# Package 'maSigPro'

May 15, 2024

Type Package
<b>Title</b> Significant Gene Expression Profile Differences in Time Course Gene Expression Data
Version 1.77.0
Author Ana Conesa and Maria Jose Nueda
Maintainer Maria Jose Nueda <mj.nueda@ua.es></mj.nueda@ua.es>
<b>Description</b> maSigPro is a regression based approach to find genes for which there are significant gene expression profile differences between experimental groups in time course microarray and RNA-Seq experiments.
<b>Depends</b> R (>= $2.3.1$ )
Imports Biobase, graphics, grDevices, venn, mclust, stats, MASS
LazyLoad yes
<b>License</b> GPL (>= 2)
biocViews Microarray, RNA-Seq, Differential Expression, TimeCourse
git_url https://git.bioconductor.org/packages/maSigPro
git_branch devel
git_last_commit 6d12731
git_last_commit_date 2024-04-30
Repository Bioconductor 3.20
Date/Publication 2024-05-15
Contents
average.rows       2         data.abiotic       3         edesign.abiotic       5         edesignCT       6         edesignDR       6         get.siggenes       7         getDS       11

2 average.rows

	getDSPatterns	12
	i.rank	14
	ISOdata	14
	ISOdesign	15
	IsoModel	16
	IsoPlot	18
	make.design.matrix	19
	e	20
	· · ·	21
	NBdesign	22
	p.vector	
	•	25
	PlotProfiles	28
		29
	e	31
	•	32
	see.genes	33
	seeDS	36
	stepback	38
	stepfor	39
	suma2Venn	41
	T.fit	42
	tableDS	44
	two.ways.stepback	46
	two.ways.stepfor	47
	two.ways.step101	7/
Index		50

## Description

average.rows

average.rows matches rownames of a matrix to a match vector and performs averaging of the rows by the index provided by an index vector.

Average rows by match and index

## Usage

```
average.rows(x, index, match, r = 0.7)
```

## Arguments

x	a matrix
index	index vector indicating how rows must be averaged
match	match vector for indexing rows
r	minimal correlation value between rows to compute average

data.abiotic 3

## **Details**

rows will be averaged only if the pearson correlation coefficient between all rows of each given index is greater than r. If not, that group of rows is discarded in the result matrix.

#### Value

a matrix of averaged rows

#### Author(s)

Ana Conesa and Maria Jose Nueda, <mj.nueda@ua.es>

## **Examples**

```
## create data matrix for row averaging x \leftarrow matrix(rnorm(30), nrow = 6, ncol = 5) rownames(x) \leftarrow paste("ID", c(1, 2, 11, 12, 19, 20), sep = "") i \leftarrow paste("g", rep(c(1:10), each = 2), sep = "") # index vector m \leftarrow paste("ID", c(1:20), sep = "") # match vector average.rows(x, i, m, r = 0)
```

data.abiotic

Gene expression data potato abiotic stress

## **Description**

data.abiotic contains gene expression of a time course microarray experiment where potato plants were submitted to 3 different abiotic stresses.

## Usage

```
data(data.abiotic)
```

## **Format**

A data frame with 1000 observations on the following 36 variables.

```
Control_3H_1 a numeric vector

Control_3H_2 a numeric vector

Control_9H_1 a numeric vector

Control_9H_2 a numeric vector

Control_9H_3 a numeric vector

Control_27H_1 a numeric vector

Control_27H_1 a numeric vector
```

4 data.abiotic

```
Control_27H_3 a numeric vector
Cold_3H_1 a numeric vector
Cold_3H_2 a numeric vector
Cold_3H_3 a numeric vector
Cold_9H_1 a numeric vector
Cold_9H_2 a numeric vector
Cold_9H_3 a numeric vector
Cold_27H_1 a numeric vector
Cold_27H_2 a numeric vector
Cold_27H_3 a numeric vector
Heat_3H_1 a numeric vector
Heat_3H_2 a numeric vector
Heat_3H_3 a numeric vector
Heat_9H_1 a numeric vector
Heat_9H_2 a numeric vector
Heat_9H_3 a numeric vector
Heat_27H_1 a numeric vector
Heat_27H_2 a numeric vector
Heat_27H_3 a numeric vector
Salt_3H_1 a numeric vector
Salt_3H_2 a numeric vector
Salt_3H_3 a numeric vector
Salt_9H_1 a numeric vector
Salt_9H_2 a numeric vector
Salt_9H_3 a numeric vector
Salt_27H_1 a numeric vector
Salt_27H_2 a numeric vector
Salt_27H_3 a numeric vector
```

#### **Details**

This data set is part of a larger experiment in wich gene expression was monitored in both roots and leaves using a 11K cDNA potato chip. This example data set contains a ramdom subset of 1000 genes of the leave study.

#### References

Rensink WA, Iobst S, Hart A, Stegalkina S, Liu J, Buell CR. Gene expression profiling of potato responses to cold, heat, and salt stress. Funct Integr Genomics. 2005 Apr 22.

## **Examples**

data(data.abiotic)

edesign.abiotic 5

edesign.abiotic

Experimental design potato abiotic stress

## **Description**

edesign.abiotic contains experimental set up of a time course microarray experiment where potato plants were submitted to 3 different abiotic stresses.

## Usage

```
data(edesign.abiotic)
```

#### **Format**

```
A matrix with 36 rows and 6 columns rows [1:36] "Control 3h 1" "Control 3h 2" "Control 3h 3" "Control 9h 1" ... columns [1:6] "Time" "Replicates" "Control" "Cold" "Heat" "Salt"
```

## **Details**

Arrays are given in rows and experiment descriptors are given in columns. Row names contain array names.

"Time" indicates the values that variable Time takes in each hybridization.

"Replicates" is an index indicating replicate hyridizations, i.e. hybridizations are numbered, giving replicates the same number.

"Control", "Cold", "Heat" and "Salt" columns indicate array assignment to experimental groups, coding with 1 and 0 whether each array belongs to that group or not.

#### References

Rensink WA, Iobst S, Hart A, Stegalkina S, Liu J, Buell CR. Gene expression profiling of potato responses to cold, heat, and salt stress. Funct Integr Genomics. 2005 Apr 22.

```
data(edesignCR)
```

6 edesignDR

edesignCT

Experimental design with a shared time

## Description

edesignCT contains the experimental set up of a time course microarray experiment where there is a common starting point for the different experimental groups.

## Usage

```
data(edesignCT)
```

#### **Format**

```
A matrix with 32 rows and 7 colums
rows [1:32] "Array1" "Array2" "Array3" "Array4" ...
columns [1:7] "Time" "Replicates" "Control" "Tissue1" "Tissue2" "Tissue3" "Tissue4"
```

## **Details**

Arrays are given in rows and experiment descriptors are given in columns. Row names contain array names.

"Time" indicates the values that variable Time takes in each hybridization. There are 4 time points, which allows an up to 3 degree regression polynome.

"Replicates" is an index indicating replicate hyridizations, i.e. hybridizations are numbered, giving replicates the same number.

"Control", "Tissue1", "Tissue2", "Tissue3" and "Tissue4" columns indicate array assignment to experimental groups, coding with 1 and 0 whether each array belongs to that group or not.

## **Examples**

```
data(edesignCT)
```

edesignDR

Experimental design with different replicates

## **Description**

edesignDR contains experimental set up of a replicated time course microarray experiment where rats were submitted to 3 different dosis of a toxic compound. A control and an placebo treatments are also present in the experiment.

## Usage

```
data(edesignDR)
```

#### **Format**

```
A matrix with 54 rows and 7 columns rows [1:54] "Array1" "Array2" "Array3" "Array4" ... columns [1:7] "Time" "Replicates" "Control" "Placebo" "Low" "Medium" "High"
```

## **Details**

Arrays are given in rows and experiment descriptors are given in columns. Row names contain array names.

"Time" indicates the values that variable Time takes in each hybridization.

"Replicates" is an index indicating replicate hyridizations, i.e. hybridizations are numbered, giving replicates the same number.

"Control", "Placebo", "Low", "Medium" and "High" columns indicate array assignment to experimental groups, coding with 1 and 0 whether each array belongs to that group or not.

#### References

Heijne, W.H.M.; Stierum, R.; Slijper, M.; van Bladeren P.J. and van Ommen B.(2003). Toxicogenomics of bromobenzene hepatotoxicity: a combined transcriptomics and proteomics approach. Biochemical Pharmacology 65 857-875.

## **Examples**

data(edesignDR)

get.siggenes Extract significant genes for sets of variables in time series gene expression experiments

## **Description**

This function creates lists of significant genes for a set of variables whose significance value has been computed with the T. fit function.

## Usage

## **Arguments**

tstep	a T. fit object	
rsq	cut-off level at the R-squared value for the stepwise regression fit. Only genes with R-squared more than rsq are selected	
add.IDs	logical indicating whether to include additional gene id's in the result	
IDs	matrix containing additional gene id information (required when add. ${\tt IDs}$ is ${\tt TRUE})$	
matchID.col	number of matching column in matrix IDs for adding genes ids	
only.names	logical. If TRUE, expression values are ommitted in the results	
vars	variables for which to extract significant genes (see details)	
significant.in	tercept	
	experimental groups for which significant intercept coefficients are considered (see details)	
groups.vector	required when vars is "groups".	
trat.repl.spots		
	treatment given to replicate spots. Possible values are "none" and "average"	
index	argument of the average.rows function to use when trat.repl.spots is "average"	
match	argument of the average.rows function to use when trat.repl.spots is "average"	
r	minimun pearson correlation coefficient for replicated spots profiles to be averaged	

#### **Details**

There are 3 possible values for the vars argument:

"all": generates one single matrix or gene list with all significant genes.

"each": generates as many significant genes extractions as variables in the general regression model. Each extraction contains the significant genes for that variable.

"groups": generates a significant genes extraction for each experimental group.

The difference between "each" and "groups" is that in the first case the variables of the same group (e.g. "TreatmentA" and "time\*TreatmentA") will be extracted separately and in the second case jointly.

When add. IDs is TRUE, a matrix of gene ids must be provided as argument of IDs, the matchID.col column of which having same levels as in the row names of sig.profiles. The option only.names is TRUE will generate a vector of significant genes or a matrix when add. IDs is set also to TRUE.

When trat.repl.spots is "average", match and index vectors are required for the average.rows function. In gene expression data context, the index vector would contain geneIDs and indicate which spots are replicates. The match vector is used to match these genesIDs to rows in the significant genes matrix, and must have the same levels as the row names of sig.profiles.

