

maSigPro

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| | |
|--------------|--|
| average.rows | <i>Average rows by match and index</i> |
|--------------|--|

Description

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.

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 |

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)

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Examples

```
## create data matrix for row averaging
x <- matrix(rnorm(30), nrow = 6, ncol = 5)
rownames(x) <- paste("ID", c(1, 2, 11, 12, 19, 20), sep = "")
i <- paste("g", rep(c(1:10), each = 2), sep = "") # index vector
m <- 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_3H_3 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_2 a numeric vector
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 which gene expression was monitored in both roots and leaves using a 11K cDNA potato chip. This example data set contains a random 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 *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 hybridizations, 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.

Examples

```
data(edesignCR)
```

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 columns

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 hybridizations, 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 hybridizations, 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)
```

| | |
|------------|---|
| edesign.OD | <i>Experimental design with a measured independent variable</i> |
|------------|---|

Description

edesign.OD contains the experimental design of a E.coli growth time course microarray experiment with a temperature shift treatment. The OD of each culture was measured and used in the experimental design as independent variable.

Usage

```
data(edesign.OD)
```

Format

A data frame with 52 rows and the following 4 variables.

OD a numeric vector. Indicates the OD value of the sampled culture

Replicate a numeric vector

37 a numeric vector. No temperature shift treatment

SHIFT a numeric vector. Temperature shift treatment

Examples

```
data(edesign.OD)
## maybe str(edesign.OD) ; plot(edesign.OD) ...
```

| | |
|--------------|---|
| get.siggenes | <i>Extract significant genes for sets of variables in time series gene expression experiments</i> |
|--------------|---|

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

```
get.siggenes(tstep, rsq = 0.7, add.IDs = FALSE, IDs = NULL, matchID.col = 1,
             only.names = FALSE, vars = c("all", "each", "groups"),
             significant.intercept = "dummy",
             groups.vector = NULL, trat.repl.spots = "none",
             index = IDs[, (matchID.col + 1)], match = IDs[, matchID.col],
             r = 0.7)
```

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.IDs is TRUE) |
| matchID.col | number of matching column in matrix IDs for adding genes ids |
| only.names | logical. If TRUE, expression values are omitted in the results |
| vars | variables for which to extract significant genes (see details) |
| significant.intercept | 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 | minimum 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 geneIDs 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 argument values.
`add.IDs = TRUE` and `only.names = TRUE` are compatible argument values.

Value

| | |
|------------------------|---|
| <code>summary</code> | a vector or matrix listing significant genes for the variables given by the function parameters |
| <code>sig.genes</code> | 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 containing: <ul style="list-style-type: none"> sig.profiles: expression values of significant genes coefficients: regression coefficients of the adjusted models groups.coeffs: regression coefficients of the implicit models of each experimental 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, aconesa@ivia.es; 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

```
#### 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,
  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)
}
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 <- 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)
rownames(tc.DATA) <- paste("feature", c(1:300), sep = "")
colnames(tc.DATA) <- paste("Array", c(1:36), sep = "")
tc.DATA [sample(c(1:(300*36)), 300)] <- NA # introduce missing values

#### CREATE EXPERIMENTAL DESIGN
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 = 0.01)
tc.tstep <- T.fit(data = tc.p , alfa = 0.05)

## This will obtain significant 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 significant genes regardless the Rsquared value.
## Replicated genes are averaged.
IDs <- rbind(paste("feature", c(1:300), sep = ""),
            rep(paste("gene", c(1:150), sep = ""), each = 2))
tc.sigs.ALL <- get.siggenes (tc.tstep, rsq = 0, vars = "all", IDs = IDs)
tc.sigs.groups <- get.siggenes (tc.tstep, rsq = 0, vars = "groups", significant.intercept

```

Description

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

Usage

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

Arguments

`x` vector

Value

Vector of ranked values

Author(s)

Ana Conesa, aconesa@ivia.es

See Also

[rank.order](#)

Examples

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

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

```
make.design.matrix(edesign, degree = 2, time.col = 1,
                   repl.col = 2, group.cols = c(3:ncol(edesign)))
```

