```
x <- matrix(rnorm(60), 2, 30)
cond <- rep(1:2, 15)
scores <- dcScore(x, cond, dc.method = 'mindy')
dcTest(scores, emat = x, condition = cond)

## Not run:
#running in parallel
num_cores = 2
cl <- parallel::makeCluster(num_cores)
doSNOW::registerDoSNOW(cl) #or doParallel
set.seed(36) #for reproducibility
dcTest(scores, emat = x, condition = cond, B = 100)
parallel::stopCluster(cl)

## End(Not run)</pre>
```

dcZscore

DC analysis using the z-score method

## Description

This function packs the entire DC analysis pipeline using the z-score method. It simplifies the implementation of the analysis and increases the flexibility of the analysis (not just limited to all pairwise comparisons).

```
dcZscore(
  emat,
  condition,
  from = NULL,
  to = NULL,
  fdrthresh = 0.1,
  cor.method = c("spearman", "pearson")
)
```

14 getSimData

# Arguments

emat a matrix, Matrix, data.frame, ExpressionSet, SummarizedExperiment or DGEa numeric, (with 1's and 2's representing a binary condition), a factor with 2 condition levels or a character representing 2 conditions from a character vector, with the names of nodes from which comparisons need to be performed. a character vector, with the names of nodes to which comparisons need to be to performed. a numeric, specifying the FDR cutoff to apply to the inferred network. fdrthresh cor.method a character, either 'spearman' (default) or 'pearson' specifying the correlation computation method to use.

#### Value

an igraph object, containing the differential coexpression network.

#### **Examples**

```
x <- matrix(rnorm(60), 10, 30)
rownames(x) = 1:10
cond <- rep(1:2, 15)
dcZscore(x, cond)
dcZscore(x, cond, to = 1:2)</pre>
```

getSimData

Get data and conditions from a given knock-down (KD)

# Description

Retrieves the simulated expression matrix and sample classification for a specific knock-down experiment.

```
getSimData(simulation, cond.name = NULL, full = FALSE)
getConditionNames(simulation)

getTrueNetwork(
   simulation,
   cond.name = NULL,
   truth.type = c("association", "influence", "direct"),
   full = FALSE
)
```

getSimData 15

#### **Arguments**

a list, storing data and results generated from simulations

a character, indicating the knock-down to use to derive conditions. Multiple knock-downs (KDs) are performed per simulation. If NULL, the first KD is chosen

full a logical, indicating whether genes associated with the condition should be excluded. Defaults to FALSE and is recommended truth.type a character, specifying which level of the true network to retrieve: 'association'

(default), 'influence' or 'direct'

#### **Details**

Genes discarded when full is FALSE are those that are solely dependent on the condition. These genes are discarded from the analysis to focus on those that are differentially co-expressed, not coordinately co-expressed.

The names of all genes knocked-out can be retrieved using getConditionNames.

The direct, influence and association networks represent different levels of true differential networks. The direct network contains differential regulatory interactions present in the original network. The influence network includes upstream interactions and the association network includes non-causative differential interactions.

#### Value

a list, containing emat, a matrix representing the expression data, condition, a numeric containing the classification of samples, and , condition\_c, a numeric containing the expression levels of the KD gene (continuous condition) for getSimData; the names of all genes that are KD for getConditionNames; and an adjacency matrix for getTrueNetwork.

#### **Functions**

- getSimData: get the expression matrix and sample classification
- getConditionNames: get names of the conditions (KDs)
- getTrueNetwork: get the true differential network

## See Also

dcScore

```
data(sim102)
KDs <- getConditionNames(sim102)
#get simulated data
simdata <- getSimData(sim102, KDs[2])
cond <- simdata$condition
emat <- simdata$emat</pre>
```

perfMethods

```
zscores <- dcScore(emat, cond)
#get the true network to evaluate against
truenet <- getTrueNetwork(sim102, KDs[2], truth.type = 'association')</pre>
```

mi.ap

Mutual information using adaptive partitioning

#### **Description**

Computes the mutual information between all pairs of variables in the matrix (along the columns). Variables are discretised using the adaptive partitioning algorithm

# Usage

```
mi.ap(mat)
```

## **Arguments**

mat

a numeric matrix

#### Value

matrix of pairwise mutual information estimates

### **Examples**

```
x <- matrix(rnorm(200), 100, 2)
mi.ap(x)</pre>
```

perfMethods

Get names of performance metric methods

# Description

Returns a list of performance metrics

### Usage

```
perfMethods()
```

#### Value

names of methods implemented

```
perfMethods()
```

performanceMeasure 17

performanceMeasure	Performance metrics to evaluate classification	
--------------------	--	--

# Description

Quantify the performance of a classification algorithm. Predictions and truth both have to be binary.