The argument significant.intercept modulates the treatment for intercept coefficients to apply for selecting significant genes when vars equals "groups". There are three possible values: "none", no significant intercept (differences) are considered for significant gene selection, "dummy", includes genes with significant intercept differences between control and experimental groups, and "all" when both significant intercept coefficient for the control group and significant intercept differences are considered for selecting significant genes.

add. IDs = TRUE and trat.repl.spots = "average" are not compatible argumet values. add. IDs = TRUE and only.names = TRUE are compatible argumet values.

#### Value

summary a vector or matrix listing significant genes for the variables given by the function

parameters

sig.genes a list with detailed information on the significant genes found for the variables

given by the function parameters. Each element of the list is also a list contain-

ing:

sig.profiles: expression values of significant genes

coefficients: regression coefficients of the adjusted models

groups.coeffs: regression coefficients of the impiclit models of each experi-

mental group

sig.pvalues: p-values of the regression coefficients for significant genes

g: number of genes

...: arguments passed by previous functions

## Author(s)

Ana Conesa and Maria Jose Nueda, <mj.nueda@ua.es>

## References

Conesa, A., Nueda M.J., Alberto Ferrer, A., Talon, T. 2006. maSigPro: a Method to Identify Significant Differential Expression Profiles in Time-Course Microarray Experiments. Bioinformatics 22, 1096-1102

```
#### GENERATE TIME COURSE DATA
## generate n random gene expression profiles of a data set with
## one control plus 3 treatments, 3 time points and r replicates per time point.
```

```
tc.GENE <- function(n, r,</pre>
             var11 = 0.01, var12 = 0.01, var13 = 0.01,
             var21 = 0.01, var22 = 0.01, var23 = 0.01,
             var31 = 0.01, var32 = 0.01, var33 = 0.01,
             var41 = 0.01, var42 = 0.01, var43 = 0.01,
             a1 = 0, a2 = 0, a3 = 0, a4 = 0,
             b1 = 0, b2 = 0, b3 = 0, b4 = 0,
             c1 = 0, c2 = 0, c3 = 0, c4 = 0)
{
 tc.dat <- NULL
 for (i in 1:n) {
   Ctl <- c(rnorm(r, a1, var11), rnorm(r, b1, var12), rnorm(r, c1, var13)) # Ctl group
   Tr1 \leftarrow c(rnorm(r, a2, var21), rnorm(r, b2, var22), rnorm(r, c2, var23)) # Tr1 group
   Tr2 <- c(rnorm(r, a3, var31), rnorm(r, b3, var32), rnorm(r, c3, var33)) # Tr2 group
   Tr3 <- c(rnorm(r, a4, var41), rnorm(r, b4, var42), rnorm(r, c4, var43)) # Tr3 group
   gene <- c(Ctl, Tr1, Tr2, Tr3)</pre>
    tc.dat <- rbind(tc.dat, gene)</pre>
 }
 tc.dat
}
## Create 270 flat profiles
flat <- tc.GENE(n = 270, r = 3)
## Create 10 genes with profile differences between Ctl and Tr1 groups
twodiff \leftarrow tc.GENE (n = 10, r = 3, b2 = 0.5, c2 = 1.3)
## Create 10 genes with profile differences between Ctl, Tr2, and Tr3 groups
threediff <- tc.GENE(n = 10, r = 3, b3 = 0.8, c3 = -1, a4 = -0.1, b4 = -0.8, c4 = -1.2)
## Create 10 genes with profile differences between Ctl and Tr2 and different variance
vardiff \leftarrow tc.GENE(n = 10, r = 3, a3 = 0.7, b3 = 1, c3 = 1.2, var32 = 0.03, var33 = 0.03)
## Create dataset
tc.DATA <- rbind(flat, twodiff, threediff, vardiff)</pre>
rownames(tc.DATA) <- paste("feature", c(1:300), sep = "")</pre>
colnames(tc.DATA) <- paste("Array", c(1:36), sep = "")</pre>
tc.DATA [sample(c(1:(300*36)), 300)] <- NA # introduce missing values
#### CREATE EXPERIMENTAL DESIGN
Time \leftarrow rep(c(rep(c(1:3), each = 3)), 4)
Replicates \leftarrow rep(c(1:12), each = 3)
Control <- c(rep(1, 9), rep(0, 27))
Treat1 <- c(rep(0, 9), rep(1, 9), rep(0, 18))
Treat2 <- c(rep(0, 18), rep(1, 9), rep(0,9))
Treat3 <- c(rep(0, 27), rep(1, 9))
edesign <- cbind(Time, Replicates, Control, Treat1, Treat2, Treat3)</pre>
rownames(edesign) <- paste("Array", c(1:36), sep = "")</pre>
tc.p <- p.vector(tc.DATA, design = make.design.matrix(edesign), Q = 0.01)
tc.tstep <- T.fit(data = tc.p , alfa = 0.05)
## This will obtain sigificant genes per experimental group
## which have a regression model Rsquared > 0.9
tc.sigs <- get.siggenes (tc.tstep, rsq = 0.9, vars = "groups")
## This will obtain all sigificant genes regardless the Rsquared value.
```

getDS 11

getDS Extract lists of significant isoforms from Differentially Spliced Genes (DSG)

## **Description**

getDS creates lists of significant isoforms from Differentially Spliced Genes (DSG)

## Usage

```
getDS(Model, vars="all", rsq=0.4)
```

## **Arguments**

Model a IsoModel object

vars argument of the get. siggenes function applied to isoforms

rsq cut-off level at the R-squared value for the stepwise regression fit. Only isoforms

with R-squared more than rsq are selected

#### **Details**

There are 3 possible values for the vars argument: "all", "each" and "groups". See get.siggenes.

#### Value

In the console a summary of the selection is printed.

Model a IsoModel object to be used in the following steps
get2 a get.siggenes object to be used in the following steps
DSG Names of the selected genes: Differentially Spliced Genes

DET Names of the selected Isoforms: Differentally Expressed Transcripts

List0 a list with the names of Differentially Spliced Genes without Isoforms with R-

squared higher than rsq

NumIso.by.gene Number of selected Isoforms for each Differentially Spliced Gene

## Author(s)

Maria Jose Nueda, <mj.nueda@ua.es>

12 getDSPatterns

#### References

Nueda, M.J., Martorell, J., Marti, C., Tarazona, S., Conesa, A. 2018. Identification and visualization of differential isoform expression in RNA-seq time series. Bioinformatics. 34, 3, 524-526.

Nueda, M.J., Tarazona, S., Conesa, A. 2014. Next maSigPro: updating maSigPro bioconductor package for RNA-seq time series. Bioinformatics, 30, 2598-602.

Conesa, A., Nueda M.J., Alberto Ferrer, A., Talon, T. 2006. maSigPro: a Method to Identify Significant Differential Expression Profiles in Time-Course Microarray Experiments. Bioinformatics 22, 1096-1102.

#### See Also

```
get.siggenes, IsoModel
```

## **Examples**

```
data(ISOdata)
data(ISOdesign)
mdis <- make.design.matrix(ISOdesign)
MyIso <- IsoModel(data=ISOdata[,-1], gen=ISOdata[,1], design=mdis, counts=TRUE)

Myget <- getDS(MyIso)
Myget$DSG
Myget$DET

see <- seeDS(Myget, cluster.all=FALSE, k=6)
table <- tableDS(see)
table$IsoTable</pre>
```

getDSPatterns

Lists of genes with Isoforms in different clusters

## **Description**

getDSPatterns is a function that makes a list with the names of genes identified with tableDS function.

## Usage

```
getDSPatterns(tableDS, Cluster.Major, Cluster.minor)
```

## **Arguments**

tableDS a tableDS object

Cluster.Major Number of the cluster where the major isoform belongs to

Cluster.minor Number(s) of the cluster(s) where the minor isoform(s) belongs to (see details)

getDSPatterns 13

#### **Details**

When minor isoforms belong to different clusters, tableDS codifies them using "&". For instance: clusters 1 and 2, will be represented as "1&2". In such cases quotation marks must be used (see examples). When minor isoforms are only in one cluster there is no need to use quotation marks.

#### Value

A vector with the names of the genes.

#### Author(s)

Maria Jose Nueda, <mj.nueda@ua.es>

#### References

Nueda, M.J., Martorell, J., Marti, C., Tarazona, S., Conesa, A. 2018. Identification and visualization of differential isoform expression in RNA-seq time series. Bioinformatics. 34, 3, 524-526.

Nueda, M.J., Tarazona, S., Conesa, A. 2014. Next maSigPro: updating maSigPro bioconductor package for RNA-seq time series. Bioinformatics, 30, 2598-602.

Conesa, A., Nueda M.J., Alberto Ferrer, A., Talon, T. 2006. maSigPro: a Method to Identify Significant Differential Expression Profiles in Time-Course Microarray Experiments. Bioinformatics 22, 1096-1102.

## See Also

```
tableDS, IsoModel
```

```
data(ISOdata)
data(ISOdesign)
mdis <- make.design.matrix(ISOdesign)
MyIso <- IsoModel(data=ISOdata[,-1], gen=ISOdata[,1], design=mdis, counts=TRUE)
Myget <- getDS(MyIso)
see <- seeDS(Myget, cluster.all=FALSE, k=6)
table <- tableDS(see)
table$IsoTable

getDSPatterns(table, 1, 4)
getDSPatterns(table, "1", "4") #will give the same result.

getDSPatterns(table, 1, "185")</pre>
```

14 ISOdata

i.rank

Ranks a vector to index

## Description

Ranks the values in a vector to sucessive values. Ties are given the same value.

## Usage

```
i.rank(x)
```

## **Arguments**

Х

vector

## Value

Vector of ranked values

## Author(s)

Ana Conesa and Maria Jose Nueda, <mj.nueda@ua.es>

## See Also

rank,order

## **Examples**

```
i.rank(c(1, 1, 1, 3, 3, 5, 7, 7, 7))
```

ISOdata

RNA-Seq dataset example for isoforms

## Description

ISOdata contains an example of RNA-Seq data at Isoform level.

## Usage

```
data(ISOdata)
```

## **Format**

A data frame with 2782 rows and 37 columns with RNA-Seq data.

ISOdesign 15

## **Details**

Rows correspond to 2782 isoforms belonging to 1000 gene.

First column is the name of the gene each isoform belongs to.

Remaining columns are the RNA-Seq data samples associated to 3 replicates of 12 experimental conditions.

## **Examples**

```
data(ISOdata)
data(ISOdesign)

mdis <- make.design.matrix(ISOdesign)

MyIso <- IsoModel(data=ISOdata[,-1], gen=ISOdata[,1], design=mdis, counts=TRUE)</pre>
```

ISOdesign

Experimental design for ISOdata dataset example

## Description

ISOdesign is the experimental design to apply ISOmaSigPro to ISOdata dataset example.