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 abscissa 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

| | |
|----------------------------|--|
| <code>dis</code> | design matrix of dummies for fitting time series |
| <code>groups.vector</code> | vector coding the experimental group to which each variable belongs to |
| <code>edesign</code> | edesign value passed as argument |

Author(s)

Ana Conesa, aconesa@ivia.es; 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
```

| | |
|----------|--|
| maSigPro | <i>Wrapping function for identifying significant differential gene expression profiles in micorarray time course experiments</i> |
|----------|--|

Description

maSigPro performs a whole maSigPro analysis for a times series gene expression experiment. The function sucesively calls the functions `make.design.matrix(optional)`, `p.vector`, `T.fit`, `get.siggenes` and `see.genes`.

Usage

```
maSigPro(data, edesign, matrix = "AUTO", groups.vector = NULL,
  degree = 2, time.col = 1, repl.col = 2, group.cols = c(3:ncol(edesign)),
  Q = 0.05, alfa = Q, nvar.correction = FALSE, step.method = "backward", rsq =
  min.obs = 3, vars = "groups", significant.intercept = "dummy", cluster.data
  add.IDs = FALSE, IDs = NULL, matchID.col = 1, only.names = FALSE, k = 9, m =
  cluster.method = "hclust", distance = "cor", agglo.method = "ward", iter.max
  summary.mode = "median", color.mode = "rainbow", trat.repl.spots = "none",
  index = IDs[, (matchID.col + 1)], match = IDs[, matchID.col], rs = 0.7,
  show.fit = TRUE, show.lines = TRUE, pdf = TRUE, cexlab = 0.8,
  legend = TRUE, main = NULL, ...)
```

Arguments

| | |
|-----------------|---|
| data | matrix with normalized gene expression data. Genes must be in rows and arrays in columns. Row names must contain geneIDs (argument of p.vector) |
| edesign | matrix of experimental design. Row names must contain arrayIDs (argument of make.design.matrix and see.genes) |
| matrix | design matrix for regression analysis. By default design is calculated with make.design.matrix (argument of p.vector and T.fit , by default computed by make.design.matrix) |
| groups.vector | vector indicating experimental group of each variable (argument of get.siggenes and see.genes , by default computed by make.design.matrix) |
| degree | the degree of the regression fit polynome. degree = 1 returns lineal regression, degree = 2 returns quadratic regression, etc... (argument of make.design.matrix) |
| time.col | column in edesign containing time values. Default is first column (argument of make.design.matrix and see.genes) |
| repl.col | column in edesign containing coding for replicates arrays. Default is second column (argument of make.design.matrix and see.genes) |
| group.cols | columns in edesign indicating the coding for each group of the experiment (see make.design.matrix) (argument of make.design.matrix and see.genes) |
| Q | level of false discovery rate (FDR) control (argument of p.vector) |
| alfa | significance level used for variable selection in the stepwise regression (argument of T.fit) |
| nvar.correction | logical for indicating correcting of stepwise regression significance level (argument of T.fit) |
| step.method | argument to be passed to the step function. Can be either "backward", "forward", "two.ways.backward" or "two.ways.forward" |
| rsq | cut-off level at the R-squared value for the stepwise regression fit. Only genes with R-squared greater than rsq are selected |

| | |
|------------------------------------|--|
| <code>min.obs</code> | genes with less than this number of true numerical values will be excluded from the analysis (argument of <code>p.vector</code> and <code>T.fit</code>) |
| <code>vars</code> | variables for which to extract significant genes (argument of <code>get.siggenes</code>) |
| <code>significant.intercept</code> | experimental groups for which significant intercept coefficients are considered (argument of <code>get.siggenes</code>) |
| <code>cluster.data</code> | Type of data used by the cluster algorithm (argument of <code>see.genes</code>) |
| <code>add.IDs</code> | logical indicating whether to include additional gene id's in the significant genes result (argument of <code>get.siggenes</code>) |
| <code>IDs</code> | matrix containing additional gene id information (required when <code>add.IDs</code> is TRUE) (argument of <code>get.siggenes</code>) |
| <code>matchID.col</code> | number of matching column in matrix IDs for adding genes ids (argument of <code>get.siggenes</code>) |
| <code>only.names</code> | logical. If TRUE, expression values are omitted in the significant genes result (argument of <code>get.siggenes</code>) |
| <code>k</code> | number of clusters (argument of <code>see.genes</code>) |
| <code>m</code> | m parameter when "mfuzz" clustering algorithm is used. See <code>mfuzz</code> (argument of <code>see.genes</code>) |
| <code>cluster.method</code> | clustering method for data partitioning (argument of <code>see.genes</code>) |
| <code>distance</code> | distance measurement function used when <code>cluster.method</code> is "hclust" (argument of <code>see.genes</code>) |
| <code>agglo.method</code> | aggregation method used when <code>cluster.method</code> is "hclust" (argument of <code>see.genes</code>) |
| <code>iter.max</code> | number of iterations when <code>cluster.method</code> is "kmeans" (argument of <code>see.genes</code>) |