### Usage

```
performanceMeasure(pred, obs, perf.method = "f.measure", ...)
```

# Arguments

pred	a logical or numeric, where 0 and FALSE represent control, and, 1 and TRUE represent cases
obs	a logical or numeric, where 0 and FALSE represent control, and, 1 and TRUE represent cases $$
perf.method	a character, specifying the method to use. Available methods can be accessed using ${\tt perfMethods}$
	additional parameters to methods. see details

## **Details**

The F-measure requires the beta parameter which can be specified using f.beta which defaults to 1 thereby computing the F1-measure.

# Value

```
a numeric, representing the performance
```

# See Also

```
perfMethods
```

```
pred <- sample(0:1, 100, replace = TRUE, prob = c(0.75, 0.25)) obs <- sample(0:1, 100, replace = TRUE, prob = c(0.75, 0.25)) #compute the F1 and F2 scores f1 <- performanceMeasure(pred, obs) f2 <- performanceMeasure(pred, obs, f.beta = 2)
```

18 plotSimNetwork

plotSimNetwork

Plot source and true differential networks from simulations

#### **Description**

Plots either the source network or the true differential network for all KDs performed in the simulation. KD nodes are coloured with their resulting differential networks coloured accordingly.

# Usage

```
plotSimNetwork(
   simulation,
   what = c("source", "direct", "influence", "association"),
   ...
)
```

## **Arguments**

#### **Details**

The direct, influence and association networks represent different levels of true differential networks. The direct network contains differential regulatory interactions present in the original network. The influence network includes upstream interactions and the association network includes non-causative differential interactions.

#### Value

a plot of the network

#### See Also

```
plot.igraph
```

```
data(sim102)
plotSimNetwork(sim102)
plotSimNetwork(sim102, what = 'direct')
plotSimNetwork(sim102, what = 'influence')
plotSimNetwork(sim102, what = 'association')
```

sim102

sim102

Simulated expression data with knock-outs

#### **Description**

A dataset containing simulated expression dataset. Data is simulated using a dynamical systems model from a network sampled from the S. Cerevisiae regulatory network. The dataset is a list containing the results from the simulation, and other information generated subsequently.

#### Usage

sim102

#### **Format**

A named list with 14 elements:

**simitr** a numeric, indicating the iteration of the simulation (a total of 1000 were performed and 812 converged)

**scores** an S4 Matrix, containing vectorised inference scores of applying the methods implemented in the package. These are precomputed predictions

**inputmodels** a named list, storing the parameters used to sample the initial values of input genes. Proportions, means and variances of each gene is stored for each gene

**staticnet** an igraph object, storing the initial regulatory network (150 node network)

**infnet** an igraph object, representing the true differential network as determined using sensitivity analysis of the model

**netlayout** a matrix (150 x 2), storing the (x, y) positions of nodes for laying out the graph

infdens a numeric, network density of the true differential association network

**numinput** a numeric, the number of input genes in the regulatory network. These are genes that have no regulators therefore need to be pre-defined

**numbimodal** a numeric, the number of input genes that are knocked-down therefore have a bimodal distribution

**numtfs** a numeric, the number of genes in the network that regulate any other gene (are TFs)

**numcotargets** a numeric, the number of genes that are co-regulated, i.e. regulated by more than one TF

**data** an S4 Matrix, the expression data with samples along the columns and genes along the rows. Condition classification (KD vs WT) are stored as attributes of this object

**triplets** a data frame, consisting of gene triplets representing TF- Target associations conditioned on the gene knocked-down. Triplets are annotated for being in either the direct, influence and association networks

sensmat an S4 Matrix, sensitivities of genes to TFs based on perturbation analysis of the simulation model

#### **Source**

LINK TO PAPERRRR

# **Index**

```
* classif
                                                    plot.igraph, 9, 18
    dcanr-package, 2
                                                    plotSimNetwork, 3, 18
* datasets
                                                    sim102, 19
    sim102, 19
* graphs
    dcanr-package, 2
* internal
    dcanr-package, 2
cor.pairs, 3, 4
dcAdjust, 3, 4, 7, 9
dcanr (dcanr-package), 2
dcanr-package, 2
dcEvaluate, 3, 5
dcMethods, 3, 6, 9, 11, 13
dcNetwork, 3, 7, 9
dcPipeline, 3, 6, 8
dcScore, 3, 7, 9, 10, 13, 15
{\tt dcScore}, {\tt data.frame-method}\,({\tt dcScore}),\, 10
dcScore, DGEList-method (dcScore), 10
dcScore,ExpressionSet-method(dcScore),
dcScore, Matrix-method (dcScore), 10
dcScore, matrix-method (dcScore), 10
{\tt dcScore}, {\tt SummarizedExperiment-method}
         (dcScore), 10
dcTest, 3, 5, 7, 9, 12
dcZscore, 13
getConditionNames, 3
getConditionNames (getSimData), 14
getSimData, 3, 14
getTrueNetwork, 3
getTrueNetwork (getSimData), 14
mi.ap, 3, 16
p.adjust, 5
perfMethods, 3, 6, 16, 17
performanceMeasure, 3, 6, 17
```