## Usage

```
data(ISOdesign)
```

#### **Format**

A matrix with 36 rows and 4 colums

```
 rownames (ISOdesign) "Gr1\_0h\_1" "Gr1\_0h\_2" "Gr1\_0h\_3" "Gr1\_2h\_1" "Gr1\_2h\_2" "Gr1\_2h\_3" "Gr1\_6h\_1" "Gr1\_6h\_2" "Gr1\_6h\_3" "Gr1\_12h\_1" "Gr1\_12h_2" "Gr1\_12h_3" "Gr1\_18h\_1" "Gr1\_18h_2" "Gr1\_18h_3" "Gr1\_24h_1" "Gr1\_24h_2" "Gr1\_24h_3" "Gr2\_0h_1" "Gr2\_0h_2" "Gr2\_0h_3" "Gr2\_2h_1" "Gr2\_2h_2" "Gr2\_2h_3" "Gr2\_6h_1" "Gr2\_6h_2" "Gr2_6h_3" "Gr2_12h_1" "Gr2\_12h_2" "Gr2_12h_3" "Gr2\_18h_1" "Gr2\_18h_2" "Gr2\_18h_3" "Gr2\_24h_1" "Gr2\_24h_2" "Gr2\_24h_3" \\ colnames (ISOdesign) "time" "replicate" "Group1" "Group2"
```

#### **Details**

Samples are given in rows and experiment descriptors are given in columns. Row names contain sample names.

"time" indicates the values that variable Time takes in each experimental condition. There are 6 time points.

<sup>&</sup>quot;replicate" is an index indicating the same experimental condition.

<sup>&</sup>quot;Group1" and "Group2" columns indicate assignment to experimental groups, coding with 1 and 0 whether each sample belongs to that group or not.

16 IsoModel

## **Examples**

```
data(ISOdata)
data(ISOdesign)
mdis <- make.design.matrix(ISOdesign)</pre>
MyIso <- IsoModel(data=ISOdata[,-1], gen=ISOdata[,1], design=mdis, counts=TRUE)</pre>
```

IsoModel Detection of genes with Isoforms with different gene expression in time course experiments

## Description

IsoModel Performs a model comparison for each gene to detect genes with different trends in time course experiments and applies maSigPro to the Isoforms belonging to selected genes.

## Usage

```
IsoModel(data, gen, design = NULL, Q = 0.05, min.obs = 6, minorFoldfilter = NULL,
    counts = FALSE, family = NULL, theta = 10, epsilon = 1e-05)
```

## **Arguments**

data	matrix containing isoform expression. Isoforms must be in rows and experimental conditions in columns
gen	vector with the name of the gene each isoform belongs to
design	design matrix for the regression fit such as that generated by the ${\tt make.design.matrix}$ function
Q	significance level
min.obs	cases with less than this number of true numerical values will be excluded from the analysis. Minimum value to estimate the model is (degree+1)xGroups+1. Default is 6.
minorFoldfilte	er
	fold expression difference between minor isoforms and the most expressed isoform to exclude minor isoforms from analysis. Default NULL
counts	a logical indicating whether your data are counts
family	the distribution function to be used in the glm model. It must be specified as a function: gaussian(), poisson(), negative.binomial(theta) If NULL family will be negative.binomial(theta) when counts=TRUE or gaussian() when counts=FALSE
theta	theta parameter for negative.binomial family
epsilon	argument to pass to glm. control, convergence tolerance in the iterative process

to estimate de glm model

IsoModel 17

## **Details**

rownames(design) and colnames(data) must be identical vectors and indicate experimental condition names.

rownames (data) should contain unique isoform IDs.

colnames(design) are the given names for the variables in the regression model.

## Value

data	input data matrix to be used in the following steps
gen	input gen vector to be used in the following steps
design	input design matrix to be used in the following steps
DSG	Names of the selected genes: Differentially Spliced Genes
pvector	p.vector output of isoforms that belong to selected.genes
Tfit	Tfit output of isoforms that belong to selected.genes

#### Author(s)

Maria Jose Nueda, <mj.nueda@ua.es>

#### References

Nueda, M.J., Martorell, J., Marti, C., Tarazona, S., Conesa, A. 2018. Identification and visualization of differential isoform expression in RNA-seq time series. Bioinformatics. 34, 3, 524-526.

Nueda, M.J., Tarazona, S., Conesa, A. 2014. Next maSigPro: updating maSigPro bioconductor package for RNA-seq time series. Bioinformatics, 30, 2598-602.

Conesa, A., Nueda M.J., Ferrer, A., Talon, T. 2006. maSigPro: a Method to Identify Significant Differential Expression Profiles in Time-Course Microarray Experiments. Bioinformatics 22, 1096-1102.

## See Also

```
p.vector, T.fit
```

```
data(ISOdata)
data(ISOdesign)
mdis <- make.design.matrix(ISOdesign)

MyIso <- IsoModel(data=ISOdata[,-1], gen=ISOdata[,1], design=mdis, counts=TRUE)
Myget <- getDS(MyIso)
see <- seeDS(Myget, cluster.all=FALSE, k=6)
table <- tableDS(see)
table$IsoTable</pre>
```

18 IsoPlot

IsoPlot Plotting the isoform profiles of a specific gene by groups	
--	--

## **Description**

This function makes a plot with the isoforms of a specific gene splitting the different experimental groups.

## Usage

```
IsoPlot(get, name, only.sig.iso=FALSE, ylim=NULL, xlab = "Time",
ylab = "Expression value", points=TRUE, cex.main=3,cex.legend=1.5)
```

#### **Arguments**

get	a getDS object a cluster of flat Isoform
name	Name of the specific gen to show in the plot
only.sig.iso	TRUE when the plot is made only with statistically significant isoforms.
ylim	Range of the y axis of the desired plot. If it is NULL it will be computed automatically.
xlab	label for the x axis
ylab	label for the y axis
points	TRUE to plot points and lines. FALSE to plot only lines.
cex.main	graphical parameter magnification to be used for main
cex.legend	graphical parameter magnification to be used for legend

## **Details**

The plot can be made with all the available isoforms or only with the statistilly significant ones.

## Value

Plot of isoform profiles of a specific gene by groups.

#### Author(s)

Maria Jose Nueda, <mj.nueda@ua.es>

## References

Nueda, M.J., Martorell, J., Marti, C., Tarazona, S., Conesa, A. 2018. Identification and visualization of differential isoform expression in RNA-seq time series. Bioinformatics. 34, 3, 524-526.

Nueda, M.J., Tarazona, S., Conesa, A. 2014. Next maSigPro: updating maSigPro bioconductor package for RNA-seq time series. Bioinformatics, 30, 2598-602.

Conesa, A., Nueda M.J., Alberto Ferrer, A., Talon, T. 2006. maSigPro: a Method to Identify Significant Differential Expression Profiles in Time-Course Microarray Experiments. Bioinformatics 22, 1096-1102.

make.design.matrix 19

## See Also

```
getDS, IsoModel
```

## **Examples**

```
data(ISOdata)
data(ISOdesign)
mdis <- make.design.matrix(ISOdesign)
MyIso <- IsoModel(data=ISOdata[,-1], gen=ISOdata[,1], design=mdis, counts=TRUE)
Myget <- getDS(MyIso)
IsoPlot(Myget, "Gene1005", only.sig.iso=FALSE,cex.main=2,cex.legend=1)</pre>
```

 ${\tt make.design.matrix}$ 

Make a design matrix for regression fit of time series gene expression experiments

## **Description**

make.design.matrix creates the design matrix of dummies for fitting time series micorarray gene expression experiments.

## Usage

#### **Arguments**

edesign	matrix describing experimental design. Rows must be arrays and columns experiment descriptors
degree	the degree of the regression fit polynome. $degree = 1$ returns linear regression, $degree = 2$ returns quadratic regression, etc
time.col	column number in edesign containing time values. Default is first column
repl.col	column number in edesign containing coding for replicate arrays. Default is second column
group.cols	column numbers in edesign indicating the coding for each experimental group (treatment, tissue,). See details

## **Details**

rownames of edesign object should contain the arrays naming (i.e. array1, array2, ...). colnames of edesign must contain the names of experiment descriptors(i.e. "Time", "Replicates", "Treatment A", "Treatment B", etc.). for each experimental group a different column must be present in edesign, coding with 1 and 0 whether each array belongs to that group or not.

make.design.matrix returns a design matrix where rows represent arrays and column variables of time, dummies and their interactions for up to the degree given. Dummies show the relative effect of each experimental group related to the first one. Single dummies indicate the abcissa component of each group. \$Time\*dummy\$ variables indicate slope changes, \$Time^2\*dummy\$ indicates curvature changes. Higher grade values could model complex responses. In case experimental groups share a initial state (i.e. common time 0), no single dummies are modeled.

#### Value

dis design matrix of dummies for fitting time series

groups.vector vector coding the experimental group to which each variable belongs to

edesign edesign value passed as argument

## Author(s)

Ana Conesa and Maria Jose Nueda, <mj.nueda@ua.es>

#### References

Conesa, A., Nueda M.J., Alberto Ferrer, A., Talon, T. 2006. maSigPro: a Method to Identify Significant Differential Expression Profiles in Time-Course Microarray Experiments. Bioinformatics 22, 1096-1102

## **Examples**

```
data(edesign.abiotic, edesignCT)
make.design.matrix(edesign.abiotic) # quadratic model
make.design.matrix(edesignCT, degree = 3) # cubic model with common starting time point
```

maSigProUsersGuide

View maSigPro User's Guide

## Description

Finds the location of the maSigPro User's Guide and opens it.

#### Usage

```
maSigProUsersGuide(view=TRUE)
```

## **Arguments**

view

logical, to specify if the document is opened using the PDF document reader.

NBdata 21

## **Details**

The function vignette("maSigPro") will find the short maSigPro Vignette which describes how to obtain the maSigPro User's Guide. The User's Guide is not itself a true vignette because it is not automatically generated using Sweave during the package build process. This means that it cannot be found using vignette, hence the need for this special function.

If the operating system is other than Windows, then the PDF viewer used is that given by Sys.getenv("R\_PDFVIEWER"). The PDF viewer can be changed using Sys.putenv(R\_PDFVIEWER=).

#### Value

If vignette(view=TRUE), the PDF document reader is started and the User's Guide is opened. If vignette(view=FALSE), returns the file location.

## **Examples**

```
maSigProUsersGuide()
maSigProUsersGuide(view=FALSE)
```

**NBdata** 

RNA-Seq dataset example

## **Description**

NBdata contains a subset of a bigger normalized negative binomial simulated dataset.

## Usage

data(NBdata)

## **Format**

A data frame with 100 observations on 36 numeric variables.

#### **Details**

This dataset is part of a larger simulated and normalized dataset with 2 experimental groups, 6 time-points and 3 replicates. Simulation has been done by using a negative binomial distribution. The first 20 genes are simulated with changes among time.