| <code>summary.mode</code> | the method to condensate expression information when more than one gene is present in the data. Possible values are "representative" and "median" (argument of <code>PlotGroups</code>) |
| <code>color.mode</code> | color scale for plotting profiles. Can be either "rainblow" or "gray" (argument of <code>PlotProfiles</code>) |
| <code>trat.repl.spots</code> | treatment given to replicate spots. Possible values are "none" and "average" (argument of <code>get.siggenes</code>) |
| <code>index</code> | argument of the <code>average.rows</code> function to use when <code>trat.repl.spots</code> is "average" (argument of <code>get.siggenes</code>) |

| | |
|-------------------------|---|
| <code>match</code> | argument of the <code>link{\average.rows}</code> function to use when <code>trat.repl.spots</code> is "average" (argument of <code>get.siggenes</code>) |
| <code>rs</code> | minimum pearson correlation coefficient for replicated spots profiles to be averaged (argument of <code>get.siggenes</code>) |
| <code>show.fit</code> | logical indicating whether regression fit curves must be plotted (argument of <code>see.genes</code>) |
| <code>show.lines</code> | logical indicating whether a line must be drawn joining plotted data points for each group (argument of <code>see.genes</code>) |
| <code>pdf</code> | logical indicating whether a pdf results file must be generated (argument of <code>see.genes</code>) |
| <code>cexlab</code> | graphical parameter magnification to be used for x labels in plotting functions |
| <code>legend</code> | logical indicating whether legend must be added when plotting profiles (argument of <code>see.genes</code>) |
| <code>main</code> | title for pdf results file |
| <code>...</code> | other graphical function arguments |

Details

maSigPro finds and display genes with significant profile differences in time series gene expression experiments. The main, compulsory, input parameters for this function are a matrix of gene expression data (see `p.vector` for details) and a matrix describing experimental design (see `make.design.matrix` or `p.vector` for details). In case extended gene ID information is wanted to be included in the result of significant genes, a third IDs matrix containing this information will be required (see `get.siggenes` for details).

Basically in the function calls subsequent steps of the maSigPro approach which is:

- Make a general regression model with dummies to indicate different experimental groups.
- Select significant genes on the basis of this general model, applying `fdr` control.
- Find significant variables for each gene, using stepwise regression.
- Extract and display significant genes for any set of variables or experimental groups.

Value

| | |
|------------------------|--|
| <code>summary</code> | a vector or matrix listing significant genes for the variables given by the function parameters |
| <code>sig.genes</code> | 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 containing: <ul style="list-style-type: none"> <code>sig.profiles</code>: expression values of significant genes. The cluster assignment of each gene is given in the last column <code>coefficients</code>: regression coefficients for significant genes <code>t.score</code>: value of the t statistics of significant genes <code>sig.pvalues</code>: p-values of the regression coefficients for significant genes <code>g</code>: number of genes <code>...</code>: arguments passed by previous functions |

| | |
|---------------|---|
| input.data | input analysis data |
| G | number of input genes |
| edesign | matrix of experimental design |
| dis | regression design matrix |
| min.obs | imputed value for minimal number of true observations |
| p.vector | vector containing the computed p-values of the general regression model for each gene |
| variables | variables in the general regression model |
| g | number of significant genes |
| p.vector.alfa | p-value at FDR = Q control |
| step.method | imputed step method for stepwise regression |
| Q | imputed value for false discovery rate (FDR) control |
| step.alfa | imputed significance level in stepwise regression |
| influ.info | data frame of genes containing influential data |

Author(s)

Ana Conesa, aconesa@ivia.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

[make.design.matrix](#), [p.vector](#), [T.fit](#), [get.siggenes](#), [see.genes](#)

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,
                   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) {
    Ct1 <- c(rnorm(r, a1, var11), rnorm(r, b1, var12), rnorm(r, c1, var13)) # Ct1 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)
  }
  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 <- 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)
rownames(tc.DATA) <- paste("feature", c(1:300), sep = "")
colnames(tc.DATA) <- paste("Array", c(1:36), sep = "")
tc.DATA[sample(c(1:(300*36)), 300)] <- NA # introduce missing values

#### CREATE EXPERIMENTAL DESIGN
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 = "")

#### RUN maSigPro
tc.test <- maSigPro(tc.DATA, edesign, degree = 2, vars = "groups", main = "Test")

tc.test$g # gives number of total significant genes
tc.test$summary # shows significant genes by experimental groups
tc.test$sig.genes$Treat1$sig.pvalues # shows pvalues of the significant coefficients
# in the regression models of the significant genes
# for Control.vs.Treat1 comparison

```

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(
  repvect = edesign[,2], show.fit = FALSE, dis = NULL, step.method = "b

```



```
min.obs = 2, alfa = 0.05, nvar.correction = FALSE, summary.mode = "me
xlab = "time", cex.xaxis = 1, ylim = NULL, main = NULL, cexlab = 0.8,
```

Arguments

| | |
|------------------------------|---|
| <code>data</code> | vector or matrix containing the gene expression data |
| <code>edesign</code> | matrix describing experimental design. Rows must be arrays and columns experiment descriptors |
| <code>time</code> | vector indicating time assignment for each array |
| <code>groups</code> | matrix indicating experimental group to which each array is assigned |
| <code>repvect</code> | index vector indicating experimental replicates |
| <code>show.fit</code> | logical indicating whether regression fit curves must be plotted |
| <code>dis</code> | regression design matrix |
| <code>step.method</code> | stepwise regression method to fit models for cluster mean profiles. It can be either "backward", "forward", "two.ways.backward" or "two.ways.forward" |
| <code>min.obs</code> | minimal number of observations for a gene to be included in the analysis |
| <code>alfa</code> | significance level used for variable selection in the stepwise regression |
| <code>nvar.correction</code> | argument for correcting stepwise regression significance level. See T.fit |
| <code>summary.mode</code> | the method to condensate expression information when more than one gene is present in the data. Possible values are "representative" and "median" |
| <code>show.lines</code> | logical indicating whether a line must be drawn joining plotted data points for reach group |
| <code>groups.vector</code> | vector indicating experimental group to which each variable belongs |
| <code>xlab</code> | label for the x axis |
| <code>cex.xaxis</code> | graphical parameter magnification to be used for x axis in plotting functions |
| <code>ylim</code> | range of the y axis |
| <code>main</code> | plot main title |
| <code>cexlab</code> | graphical parameter magnification to be used for x axis label in plotting functions |
| <code>legend</code> | logical indicating whether legend must be added when plotting profiles |
| <code>sub</code> | plot subtitle |

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 than 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.

Author(s)

Ana Conesa, <aconesa@ivia.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

[PlotProfiles](#)

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,
                   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) {
    Ct1 <- c(rnorm(r, a1, var11), rnorm(r, b1, var12), rnorm(r, c1, var13)) # Ct1 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(Ct1, Tr1, Tr2, Tr3)
    tc.dat <- rbind(tc.dat, gene)
  }
  tc.dat
}

## create 10 genes with profile differences between Ct1, Tr2, and Tr3 groups
tc.DATA <- 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 = "")
colnames(tc.DATA) <- paste("Array", c(1:36), sep = "")

##### CREATE EXPERIMENTAL DESIGN
Time <- rep(c(rep(c(1:3), each = 3)), 4)
Replicates <- rep(c(1:12), each = 3)
Ct1 <- c(rep(1, 9), rep(0, 27))
Tr1 <- c(rep(0, 9), rep(1, 9), rep(0, 18))
Tr2 <- c(rep(0, 18), rep(1, 9), rep(0, 9))
```

```
Tr3 <- c(rep(0, 27), rep(1, 9))
```

```
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, main = NULL, cex.xaxis = 0.5, ylim = NULL,
             repvect, sub = NULL, color.mode = "rainbow")
```