## **Examples**

data(NBdata)

p.vector

NBdesign

Experimental design for RNA-Seq example

## **Description**

NBdesign contains a subset of a bigger normalized negative binomial simulated dataset.

## Usage

```
data(NBdesign)
```

#### **Format**

```
A matrix with 36 rows and 4 colums rows [1:36] "G1.T1.1" "G1.T1.2" "G1.T1.3" "G1.T2.1" ... columns [1:6] [1] "Time" "Replicates" "Group.1" "Group.2"
```

#### **Details**

Samples are given in rows and experiment descriptors are given in columns. Row names contain sample names.

"Time" indicates the values that variable Time takes in each experimental condition. There are 6 time points.

"Replicates" is an index indicating the same experimental condition.

"Group.1" and "Group.2" columns indicate assignment to experimental groups, coding with 1 and 0 whether each sample belongs to that group or not.

## **Examples**

```
data(NBdesign)
```

p.vector

Make regression fit for time series gene expression experiments

## **Description**

p. vector performs a regression fit for each gene taking all variables present in the model given by a regression matrix and returns a list of FDR corrected significant genes.

#### Usage

```
p.vector(data, design, Q = 0.05, MT.adjust = "BH", min.obs = 6,
counts=FALSE, family=NULL, theta=10, epsilon=0.00001, item="gene")
```

p.vector 23

## Arguments

data	matrix containing normalized gene expression data. Genes must be in rows and arrays in columns
design	design matrix for the regression fit such as that generated by the ${\tt make.design.matrix}$ function
Q	significance level
MT.adjust	argument to pass to p.adjust function indicating the method for multiple testing adjustment of p.value
min.obs	genes with less than this number of true numerical values will be excluded from the analysis. Minimum value to estimate the model is (degree+1)xGroups+1. Default is 6.
counts	a logical indicating whether your data are counts
family	the distribution function to be used in the glm model. It must be specified as a function: gaussian(), poisson(), negative.binomial(theta) If NULL family will be negative.binomial(theta) when counts=TRUE or gaussian() when counts=FALSE
theta	theta parameter for negative.binomial family
epsilon	argument to pass to glm. control, convergence tolerance in the iterative process to estimate de glm model
item	Name of the analysed item to show in the screen while p.vector is in process

## **Details**

rownames(design) and colnames(data) must be identical vectors and indicate array naming. rownames(data) should contain unique gene IDs. colnames(design) are the given names for the variables in the regression model.

## Value

SELEC	matrix containing the expression values for significant genes
p.vector	vector containing the computed p-values
G	total number of input genes
g	number of genes taken in the regression fit
FDR	p-value at FDR Q control when Benajamini & Holderberg (BH) correction is used
i	number of significant genes
dis	design matrix used in the regression fit
dat	matrix of expression value data used in the regression fit
	additional values from input parameters

## Author(s)

Ana Conesa and Maria Jose Nueda, <mj.nueda@ua.es>

24 p.vector

#### References

Conesa, A., Nueda M.J., Alberto Ferrer, A., Talon, T. 2006. maSigPro: a Method to Identify Significant Differential Expression Profiles in Time-Course Microarray Experiments. Bioinformatics 22, 1096-1102

#### See Also

```
T.fit.lm
```

#### CREATE EXPERIMENTAL DESIGN

```
#### GENERATE TIME COURSE DATA
## generates n random gene expression profiles of a data set with
## one control plus 3 treatments, 3 time points and r replicates per time point.
tc.GENE <- function(n, r,</pre>
             var11 = 0.01, var12 = 0.01, var13 = 0.01,
             var21 = 0.01, var22 = 0.01, var23 = 0.01,
             var31 = 0.01, var32 = 0.01, var33 = 0.01,
             var41 = 0.01, var42 = 0.01, var43 = 0.01,
             a1 = 0, a2 = 0, a3 = 0, a4 = 0,
             b1 = 0, b2 = 0, b3 = 0, b4 = 0,
             c1 = 0, c2 = 0, c3 = 0, c4 = 0)
{
 tc.dat <- NULL
 for (i in 1:n) {
   Ctl <- c(rnorm(r, a1, var11), rnorm(r, b1, var12), rnorm(r, c1, var13)) # Ctl group
   Tr1 <- c(rnorm(r, a2, var21), rnorm(r, b2, var22), rnorm(r, c2, var23)) # Tr1 group
   Tr2 <- c(rnorm(r, a3, var31), rnorm(r, b3, var32), rnorm(r, c3, var33)) # Tr2 group
   Tr3 <- c(rnorm(r, a4, var41), rnorm(r, b4, var42), rnorm(r, c4, var43)) # Tr3 group
   gene <- c(Ctl, Tr1, Tr2, Tr3)
    tc.dat <- rbind(tc.dat, gene)</pre>
 tc.dat
}
## Create 270 flat profiles
flat <- tc.GENE(n = 270, r = 3)
## Create 10 genes with profile differences between Ctl and Tr1 groups
twodiff \leftarrow tc.GENE (n = 10, r = 3, b2 = 0.5, c2 = 1.3)
## Create 10 genes with profile differences between Ctl, Tr2, and Tr3 groups
threediff <- tc.GENE(n = 10, r = 3, b3 = 0.8, c3 = -1, a4 = -0.1, b4 = -0.8, c4 = -1.2)
## Create 10 genes with profile differences between Ctl and Tr2 and different variance
vardiff < -tc.GENE(n = 10, r = 3, a3 = 0.7, b3 = 1, c2 = 1.3, var32 = 0.03, var33 = 0.03)
## Create dataset
tc.DATA <- rbind(flat, twodiff, threediff, vardiff)</pre>
rownames(tc.DATA) <- paste("feature", c(1:300), sep = "")
colnames(tc.DATA) <- paste("Array", c(1:36), sep = "")</pre>
tc.DATA [sample(c(1:(300*36)), 300)] <- NA # introduce missing values
```

PlotGroups 25

```
Time <- rep(c(rep(c(1:3), each = 3)), 4)
Replicates <- rep(c(1:12), each = 3)
Control <- c(rep(1, 9), rep(0, 27))
Treat1 <- c(rep(0, 9), rep(1, 9), rep(0, 18))
Treat2 <- c(rep(0, 18), rep(1, 9), rep(0,9))
Treat3 <- c(rep(0, 27), rep(1, 9))
edesign <- cbind(Time, Replicates, Control, Treat1, Treat2, Treat3)
rownames(edesign) <- paste("Array", c(1:36), sep = "")

tc.p <- p.vector(tc.DATA, design = make.design.matrix(edesign), Q = 0.05)
tc.p$i # number of significant genes
tc.p$SELEC # expression value of significant genes
tc.p$FDR # p.value at FDR control
tc.p$p.adjusted# adjusted p.values</pre>
```

PlotGroups

Function for plotting gene expression profile at different experimental groups

## **Description**

This function displays the gene expression profile for each experimental group in a time series gene expression experiment.

## Usage

```
PlotGroups(data, edesign = NULL, time = edesign[, 1], groups = edesign[,c(3:ncol(edesign))], repvect = edesign[, 2], show.lines = TRUE, show.fit = FALSE, dis = NULL, step.method = "backward", min.obs = 2, alfa = 0.05, nvar.correction = FALSE, summary.mode = "median", groups.vector = NULL, main = NULL, sub = NULL, xlab = "Time", ylab = "Expression value", item = NULL, ylim = NULL, pch = 21, col = NULL, legend = TRUE, cex.legend = 1,lty.legend = NULL,...)
```

#### **Arguments**

data	vector or matrix containing the gene expression data
edesign	matrix describing experimental design. Rows must be arrays and columns experiment descriptors
time	vector indicating time assignment for each array
groups	matrix indicating experimental group to which each array is assigned
repvect	index vector indicating experimental replicates
show.lines	logical indicating whether a line must be drawn joining plotted data points for reach group
show.fit	logical indicating whether regression fit curves must be plotted

dis regression design matrix

step.method stepwise regression method to fit models for cluster mean profiles. It can be ei-

ther "backward", "forward", "two.ways.backward" or "two.ways.forward"

min.obs minimal number of observations for a gene to be included in the analysis alfa significance level used for variable selection in the stepwise regression

nvar.correction

argument for correcting stepwise regression significance level. See T. fit

summary.mode the method to condensate expression information when more than one gene is

present in the data. Possible values are "representative" and "median"

groups.vector vector indicating experimental group to which each variable belongs

main plot main title
sub plot subtitle
vlab label for the x a

xlab label for the x axis ylab label for the y axis

item name of the analysed items to show

ylim range of the y axis

pch integer specifying type of points to plot

col a vector specifying colours to plot. If missing first naturals will be used legend logical indicating whether legend must be added when plotting profiles

cex.legend Expansion factor for legend

1ty.legend To add a coloured line in the legend ... other graphical function argument

#### **Details**

To compute experimental groups either a edesign object must be provided, or separate values must be given for the time, repvect and groups arguments.

When data is a matrix, the average expression value is displayed.

When there are array replicates in the data (as indicated by repvect), values are averaged by repvect.

PlotGroups plots one single expression profile for each experimental group even if there are more that one genes in the data set. The way data is condensated for this is given by summary.mode. When this argument takes the value "representative", the gene with the lowest distance to all genes in the cluster will be plotted. When the argument is "median", then median expression value is computed.

When show, fit is TRUE the stepwise regression fit for the data will be computed and the regression curves will be displayed.

If data is a matrix of genes and summary.mode is "median", the regression fit will be computed for the median expression value.

## Value

Plot of gene expression profiles by-group.

PlotGroups 27

#### Author(s)

Ana Conesa and Maria Jose Nueda, <mj.nueda@ua.es>

#### References

Conesa, A., Nueda M.J., Alberto Ferrer, A., Talon, T. 2005. maSigPro: a Method to Identify Significant Differential Expression Profiles in Time-Course Microarray Experiments.