Arguments

| | |
|------------|---|
| data | a matrix containing the gene expression data |
| cond | vector for x axis labeling, typically array names |
| main | plot main title |
| cex.xaxis | graphical parameter magnification to be used for x axis in plotting functions |
| ylim | index vector indicating experimental replicates |
| repvect | index vector indicating experimental replicates |
| sub | plot subtitle |
| color.mode | color scale for plotting profiles. Can be either "rainblow" or "gray" |

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, aconesa@ivia.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

[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,
  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)
  }
  tc.dat
}

## create 10 genes with profile differences between Ctl, Tr2, and Tr3 groups
tc.DATA <- 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 = "")
colnames(tc.DATA) <- paste("Array", c(1:36), sep = "")

PlotProfiles (tc.DATA, cond = colnames(tc.DATA), main = "Time Course",
  repvect = rep(c(1:12), each = 3))
```

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, aconesa@ivia.es

Examples

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

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 = NULL, Q = 0.05, MT.adjust = "BH", min.obs = 3)
```

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 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. Default is 3 (minimum value for a quadratic fit) |

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 |
| <i>p.vector</i> | vector containing the computed p-values |
| G | total number of input genes |
| g | number of genes taken in the regression fit |
| BH.alfa | p-value at FDR Q control when Benjamini & 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, aconesa@ivia.es; 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

See Also

[T.fit](#), [lm](#)

Examples

```
#### 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,
  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) {
    Ct1 <- c(rnorm(r, a1, var11), rnorm(r, b1, var12), rnorm(r, c1, var13)) # Ct1 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(Ct1, Tr1, Tr2, Tr3)
    tc.dat <- rbind(tc.dat, gene)
  }
  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 <- 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)
rownames(tc.DATA) <- paste("feature", c(1:300), sep = "")
colnames(tc.DATA) <- paste("Array", c(1:36), sep = "")
tc.DATA [sample(c(1:(300*36)), 300)] <- NA # introduce missing values

#### CREATE EXPERIMENTAL DESIGN
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$BH.alfa # p.value at FDR control
tc.p$p.adjusted# adjusted p.values

```

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
group)
```

Arguments

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

`indepen` independent 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 term, x^2 term, x^3 term, and so on...

Value

reg.coef `vector of calculated regression coefficients`

Author(s)

Ana Conesa, aconesa@ivia.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.

Examples

```
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.coefcs(coefficients, groups.vector = groups.vector, group = "T1")
```

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 partitioning 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, m = 1.45,
  cluster.method = "hclust", distance = "cor", aggllo.method = "ward",
  show.fit = FALSE, dis = NULL, step.method = "backward", min.obs = 3,
  alfa = 0.05, nvar.correction = FALSE, show.lines = TRUE, iter.max = 500,
  summary.mode = "median", color.mode = "rainbow", cexlab = 1, legend = TRUE,
  newX11 = TRUE, ylim = NULL, main = NULL, ...)
```


Arguments

| | |
|------------------------------|--|
| <code>data</code> | either matrix or a list containing the gene expression data, typically a get.siggenes object |
| <code>edesign</code> | matrix of experimental design |
| <code>time.col</code> | column in <code>edesign</code> containing time values. Default is first column |
| <code>repl.col</code> | column in <code>edesign</code> containing coding for replicates arrays. Default is second column |
| <code>group.cols</code> | columns indicating the coding for each group (treatment, tissue,...) in the experiment (see details) |
| <code>names.groups</code> | names for experimental groups |
| <code>cluster.data</code> | type of data used by the cluster algorithm (see details) |
| <code>groups.vector</code> | vector indicating the experimental group to which each variable belongs |
| <code>k</code> | number of clusters for data partitioning |
| <code>m</code> | m parameter when "mfuzz" clustering algorithm is used. See mfuzz |
| <code>cluster.method</code> | clustering method for data partitioning. Currently "hclust", "kmeans" and "mfuzz" are supported |
| <code>distance</code> | distance measurement function for when <code>cluster.method</code> is hclust |
| <code>agglo.method</code> | aggregation method used when <code>cluster.method</code> is hclust |
| <code>show.fit</code> | logical indicating whether regression fit curves must be plotted |
| <code>dis</code> | regression design matrix |
| <code>step.method</code> | stepwise regression method to fit models for cluster mean profiles. Can be either "backward", "forward", "two.ways.backward" or "two.ways.forward" |
| <code>min.obs</code> | minimal number of observations for a gene to be included in the analysis |
| <code>alfa</code> | significance level used for variable selection in the stepwise regression |
| <code>nvar.correction</code> | argument for correcting <code>T.fitsignificance</code> level. See T.fit |
| <code>show.lines</code> | logical indicating whether a line must be drawn joining plotted data points for reach group |
| <code>iter.max</code> | maximum number of iterations when <code>cluster.method</code> is kmeans |
| <code>summary.mode</code> | the method PlotGroups takes to condensate expression information when more than one gene is present in the data. Possible values are "representative" and "median" |
| <code>color.mode</code> | color scale for plotting profiles. Can be either "rainblow" or "gray" |
| <code>cexlab</code> | graphical parameter magnification to be used for x labels in plotting functions |
| <code>legend</code> | logical indicating whether legend must be added when plotting profiles |
| <code>main</code> | plot title |
| <code>ylim</code> | range of the y axis to be used by PlotProfiles and PlotGroups |
| <code>newX11</code> | when TRUE, plot each type of plot in a diferent graphical device |
| <code>...</code> | 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.

| | |
|--------------------------|---|
| <code>cut</code> | vector indicating gene partitioning into clusters |
| <code>c.algo.used</code> | clustering algorithm used for data partitioning |
| <code>groups</code> | groups matrix used for plotting functions |

Author(s)

Ana Conesa, aconesa@ivia.es; 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

See Also

[PlotProfiles](#), [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,
                   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) {
    Ct1 <- c(rnorm(r, a1, var11), rnorm(r, b1, var12), rnorm(r, c1, var13)) # Ct1 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)
  }
  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 <- 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)
rownames(tc.DATA) <- paste("feature", c(1:300), sep = "")
colnames(tc.DATA) <- paste("Array", c(1:36), sep = "")
tc.DATA [sample(c(1:(300*36)), 300)] <- NA # introduce missing values

#### CREATE EXPERIMENTAL DESIGN
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 = "")

see.genes(tc.DATA, edesign = edesign, k = 4, main = "Time Course")

# This will show the regression fit curve
dise <- make.design.matrix(edesign)
see.genes(tc.DATA, edesign = edesign, k = 4, main = "Time Course", show.fit = TRUE,
          dis = dise$dis, groups.vector = dise$groups.vector, distance = "euclidean")

```

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)
```

Arguments

| | |
|------|---|
| y | 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 |

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 `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, aconesa@ivia.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`, `stepfor`, `two.ways.stepback`, `two.ways.stepfor`

Examples

```
## 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))
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, -0.055, 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,

s.fit <- 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)
```

Arguments

| | |
|-------------------|---|
| <code>y</code> | dependent variable |
| <code>d</code> | data frame containing by columns the set of variables that could be in the selected model |
| <code>alfa</code> | significance level to decide if a variable stays or not in the 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.