#### See Also

**PlotProfiles** 

```
#### GENERATE TIME COURSE DATA
## generate n random gene expression profiles of a data set with
## one control plus 3 treatments, 3 time points and r replicates per time point.
tc.GENE <- function(n, r,</pre>
             var11 = 0.01, var12 = 0.01, var13 = 0.01,
             var21 = 0.01, var22 = 0.01, var23 = 0.01,
             var31 = 0.01, var32 = 0.01, var33 = 0.01,
             var41 = 0.01, var42 = 0.01, var43 = 0.01,
             a1 = 0, a2 = 0, a3 = 0, a4 = 0,
             b1 = 0, b2 = 0, b3 = 0, b4 = 0,
             c1 = 0, c2 = 0, c3 = 0, c4 = 0)
{
  tc.dat <- NULL
  for (i in 1:n) {
    Ctl <- c(rnorm(r, a1, var11), rnorm(r, b1, var12), rnorm(r, c1, var13)) # Ctl group
    Tr1 <- c(rnorm(r, a2, var21), rnorm(r, b2, var22), rnorm(r, c2, var23)) # Tr1 group
    Tr2 <- c(rnorm(r, a3, var31), rnorm(r, b3, var32), rnorm(r, c3, var33)) # Tr2 group
    Tr3 <- c(rnorm(r, a4, var41), rnorm(r, b4, var42), rnorm(r, c4, var43)) # Tr3 group
    gene <- c(Ctl, Tr1, Tr2, Tr3)
    tc.dat <- rbind(tc.dat, gene)</pre>
  }
  tc.dat
}
## create 10 genes with profile differences between Ctl, Tr2, and Tr3 groups
tc.DATA \leftarrow tc.GENE(n = 10,r = 3, b3 = 0.8, c3 = -1, a4 = -0.1, b4 = -0.8, c4 = -1.2)
rownames(tc.DATA) <- paste("gene", c(1:10), sep = "")</pre>
colnames(tc.DATA) <- paste("Array", c(1:36), sep = "")</pre>
#### CREATE EXPERIMENTAL DESIGN
Time <- rep(c(rep(c(1:3), each = 3)), 4)
Replicates \leftarrow rep(c(1:12), each = 3)
Ctl <- c(rep(1, 9), rep(0, 27))
Tr1 \leftarrow c(rep(0, 9), rep(1, 9), rep(0, 18))
Tr2 \leftarrow c(rep(0, 18), rep(1, 9), rep(0, 9))
Tr3 <- c(rep(0, 27), rep(1, 9))
```

28 PlotProfiles

```
PlotGroups (tc.DATA, time = Time, repvect = Replicates, groups = cbind(Ctl, Tr1, Tr2, Tr3))
```

PlotProfiles

Function for visualization of gene expression profiles

## Description

PlotProfiles displays the expression profiles of a group of genes.

## Usage

```
PlotProfiles(data, cond, cex.axis = 0.5, ylim = NULL, repvect, main = NULL, sub = NULL, color.mode = "rainbow", item = NULL)
```

## **Arguments**

data a matrix containing the gene expression data cond vector for x axis labeling, typically array names

cex.axis graphical parameter maginfication to be used for x axis in plotting functions

ylim index vector indicating experimental replicates repvect index vector indicating experimental replicates

main plot main title sub plot subtitle

color.mode color scale for plotting profiles. Can be either "rainblow" or "gray"

item Name of the analysed items to show

#### **Details**

The repvect argument is used to indicate with vertical lines groups of replicated arrays.

#### Value

Plot of experiment-wide gene expression profiles.

## Author(s)

Ana Conesa and Maria Jose Nueda, <mj.nueda@ua.es>

## References

Conesa, A., Nueda M.J., Alberto Ferrer, A., Talon, T. 2005. maSigPro: a Method to Identify Significant Differential Expression Profiles in Time-Course Microarray Experiments.

PodiumChange 29

## See Also

**PlotGroups** 

## **Examples**

```
#### GENERATE TIME COURSE DATA
## generate n random gene expression profiles of a data set with
## one control plus 3 treatments, 3 time points and r replicates per time point.
tc.GENE <- function(n, r,</pre>
             var11 = 0.01, var12 = 0.01, var13 = 0.01,
             var21 = 0.01, var22 = 0.01, var23 = 0.01,
             var31 = 0.01, var32 = 0.01, var33 = 0.01,
             var41 = 0.01, var42 = 0.01, var43 = 0.01,
             a1 = 0, a2 = 0, a3 = 0, a4 = 0,
             b1 = 0, b2 = 0, b3 = 0, b4 = 0,
             c1 = 0, c2 = 0, c3 = 0, c4 = 0)
{
  tc.dat <- NULL
  for (i in 1:n) {
   Ctl <- c(rnorm(r, a1, var11), rnorm(r, b1, var12), rnorm(r, c1, var13)) # Ctl group
   Tr1 <- c(rnorm(r, a2, var21), rnorm(r, b2, var22), rnorm(r, c2, var23)) # Tr1 group
   Tr2 <- c(rnorm(r, a3, var31), rnorm(r, b3, var32), rnorm(r, c3, var33)) # Tr2 group
   Tr3 <- c(rnorm(r, a4, var41), rnorm(r, b4, var42), rnorm(r, c4, var43)) # Tr3 group
   gene <- c(Ctl, Tr1, Tr2, Tr3)
    tc.dat <- rbind(tc.dat, gene)</pre>
  }
  tc.dat
}
## create 10 genes with profile differences between Ctl, Tr2, and Tr3 groups
tc.DATA \leftarrow tc.GENE(n = 10, r = 3, b3 = 0.8, c3 = -1, a4 = -0.1, b4 = -0.8, c4 = -1.2)
rownames(tc.DATA) <- paste("gene", c(1:10), sep = "")</pre>
colnames(tc.DATA) <- paste("Array", c(1:36), sep = "")</pre>
PlotProfiles (tc.DATA, cond = colnames(tc.DATA), main = "Time Course",
              repvect = rep(c(1:12), each = 3))
```

PodiumChange

Detection of Genes with switchs of their major isoforms

#### **Description**

This function provides lists of genes that have different Major isoforms (most expressed) when different intervals of the experimental conditions are considered.

The subrange of the experimental conditions can be chosen as a specific point, all the points of a specific experimental group or at any point.

30 PodiumChange

#### Usage

```
PodiumChange(get, only.sig.iso=FALSE, comparison=c("any",
"groups", "specific"), group.name="Ctr", time.points=0)
```

#### **Arguments**

get a getDS object a cluster of flat Isoform

only.sig.iso TRUE when changes are looked for only through statistically significant iso-

forms.

comparison Type of search to do: any, groups or specific (see details).

group.name required when comparison is "specific".

time.points required when comparison is "specific".

#### **Details**

There are 3 possible values for the comparison argument:

"any": Detects genes with Major Isoform changes in at least one experimental condition.

"groups": Detects genes with different Major Isoform for different experimental groups.

"specific": Detects genes with Major Isoform changes in a specific time interval, especified in time.points argument and a specific experimental group, especified in group.name argument.

#### Value

Names of the genes with PodiumChange Isoforms

data.L Data values of all the isoforms belonging to the genes in L gen.L gen vector with the name of the gene of each isoform

edesign matrix describing experimental design needed to visualize PodiumChange se-

lection with IsoPlot function. It is the input of make.design.matrix.

#### Author(s)

Maria Jose Nueda, <mj.nueda@ua.es>

## References

Nueda, M.J., Martorell, J., Marti, C., Tarazona, S., Conesa, A. 2018. Identification and visualization of differential isoform expression in RNA-seq time series. Bioinformatics. 34, 3, 524-526. Nueda, M.J., Tarazona, S., Conesa, A. 2014. Next maSigPro: updating maSigPro bioconductor package for RNA-seq time series. Bioinformatics, 30, 2598-602.

Conesa, A., Nueda M.J., Alberto Ferrer, A., Talon, T. 2006. maSigPro: a Method to Identify Significant Differential Expression Profiles in Time-Course Microarray Experiments. Bioinformatics 22, 1096-1102.

#### See Also

```
see.genes, IsoModel
```

position 31

## **Examples**

```
data(ISOdata)
data(ISOdesign)
mdis <- make.design.matrix(ISOdesign)
MyIso <- IsoModel(data=ISOdata[,-1], gen=ISOdata[,1], design=mdis,
counts=TRUE)
Myget <- getDS(MyIso)

PC <- PodiumChange(Myget, only.sig.iso=TRUE,
comparison="specific", group.name="Group2", time.points=c(18,24))
PC$L</pre>
```

position

Column position of a variable in a data frame

## Description

Finds the column position of a character variable in the column names of a data frame.

## Usage

```
position(matrix, vari)
```

## **Arguments**

matrix matrix or data.frame with character column names
vari character variable

## Value

numerical. Column position for the given variable.

## Author(s)

Ana Conesa and Maria Jose Nueda, <mj.nueda@ua.es>

```
x <- matrix(c(1, 1, 2, 2, 3, 3),ncol = 3,nrow = 2)
colnames(x) <- c("one", "two", "three")
position(x, "one")</pre>
```

reg.coeffs

reg.coeffs Calculate true variables regression coefficients
---

## **Description**

reg.coeffs calculates back regression coefficients for true variables (experimental groups) from dummy variables regression coefficients.

## Usage

```
reg.coeffs(coefficients,
indepen = groups.vector[nchar(groups.vector)==min(nchar(groups.vector))][1],
groups.vector, group)
```

## **Arguments**

coefficients vector of regression coefficients obtained from a regression model with dummy

variables

indepen idependent variable of the regression formula

groups.vector vector indicating the true variable of each variable in coefficients group true variable for which regression coefficients are to be computed

## **Details**

regression coefficients in coefficients vector should be ordered by polynomial degree in a regression formula, ie: intercept,  $x^2$  term,  $x^3$  term, and so on...

#### Value

```
reg.coeff vector of calculated regression coefficients
```

## Author(s)

Ana Conesa and Maria Jose Nueda, <mj.nueda@ua.es>

#### References

Conesa, A., Nueda M.J., Alberto Ferrer, A., Talon, T. 2005. maSigPro: a Method to Identify Significant Differential Expression Profiles in Time-Course Microarray Experiments.

```
groups.vector <-c("CT", "T1vsCT", "T2vsCT", "CT", "T1vsCT", "T2vsCT", "CT", "T1vsCT", "T2vsCT")
coefficients <- c(0.1, 1.2, -0.8, 1.7, 3.3, 0.4, 0.0, 2.1, -0.9)
## calculate true regression coefficients for variable "T1"
reg.coeffs(coefficients, groups.vector = groups.vector, group = "T1")</pre>
```

see.genes 33

see.genes	Wrapper function for visualization of gene expression values of time
	course experiments

## **Description**

This function provides visualisation tools for gene expression values in a time course experiment. The function first calls the heatmap function for a general overview of experiment results. Next a partioning of the data is generated using a clustering method. The results of the clustering are visualized both as gene expression profiles extended along all arrays in the experiment, as provided by the plot.profiles function, and as summary expression profiles for comparison among experimental groups.

## Usage

```
see.genes(data, edesign = data$edesign, time.col = 1, repl.col = 2,
  group.cols = c(3:ncol(edesign)), names.groups = colnames(edesign)[3:ncol(edesign)],
  cluster.data = 1, groups.vector = data$groups.vector, k = 9, k.mclust=FALSE,
  cluster.method = "hclust", distance = "cor", agglo.method = "ward.D",
  show.lines = TRUE, show.fit = FALSE, dis = NULL, step.method = "backward",
  min.obs = 3, alfa = 0.05, nvar.correction = FALSE, iter.max = 500,
  summary.mode = "median", color.mode = "rainbow", ylim = NULL, item = "genes",
  legend = TRUE, cex.legend = 1, lty.legend = NULL,...)
```

## Arguments

data	either matrix or a list containing the gene expression data, typically a get.siggenes object
edesign	matrix of experimental design
time.col	column in edesign containing time values. Default is first column
repl.col	column in edesign containing coding for replicates arrays. Default is second column
group.cols	columns indicating the coding for each group (treatment, tissue,) in the experiment (see details)
names.groups	names for experimental groups
cluster.data	type of data used by the cluster algorithm (see details)
groups.vector	vector indicating the experimental group to which each variable belongs
k	number of clusters for data partioning
k.mclust	TRUE for computing the optimal number of clusters with Mclust algorithm
cluster.method	clustering method for data partioning. Currently "hclust", "kmeans" and "Mclust" are supported
distance	distance measurement function when cluster.method is hclust
agglo.method	aggregation method used when cluster.method is hclust

34 see.genes

show.lines logical indicating whether a line must be drawn joining plotted data points for

reach group

show. fit logical indicating whether regression fit curves must be plotted

dis regression design matrix

step.method stepwise regression method to fit models for cluster mean profiles. Can be either

"backward", "forward", "two.ways.backward" or "two.ways.forward"

min.obs minimal number of observations for a gene to be included in the analysis alfa significance level used for variable selection in the stepwise regression

nvar.correction

argument for correcting T. fitsignificance level. See T. fit

iter.max maximum number of iterations when cluster.method is kmeans

summary.mode the method PlotGroups takes to condensate expression information when more

than one gene is present in the data. Possible values are "representative" and

"median"

color.mode color scale for plotting profiles. Can be either "rainblow" or "gray" ylim range of the y axis to be used by PlotProfiles and PlotGroups

item Name of the analysed items to show

legend logical indicating whether legend must be added when plotting profiles

cex.legend Expansion factor for legend

1ty.legend To add a coloured line in the legend ... other graphical function argument

## **Details**

Data can be provided either as a single data matrix of expression values, or a get.siggenes object. In the later case the other argument of the fuction can be taken directly from data.

Data clustering can be done on the basis of either the original expression values, the regression coefficients, or the t.scores. In case data is a get.siggenes object, this is given by providing the element names of the list c("sig.profiles", "coefficients", "t.score") of their list position (1,2 or 3).

#### Value

Experiment wide gene profiles and by group profiles plots are generated for each data cluster in the graphical device.

cut vector indicating gene partioning into clusters
c.algo.used clustering algorith used for data partioning
groups groups matrix used for plotting functions

## Author(s)

Ana Conesa and Maria Jose Nueda, <mj.nueda@ua.es>

see.genes 35

#### References

Conesa, A., Nueda M.J., Alberto Ferrer, A., Talon, T. 2006. maSigPro: a Method to Identify Significant Differential Expression Profiles in Time-Course Microarray Experiments. Bioinformatics 22, 1096-1102

#### See Also

```
PlotProfiles, PlotGroups
```

```
#### GENERATE TIME COURSE DATA
## generate n random gene expression profiles of a data set with
## one control plus 3 treatments, 3 time points and r replicates per time point.
tc.GENE <- function(n, r,</pre>
             var11 = 0.01, var12 = 0.01, var13 = 0.01,
             var21 = 0.01, var22 = 0.01, var23 = 0.01,
             var31 = 0.01, var32 = 0.01, var33 = 0.01,
             var41 = 0.01, var42 = 0.01, var43 = 0.01,
             a1 = 0, a2 = 0, a3 = 0, a4 = 0,
             b1 = 0, b2 = 0, b3 = 0, b4 = 0,
             c1 = 0, c2 = 0, c3 = 0, c4 = 0)
{
 tc.dat <- NULL
 for (i in 1:n) {
   Ctl <- c(rnorm(r, a1, var11), rnorm(r, b1, var12), rnorm(r, c1, var13)) # Ctl group
   Tr1 <- c(rnorm(r, a2, var21), rnorm(r, b2, var22), rnorm(r, c2, var23)) # Tr1 group
   Tr2 <- c(rnorm(r, a3, var31), rnorm(r, b3, var32), rnorm(r, c3, var33)) # Tr2 group
   Tr3 <- c(rnorm(r, a4, var41), rnorm(r, b4, var42), rnorm(r, c4, var43)) # Tr3 group
   gene <- c(Ctl, Tr1, Tr2, Tr3)
    tc.dat <- rbind(tc.dat, gene)</pre>
 tc.dat
}
## Create 270 flat profiles
flat <- tc.GENE(n = 270, r = 3)
## Create 10 genes with profile differences between Ctl and Tr1 groups
twodiff \leftarrow tc.GENE (n = 10, r = 3, b2 = 0.5, c2 = 1.3)
## Create 10 genes with profile differences between Ctl, Tr2, and Tr3 groups
threediff < tc.GENE(n = 10, r = 3, b3 = 0.8, c3 = -1, a4 = -0.1, b4 = -0.8, c4 = -1.2)
## Create 10 genes with profile differences between Ctl and Tr2 and different variance
vardiff < -tc.GENE(n = 10, r = 3, a3 = 0.7, b3 = 1, c3 = 1.2, var32 = 0.03, var33 = 0.03)
## Create dataset
tc.DATA <- rbind(flat, twodiff, threediff, vardiff)</pre>
rownames(tc.DATA) <- paste("feature", c(1:300), sep = "")
colnames(tc.DATA) <- paste("Array", c(1:36), sep = "")</pre>
tc.DATA [sample(c(1:(300*36)), 300)] <- NA # introduce missing values
#### CREATE EXPERIMENTAL DESIGN
```

36 seeDS

seeDS

Wrapper function for visualization of significant isoforms from Differentially Spliced Genes

## **Description**

seeDS This function provides visualisation tools for Significant Isoforms in a time course experiment. The function calls the see.genes function for selected Isoforms. This cluster will be the reference in tableDS function to identify the trends that follows the isoforms of a specific gene.

## Usage

```
seeDS(get, rsq=0.4, cluster.all=TRUE, plot.mDSG=FALSE, k=6,
cluster.method="hclust", k.mclust=FALSE, ...)
```

## **Arguments**

get	a getDS object a cluster of flat Isoform
rsq	Required when cluster.all=TRUE. It is the cut-off level at the R-squared value for detecting significant isoforms of all the genome.
cluster.all	TRUE to make the cluster with significant isoforms of all the genome. FALSE to make the cluster with significant isoforms of Differentially Spliced Genes.
plot.mDSG	TRUE to make a cluster with the Isoforms belonging to monoIsoform genes
k	number of clusters for data partioning
cluster.method	clustering method for data partioning. Currently "hclust", "kmeans" and "Mclust" are supported
k.mclust	TRUE for computing the optimal number of clusters with Mclust algorithm
	other graphical function argument

seeDS 37

#### **Details**

The cluster reference can be made with significant isoforms of all the genome or with the isoforms belonging to the Differentially Spliced Genes.

Alternatively a cluster of monoIsoforms can be asked.

Next a partioning of the data is generated using a clustering method.

The results of the clustering are visualized in two plots as in see.genes.

## Value

Experiment wide Isoform profiles and by group profiles plots are generated for each data cluster in the graphical device.

Model a IsoModel object to be used in the following steps get a get.siggenes object to be used in the following steps

NumIso.by.gene Number of selected Isoforms for each Differentially Spliced Gene

cut vector indicating gene partioning into clusters

names genes vector with the name of the gene each selected isoform belongs to

#### Author(s)

Maria Jose Nueda, <mj.nueda@ua.es>

#### References

Nueda, M.J., Martorell, J., Marti, C., Tarazona, S., Conesa, A. 2018. Identification and visualization of differential isoform expression in RNA-seq time series. Bioinformatics. 34, 3, 524-526.

Nueda, M.J., Tarazona, S., Conesa, A. 2014. Next maSigPro: updating maSigPro bioconductor package for RNA-seq time series. Bioinformatics, 30, 2598-602.

Conesa, A., Nueda M.J., Alberto Ferrer, A., Talon, T. 2006. maSigPro: a Method to Identify Significant Differential Expression Profiles in Time-Course Microarray Experiments. Bioinformatics 22, 1096-1102.

#### See Also

```
see.genes, IsoModel
```

```
data(ISOdata)
data(ISOdesign)
mdis <- make.design.matrix(ISOdesign)
MyIso <- IsoModel(data=ISOdata[,-1], gen=ISOdata[,1], design=mdis, counts=TRUE)
Myget <- getDS(MyIso)
see <- seeDS(Myget, cluster.all=FALSE, k=6)

table <- tableDS(see)
table$IsoTable</pre>
```

38 stepback

stepback	Fitting a linear model by backward-stepwise regression

# **Description**

stepback fits a linear regression model applying a backward-stepwise strategy.

## Usage

```
stepback(y = y, d = d, alfa = 0.05, family = gaussian(), epsilon=0.00001)
```

# Arguments

У	dependent variable
d	data frame containing by columns the set of variables that could be in the selected model
alfa	significance level to decide if a variable stays or not in the model
family	the distribution function to be used in the glm model
epsilon	argument to pass to glm. control, convergence tolerance in the iterative process to estimate de glm model

#### **Details**

The strategy begins analysing a model with all the variables included in d. If all variables are statistically significant (all variables have a p-value less than alfa) this model will be the result. If not, the less statistically significant variable will be removed and the model is re-calculated. The process is repeated up to find a model with all the variables statistically significant.

## Value

stepback returns an object of the class 1m, where the model uses y as dependent variable and all the selected variables from d as independent variables.

The function summary are used to obtain a summary and analysis of variance table of the results. The generic accessor functions coefficients, effects, fitted.values and residuals extract various useful features of the value returned by lm.

## Author(s)

Ana Conesa, aconesa@cipf.es; Maria Jose Nueda, mj.nueda@ua.es

## References

Conesa, A., Nueda M.J., Alberto Ferrer, A., Talon, T. 2005. maSigPro: a Method to Identify Significant Differential Expression Profiles in Time-Course Microarray Experiments.

stepfor 39

## See Also

```
lm, step, stepfor, two.ways.stepback, two.ways.stepfor
```

# **Examples**

```
## create design matrix
Time \leftarrow rep(c(rep(c(1:3), each = 3)), 4)
Replicates <- rep(c(1:12), each = 3)
Control <- c(rep(1, 9), rep(0, 27))
Treat1 <- c(rep(0, 9), rep(1, 9), rep(0, 18))
Treat2 <- c(rep(0, 18), rep(1, 9), rep(0,9))
Treat3 <- c(rep(0, 27), rep(1, 9))
edesign <- cbind(Time, Replicates, Control, Treat1, Treat2, Treat3)</pre>
rownames(edesign) <- paste("Array", c(1:36), sep = "")</pre>
dise <- make.design.matrix(edesign)</pre>
dis <- as.data.frame(dise$dis)</pre>
## expression vector
y < -c(0.082, 0.021, 0.010, 0.113, 0.013, 0.077, 0.068, 0.042, -0.056, -0.232, -0.014, -0.040,
-0.055, 0.150, -0.027, 0.064, -0.108, -0.220, 0.275, -0.130, 0.130, 1.018, 1.005, 0.931,
-1.009, -1.101, -1.014, -0.045, -0.110, -0.128, -0.643, -0.785, -1.077, -1.187, -1.249, -1.463)
s.fit \leftarrow stepback(y = y, d = dis)
summary(s.fit)
```

stepfor

Fitting a linear model by forward-stepwise regression

# **Description**

stepfor fits a linear regression model applying forward-stepwise strategy.