Value

`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, aconesa@ivia.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`

Examples

```

## 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))
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, -0.055, 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,

s.fit <- stepfor(y = y, d = dis)
summary(s.fit)

```

suma2Venn

Creates a Venn Diagram from a matrix of characters

Description

suma2Venn transforms a matrix of characters into a binary matrix and creates a vennDiagram of the common elements between columns

Usage

```
suma2Venn(x, ...)
```

Arguments

| | |
|-----|---|
| x | data frame of character values |
| ... | plotting arguments for the vennDiagram function |

Details

suma2Venn creates a list of all elements of a matrix or data frame of characters and computes the presence/absence of each element in each column of the matrix. This results is a numeric matrix of 1 and 0 which can be taken by the [vennDiagram](#) to generate a Venn Plot

Value

suma2Venn returns a Venn Plot such as that created by the [vennDiagram](#) funcion

Author(s)

Ana Conesa, aconesa@ivia.es

See Also[vennDiagram](#)**Examples**

```

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

```

| | |
|-------|--|
| T.fit | <i>Makes a stepwise regression fit for time series gene expression experiments</i> |
|-------|--|

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)

```

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 |

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

Value

| | |
|----------------------------|---|
| <code>sol</code> | matrix for summary results of the stepwise regression. For each selected gene the following values are given: <ul style="list-style-type: none"> • p-value of the regression ANOVA • R-squared of the model • p-value of the regression coefficients of the selected variables |
| <code>sig.profiles</code> | expression values for the genes contained in <code>sol</code> |
| <code>coefficients</code> | matrix containing regression coefficients for the adjusted models |
| <code>groups.coeffs</code> | matrix containing the coefficients of the implicit models of each experimental group |
| <code>variables</code> | variables in the complete regression model |
| <code>G</code> | total number of input genes |
| <code>g</code> | number of genes taken in the regression fit |
| <code>dat</code> | input analysis data matrix |
| <code>dis</code> | regression design matrix |
| <code>step.method</code> | imputed step method for stepwise regression |
| <code>edesign</code> | matrix of experimental design |
| <code>influ.info</code> | data frame of genes containing influential data |

Author(s)

Ana Conesa, <aconesa@ivia.es>; 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

See Also

[p.vector](#), [step](#)

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,
  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) {
  Ct1 <- c(rnorm(r, a1, var11), rnorm(r, b1, var12), rnorm(r, c1, var13)) # Ct1 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(Ct1, Tr1, Tr2, Tr3)
  tc.dat <- rbind(tc.dat, gene)
}
tc.dat
}

## Create 270 flat profiles
flat <- tc.GENE(n = 270, r = 3)
## Create 10 genes with profile differences between Ct1 and Tr1 groups
twodiff <- tc.GENE (n = 10, r = 3, b2 = 0.5, c2 = 1.3)
## Create 10 genes with profile differences between Ct1, 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 Ct1 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)
rownames(tc.DATA) <- paste("feature", c(1:300), sep = "")
colnames(tc.DATA) <- paste("Array", c(1:36), sep = "")
tc.DATA [sample(c(1:(300*36)), 300)] <- NA # introduce missing values

#### CREATE EXPERIMENTAL DESIGN
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 = "")

## run T.fit from a p.vector object
tc.p <- p.vector(tc.DATA, design = make.design.matrix(edesign), Q = 0.01)
tc.tstep <- T.fit(data = tc.p , alfa = 0.05)

## run T.fit from a data matrix and a design matrix
dise <- make.design.matrix(edesign)
tc.tstep <- T.fit (data = tc.DATA[271:300,], design = dise$dis,
                 step.method = "two.ways.backward", min.obs = 10, alfa = 0.05)
tc.tstep$sol # gives the p.values of the significant
             # regression coefficients of the optimized models

```

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)
```

Arguments

| | |
|------|---|
| y | 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 |

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 `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, aconesa@ivia.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`, `stepfor`, `stepback`, `two.ways.stepfor`

Examples

```
## 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))
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, -0.055, 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,

s.fit <- two.ways.stepback(y = y, d = dis)
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)
```

Arguments

| | |
|-------------------|---|
| <code>y</code> | dependent variable |
| <code>d</code> | data frame containing by columns the set of variables that could be in the selected model |
| <code>alfa</code> | significance level to decide if a variable stays or not in the 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 variables 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, aconesa@ivia.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](#), [stepfor](#), [two.ways.stepback](#)

Examples

```
## 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))
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, -0.055, 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,

s.fit <- two.ways.stepfor(y = y, d = dis)
summary(s.fit)
```

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