# Usage

```
stepfor(y = y, d = d, alfa = 0.05, family = gaussian(), epsilon=0.00001 )
```

# **Arguments**

У	dependent variable
d	data frame containing by columns the set of variables that could be in the selected model
alfa	significance level to decide if a variable stays or not in the model
family	the distribution function to be used in the glm model
epsilon	argument to pass to glm.control, convergence tolerance in the iterative process to estimate de glm model

40 stepfor

#### **Details**

The strategy begins analysing all the possible models with only one of the variables included in d. The most statistically significant variable (with the lowest p-value) is included in the model and then it is considered to introduce in the model another variable analysing all the possible models with two variables (the selected variable in the previous step plus a new variable). Again the most statistically significant variable (with lowest p-value) is included in the model. The process is repeated till there are no more statistically significant variables to include.

#### Value

stepfor returns an object of the class 1m, where the model uses y as dependent variable and all the selected variables from d as independent variables.

The function summary are used to obtain a summary and analysis of variance table of the results. The generic accessor functions coefficients, effects, fitted.values and residuals extract various useful features of the value returned by lm.

#### Author(s)

Ana Conesa, aconesa@cipf.es; Maria Jose Nueda, mj.nueda@ua.es

#### References

Conesa, A., Nueda M.J., Alberto Ferrer, A., Talon, T. 2005. maSigPro: a Method to Identify Significant Differential Expression Profiles in Time-Course Microarray Experiments.

#### See Also

```
lm, step, stepback, two.ways.stepback, two.ways.stepfor
```

```
## create design matrix
Time \leftarrow rep(c(rep(c(1:3), each = 3)), 4)
Replicates \leftarrow rep(c(1:12), each = 3)
Control <- c(rep(1, 9), rep(0, 27))
Treat1 <- c(rep(0, 9), rep(1, 9), rep(0, 18))
Treat2 <- c(rep(0, 18), rep(1, 9), rep(0,9))
Treat3 <- c(rep(0, 27), rep(1, 9))
edesign <- cbind(Time, Replicates, Control, Treat1, Treat2, Treat3)</pre>
rownames(edesign) <- paste("Array", c(1:36), sep = "")</pre>
dise <- make.design.matrix(edesign)</pre>
dis <- as.data.frame(dise$dis)</pre>
## expression vector
y <- c(0.082, 0.021, 0.010, 0.113, 0.013, 0.077, 0.068, 0.042, -0.056, -0.232, -0.014, -0.040,
-0.055, 0.150, -0.027, 0.064, -0.108, -0.220, 0.275, -0.130, 0.130, 1.018, 1.005, 0.931,
-1.009, -1.101, -1.014, -0.045, -0.110, -0.128, -0.643, -0.785, -1.077, -1.187, -1.249, -1.463)
s.fit <- stepfor(y = y, d = dis)
```

suma2Venn 41

```
summary(s.fit)
```

suma2Venn

Creates a Venn Diagram from a matrix of characters

# Description

suma2Venn transforms a matrix or a data frame with characters into a list to draw and display a Venn diagram with up to 7 sets

# Usage

```
suma2Venn(x, size = 30, cexil = 0.9, cexsn = 1, zcolor = heat.colors(ncol(x)), ...)
```

# **Arguments**

X	matrix or data frame of character values
size	Plot size, in centimeters
cexil	Character expansion for the intersection labels
cexsn	Character expansion for the set names
zcolor	A vector of colors for the custom zones
	Additional plotting arguments for the venn function

# **Details**

suma2Venn creates a list with the columns of a matrix or a data frame of characters which can be taken by the venn to generate a Venn Diagram

#### Value

suma2Venn returns a Venn Plot such as that created by the venn function

# Author(s)

Ana Conesa and Maria Jose Nueda, <mj.nueda@ua.es>

#### See Also

venn

```
A <- c("a","b","c", "d", "e", NA, NA)
B <- c("a","b","f", NA, NA, NA, NA)
C <- c("a","b","e","f", "h", "i", "j", "k")
x <- cbind(A, B, C)
suma2Venn(x)
```

42 T.fit

T.fit Makes a stepwise regression fit for time series gene expression iments	ion exper-
--	------------

# **Description**

T. fit selects the best regression model for each gene using stepwise regression.

# Usage

```
T.fit(data, design = data$dis, step.method = "backward",
    min.obs = data$min.obs, alfa = data$Q,
    nvar.correction = FALSE, family = gaussian(),
    epsilon=0.00001, item="gene")
```

## **Arguments**

data	can either be a p. vector object or a matrix containing expression data with the same requirements as for the p. vector function	
design	design matrix for the regression fit such as that generated by the make.design.matrix function. If data is a p.vector object, the same design matrix is used by default	
step.method	argument to be passed to the step function. Can be either "backward", "forward", "two.ways.backward" or "two.ways.forward"	
min.obs	genes with less than this number of true numerical values will be excluded from the analysis	
alfa	significance level used for variable selection in the stepwise regression	
nvar.correction		
	argument for correcting T.fit significance level. See details	
family	the distribution function to be used in the glm model. It must be the same used in p.vector	
epsilon	argument to pass to glm. control, convergence tolerance in the iterative process to estimate de glm model	
item	Name of the analysed item to show in the screen while T.fit is in process	

# **Details**

In the maSigPro approach p.vector and T.fit are subsequent steps, meaning that significant genes are first selected on the basis of a general model and then the significant variables for each gene are found by step-wise regression.

The step regression can be "backward" or "forward" indicating whether the step procedure starts from the model with all or none variables. With the "two.ways.backward" or "two.ways.forward" options the variables are both allowed to get in and out. At each step the p-value of each variable is computed and variables get in/out the model when this p-value is lower or higher than given threshold alfa. When nva.correction is TRUE the given significance level is corrected by the number of variables in the model

T.fit

#### Value

sol matrix for summary results of the stepwise regression. For each selected gene the following values are given: • p-value of the regression ANOVA • R-squared of the model • p-value of the regression coefficients of the selected variables sig.profiles expression values for the genes contained in sol coefficients matrix containing regression coefficients for the adjusted models groups.coeffs matrix containing the coefficients of the impiclit models of each experimental group variables variables in the complete regression model G total number of input genes number of genes taken in the regression fit g dat input analysis data matrix dis regression design matrix step.method imputed step method for stepwise regression edesign matrix of experimental design

#### Author(s)

influ.info

Ana Conesa and Maria Jose Nueda, <mj.nueda@ua.es>

# References

Conesa, A., Nueda M.J., Alberto Ferrer, A., Talon, T. 2006. maSigPro: a Method to Identify Significant Differential Expression Profiles in Time-Course Microarray Experiments. Bioinformatics 22, 1096-1102

data frame of genes containing influencial data

#### See Also

```
p.vector, step
```

44 tableDS

```
c1 = 0, c2 = 0, c3 = 0, c4 = 0)
{
 tc.dat <- NULL
 for (i in 1:n) {
   Ctl <- c(rnorm(r, a1, var11), rnorm(r, b1, var12), rnorm(r, c1, var13)) # Ctl group
   Tr1 <- c(rnorm(r, a2, var21), rnorm(r, b2, var22), rnorm(r, c2, var23)) # Tr1 group
   Tr2 <- c(rnorm(r, a3, var31), rnorm(r, b3, var32), rnorm(r, c3, var33)) # Tr2 group
   Tr3 <- c(rnorm(r, a4, var41), rnorm(r, b4, var42), rnorm(r, c4, var43)) # Tr3 group
    gene <- c(Ctl, Tr1, Tr2, Tr3)
    tc.dat <- rbind(tc.dat, gene)</pre>
 }
 tc.dat
}
## Create 270 flat profiles
flat <- tc.GENE(n = 270, r = 3)
## Create 10 genes with profile differences between Ctl and Tr1 groups
twodiff \leftarrow tc.GENE (n = 10, r = 3, b2 = 0.5, c2 = 1.3)
## Create 10 genes with profile differences between Ctl, Tr2, and Tr3 groups
threediff < tc.GENE(n = 10, r = 3, b3 = 0.8, c3 = -1, a4 = -0.1, b4 = -0.8, c4 = -1.2)
## Create 10 genes with profile differences between Ctl and Tr2 and different variance
vardiff < -tc.GENE(n = 10, r = 3, a3 = 0.7, b3 = 1, c3 = 1.2, var32 = 0.03, var33 = 0.03)
## Create dataset
tc.DATA <- rbind(flat, twodiff, threediff, vardiff)</pre>
rownames(tc.DATA) <- paste("feature", c(1:300), sep = "")</pre>
colnames(tc.DATA) <- paste("Array", c(1:36), sep = "")</pre>
tc.DATA [sample(c(1:(300*36)), 300)] <- NA # introduce missing values
#### CREATE EXPERIMENTAL DESIGN
Time \leftarrow rep(c(rep(c(1:3), each = 3)), 4)
Replicates \leftarrow rep(c(1:12), each = 3)
Control <- c(rep(1, 9), rep(0, 27))
Treat1 <- c(rep(0, 9), rep(1, 9), rep(0, 18))
Treat2 <- c(rep(0, 18), rep(1, 9), rep(0,9))
Treat3 <- c(rep(0, 27), rep(1, 9))
edesign <- cbind(Time, Replicates, Control, Treat1, Treat2, Treat3)
rownames(edesign) <- paste("Array", c(1:36), sep = "")</pre>
## run T.fit from a p.vector object
tc.p <- p.vector(tc.DATA, design = make.design.matrix(edesign), Q = 0.01)</pre>
tc.tstep <- T.fit(data = tc.p , alfa = 0.05)</pre>
## run T.fit from a data matrix and a design matrix
dise <- make.design.matrix(edesign)</pre>
tc.tstep <- T.fit (data = tc.DATA[271:300,], design = dise$dis,</pre>
                   step.method = "two.ways.backward", min.obs = 10, alfa = 0.05)
tc.tstep$sol # gives the p.values of the significant
             # regression coefficients of the optimized models
```

tableDS 45

# **Description**

tableDS identifies for each Differentialy Spliced Gene (DSG) the clusters where their isoforms belong to, labelling gene transcripts as mayor (or most expressed) and minor.

# Usage

tableDS(seeDS)

# **Arguments**

seeDS

a seeDS object

#### **Details**

This table includes DSG with 2 or more Isoforms. Mono isoform genes are useful to determine the trends of the cluster. However, as they have only one Isoform, there is not the possibility of comparing minor and major DETs.

#### Value

IsoTable A classification table that indicates the distribution of isoforms across different

clusters

IsoClusters A data frame with genes in rows and two columns: first indicates the number of

cluster of the major isoform and second the number(s) of cluster(s) of the minor

isoforms.

#### Author(s)

Maria Jose Nueda, <mj.nueda@ua.es>

#### References

Nueda, M.J., Martorell, J., Marti, C., Tarazona, S., Conesa, A. 2018. Identification and visualization of differential isoform expression in RNA-seq time series. Bioinformatics. 34, 3, 524-526.

Nueda, M.J., Tarazona, S., Conesa, A. 2014. Next maSigPro: updating maSigPro bioconductor package for RNA-seq time series. Bioinformatics, 30, 2598-602.

Conesa, A., Nueda M.J., Alberto Ferrer, A., Talon, T. 2006. maSigPro: a Method to Identify Significant Differential Expression Profiles in Time-Course Microarray Experiments. Bioinformatics 22, 1096-1102.

#### See Also

seeDS, IsoModel

46 two.ways.stepback

## **Examples**

```
data(ISOdata)
data(ISOdesign)
mdis <- make.design.matrix(ISOdesign)

MyIso <- IsoModel(data=ISOdata[,-1], gen=ISOdata[,1], design=mdis, counts=TRUE)
Myget <- getDS(MyIso)
see <- seeDS(Myget, cluster.all=FALSE, k=6)
table <- tableDS(see)
table$IsoTable</pre>
```

two.ways.stepback

Fitting a linear model by backward-stepwise regression

# **Description**

two.ways.stepback fits a linear regression model applying backward-stepwise strategy.

#### Usage

```
two.ways.stepback(y = y, d = d, alfa = 0.05, family = gaussian(), epsilon=0.00001)
```

# Arguments

У	dependent variable
d	data frame containing by columns the set of variables that could be in the selected model
alfa	significance level to decide if a variable stays or not in the model
family	the distribution function to be used in the glm model
epsilon	argument to pass to glm.control, convergence tolerance in the iterative process to estimate de glm model

#### **Details**

The strategy begins analysing a model with all the variables included in d. If all the variables are statistically significant (all the variables have a p-value less than alfa) this model will be the result. If not, the less statistically significant variable will be removed and the model is re-calculated. The process is repeated up to find a model with all the variables statistically significant (p-value < alpha). Each time that a variable is removed from the model, it is considered the possibility of one or more removed variables to come in again.

#### Value

 $two.ways.stepback\ returns\ an\ object\ of\ the\ class\ 1m,\ where\ the\ model\ uses\ y\ as\ dependent\ variable$  and all the selected variables from d as independent variables.

The function summary are used to obtain a summary and analysis of variance table of the results. The generic accessor functions coefficients, effects, fitted.values and residuals extract various useful features of the value returned by lm.

two.ways.stepfor 47

#### Author(s)

Ana Conesa and Maria Jose Nueda, <mj.nueda@ua.es>

#### References

Conesa, A., Nueda M.J., Alberto Ferrer, A., Talon, T. 2005. maSigPro: a Method to Identify Significant Differential Expression Profiles in Time-Course Microarray Experiments.

#### See Also

```
lm, step, stepfor, stepback, two.ways.stepfor
```

# **Examples**

```
## create design matrix
Time \leftarrow rep(c(rep(c(1:3), each = 3)), 4)
Replicates \leftarrow rep(c(1:12), each = 3)
Control <- c(rep(1, 9), rep(0, 27))
Treat1 <- c(rep(0, 9), rep(1, 9), rep(0, 18))
Treat2 <- c(rep(0, 18), rep(1, 9), rep(0,9))
Treat3 <- c(rep(0, 27), rep(1, 9))
edesign <- cbind(Time, Replicates, Control, Treat1, Treat2, Treat3)</pre>
rownames(edesign) <- paste("Array", c(1:36), sep = "")</pre>
dise <- make.design.matrix(edesign)</pre>
dis <- as.data.frame(dise$dis)</pre>
## expression vector
y < -c(0.082, 0.021, 0.010, 0.113, 0.013, 0.077, 0.068, 0.042, -0.056, -0.232, -0.014, -0.040,
-0.055, 0.150, -0.027, 0.064, -0.108, -0.220, 0.275, -0.130, 0.130, 1.018, 1.005, 0.931,
-1.009, -1.101, -1.014, -0.045, -0.110, -0.128, -0.643, -0.785, -1.077, -1.187, -1.249, -1.463)
s.fit <- two.ways.stepback(y = y, d = dis)</pre>
summary(s.fit)
```

two.ways.stepfor

Fitting a linear model by forward-stepwise regression

# **Description**

two.ways.stepfor fits a linear regression model applying forward-stepwise strategy.

#### Usage

```
two.ways.stepfor(y = y, d = d, alfa = 0.05, family = gaussian(), epsilon=0.00001)
```

48 two.ways.stepfor

## Arguments

У	dependent variable
d	data frame containing by columns the set of variables that could be in the selected model
alfa	significance level to decide if a variable stays or not in the model
family	the distribution function to be used in the glm model
epsilon	argument to pass to glm.control, convergence tolerance in the iterative process to estimate de glm model

#### Details

The strategy begins analysing all the possible models with only one of the variables included in d. The most statistically significant variable (with the lowest p-value) is included in the model and then it is considered to introduce in the model another variable analysing all the possible models with two variables (the selected variable in the previous step plus a new variable). Again the most statistically significant variable (with lowest p-value) is included in the model. The process is repeated till there are no more statistically significant variables to include. Each time that a variable enters the model, the p-values of the current model vairables is recalculated and non significant variables will be removed.

## Value

two.ways.stepfor returns an object of the class lm, where the model uses y as dependent variable and all the selected variables from d as independent variables.

The function summary are used to obtain a summary and analysis of variance table of the results. The generic accessor functions coefficients, effects, fitted.values and residuals extract various useful features of the value returned by lm.

# Author(s)

Ana Conesa and Maria Jose Nueda, <mj.nueda@ua.es>

# References

Conesa, A., Nueda M.J., Alberto Ferrer, A., Talon, T. 2005. maSigPro: a Method to Identify Significant Differential Expression Profiles in Time-Course Microarray Experiments.

#### See Also

```
lm, step, stepback, stepfor, two.ways.stepback
```

```
## create design matrix
Time <- rep(c(rep(c(1:3), each = 3)), 4)
Replicates <- rep(c(1:12), each = 3)
Control <- c(rep(1, 9), rep(0, 27))
Treat1 <- c(rep(0, 9), rep(1, 9), rep(0, 18))</pre>
```

two.ways.stepfor 49

```
Treat2 <- c(rep(0, 18), rep(1, 9), rep(0,9))
Treat3 <- c(rep(0, 27), rep(1, 9))
edesign <- cbind(Time, Replicates, Control, Treat1, Treat2, Treat3)
rownames(edesign) <- paste("Array", c(1:36), sep = "")
dise <- make.design.matrix(edesign)
dis <- as.data.frame(dise$dis)

## expression vector
y <- c(0.082, 0.021, 0.010, 0.113, 0.013, 0.077, 0.068, 0.042, -0.056, -0.232, -0.014, -0.040, -0.055, 0.150, -0.027, 0.064, -0.108, -0.220, 0.275, -0.130, 0.130, 1.018, 1.005, 0.931, -1.009, -1.101, -1.014, -0.045, -0.110, -0.128, -0.643, -0.785, -1.077, -1.187, -1.249, -1.463)
s.fit <- two.ways.stepfor(y = y, d = dis)
summary(s.fit)</pre>
```

# **Index**

* UsersGuide	coefficients, 38, 40, 46, 48
maSigProUsersGuide, 20	
* aplot	data.abiotic,3
PlotGroups, 25	
PlotProfiles, 28	edesign.abiotic,5
see.genes, 33	edesignCT, 6
suma2Venn, 41	edesignDR, 6
* arith	effects, 38, 40, 46, 48
average.rows, 2	fitted.values, 38, 40, 46, 48
i.rank, 14	11tted. values, 30, 40, 40, 40
position, 31	get.siggenes, 7, 11, 12, 33, 34
* datasets	getDS, 11, 19
data.abiotic, 3	getDSPatterns, 12
edesign.abiotic,5	getser determs, 12
edesignCT, 6	hclust, 33
edesignDR, 6	,
ISOdata, 14	i.rank, 14
ISOdesign, 15	ISOdata, 14
NBdata, 21	ISOdesign, 15
NBdesign, 22	IsoModel, 12, 13, 16, 19, 30, 37, 45
* design	IsoPlot, 18
make.design.matrix, 19	
* manip	kmeans, 34
get.siggenes,7	1 24 29 40 46 49
see.genes, 33	lm, 24, 38–40, 46–48
* misc	make.design.matrix, <i>16</i> , 19, <i>23</i> , <i>42</i>
reg.coeffs, 32	maSigProUsersGuide, 20
* models	masign rooser source, 20
T. fit, 42	NBdata, 21
* regression	NBdesign, 22
make.design.matrix, 19	3 7
p.vector, 22	order, <i>14</i>
stepback, 38	
stepfor, 39	p.vector, <i>17</i> , 22, <i>42</i> , <i>43</i>
T.fit, 42	PlotGroups, 25, 29, 34, 35
two.ways.stepback, 46	PlotProfiles, 27, 28, 34, 35
two.ways.stepback,40 two.ways.stepfor,47	PodiumChange, 29
two.ways.Stepior, 4/	position, 31
average.rows, 2, 8, 9	rank, <i>14</i>
a. c. a <sub>0</sub> c 3no, 2, 0, >	,

INDEX 51

```
reg.coeffs, 32
residuals, 38, 40, 46, 48
see.genes, 30, 33, 37
seeDS, 36, 45
step, 39, 40, 43, 47, 48
stepback, 38, 40, 47, 48
stepfor, 39, 39, 47, 48
suma2Venn, 41
summary, 38, 40, 46, 48
T.fit, 17, 24, 26, 34, 42, 42
tableDS, 13, 44
two.ways.stepback, 39, 40, 46, 48
two.ways.stepfor, 39, 40, 47, 47
venn, 41
```