Package: supclust (via r-universe)

September 13, 2024

Title Supervised Clustering of Predictor Variables Such as Genes

Version 1.1-2

Date 2024-08-14

VersionNote Last CRAN 1.1-1 on 2021-09-25

Description Methodology for supervised grouping aka ``clustering" of potentially many predictor variables, such as genes etc, implementing algorithms 'PELORA' and 'WILMA'.

Imports graphics, stats, rpart, class

URL https://github.com/mmaechler/supclust

Encoding UTF-8

License GPL-3

Repository https://mmaechler.r-universe.dev

RemoteUrl https://github.com/mmaechler/supclust

RemoteRef HEAD

RemoteSha 4268e26e82f9868d2d8be5acd8910bbbbdbe3948

Contents

back.search																												2
coef.pelora																												2
dlda																												3
fitted.pelora			•			•	•	•					•		•	•	•	•		•		•	•	•	•	•	•	4
fitted.wilma																												5
leukemia			•			•	•	•					•		•	•	•	•		•		•	•	•	•	•	•	6
margin																												
pelora			•			•	•	•					•		•	•	•	•		•		•	•	•	•	•		8
plot.pelora .																												
plot.wilma .																												
predict.pelora																												
predict.wilma	a		•			•	•	•					•		•	•	•	•		•		•	•	•	•	•		14
print.pelora															•													16

coef.pelora

print.wilma	 														17
score	 														18
sign.change	 														19
sign.flip	 														20
standardize.genes	 														21
summary.pelora	 														22
summary.wilma	 														23
wilma	 														23
															26

Index

back.search

Internal functions for Supervised Grouping

Description

These are not to be called by the user.

See Also

pelora and wilma

coef.pelora

Extract the Model Coefficients of Pelora

Description

Yields the coefficients of the penalized logistic regression model that is fitted by pelora with its groups of predictor variables (genes) as input

Usage

S3 method for class 'pelora'
coef(object, ...)

Arguments

object	an R object of class "pelora", typically the result of pelora().
	further arguments passed to and from methods.

Value

A numeric vector of length noc + 1, giving the penalized logistic regression coefficients for the intercept and the noc groups and/or single variables identified by pelora.

dlda

Author(s)

Marcel Dettling, <dettling@stat.math.ethz.ch>

See Also

pelora, also for references.

Examples

```
## Running the examples of Pelora's help page
example(pelora, echo = FALSE)
coef(fit)
```

dlda

Classification with Wilma's Clusters

Description

The four functions nnr (nearest neighbor rule), dlda (diagonal linear discriminant analysis), logreg (logistic regression) and aggtrees (aggregated trees) are used for binary classification with the cluster representatives of Wilma's output.

Usage

```
dlda (xlearn, xtest, ylearn)
nnr (xlearn, xtest, ylearn)
logreg (xlearn, xtest, ylearn)
aggtrees(xlearn, xtest, ylearn)
```

Arguments

xlearn	Numeric matrix of explanatory variables (q variables in columns, n cases in rows), containing the learning or training data. Typically, these are the (gene) cluster representatives of Wilma's output.
xtest	A numeric matrix of explanatory variables (q variables in columns, m cases in rows), containing the test or validation data. Typically, these are the fitted (gene) cluster representatives of Wilma's output for the training data, obtained from predict.wilma.
ylearn	Numeric vector of length n containing the class labels for the training observations. These labels have to be coded by 0 and 1.

Details

nnr implements the 1-nearest-neighbor-rule with Euclidean distance function. dlda is linear discriminant analysis, using the restriction that the covariance matrix is diagonal with equal variance for all predictors. logreg is default logistic regression. aggtrees fits a default stump (a classification tree with two terminal nodes) by rpart for every predictor variable and uses majority voting to determine the final classifier.

Value

Numeric vector of length m, containing the predicted class labels for the test observations. The class labels are coded by 0 and 1.

Author(s)

Marcel Dettling

References

see those in wilma.

See Also

wilma

Examples

```
## Generating random learning data: 20 observations and 10 variables (clusters)
set.seed(342)
xlearn <- matrix(rnorm(200), nrow = 20, ncol = 10)
## Generating random test data: 8 observations and 10 variables(clusters)
xtest <- matrix(rnorm(80), nrow = 8, ncol = 10)
## Generating random class labels for the learning data
ylearn <- as.numeric(runif(20)>0.5)
## Predicting the class labels for the test data
nnr(xlearn, xtest, ylearn)
dlda(xlearn, xtest, ylearn)
logreg(xlearn, xtest, ylearn)
aggtrees(xlearn, xtest, ylearn)
```

fitted.pelora Extract the Fitted Values of Pelora

Description

Yields the fitted values, i.e., the centroids of the (gene) groups that have been identified by pelora.

Usage

```
## S3 method for class 'pelora'
fitted(object, ...)
```

fitted.wilma

Arguments

object	An R object of class "pelora", typically the result of pelora().
• • •	Further arguments passed to and from methods.

Value

Numeric matrix of fitted values (for n cases in rows, and noc group centroids in columns).

Author(s)

Marcel Dettling, <dettling@stat.math.ethz.ch>

See Also

pelora, also for references.

Examples

```
## Running the examples of Pelora's help page
example(pelora, echo = FALSE)
fitted(fit)
```

fitted.wilma Extract the Fitted Values of Wilma

Description

Yields the fitted values, i.e. the centroids of the (gene) clusters that have been found by wilma.

Usage

```
## S3 method for class 'wilma'
fitted(object, ...)
```

Arguments

object	An R object of class "wilma", typically the result of wilma().
	further arguments passed to and from methods.

Value

Numeric matrix of fitted values (for n cases in rows, and noc group centroids in columns).

Author(s)

Marcel Dettling, <dettling@stat.math.ethz.ch>

See Also

wilma, also for references.

Examples

```
## Running the examples of Wilma's help page
example(wilma, echo = FALSE)
fitted(fit)
```

leukemia

A part of the Golub's famous AML/ALL-leukemia dataset

Description

Part of the training set of the famous AML/ALL-leukemia dataset from the Whitehead Institute. It has been reduced to 250 genes, about half of which are very informative for classification, whereas the other half was chosen randomly.

Usage

data(leukemia)

Format

Contains three R-objects: The expression (38×250) matrix leukemia.x, the associated binary (0, 1) response variable leukemia.y, and the associated 3-class response variable leukemia.z with values in 0, 1, 2.

Author(s)

Marcel Dettling

Source

originally at http://www.genome.wi.mit.edu/MPR/, (which is not a valid URL any more).

References

First published in Golub et al. (1999) Molecular Classification of Cancer: Class Discovery and Class Prediction by Gene Expression Monitoring. *Science* **286**, 531–538.

6

margin

Examples

```
data(leukemia, package="supclust")
str(leukemia.x)
str(leukemia.y)
str(leukemia.z)
op <- par(mfrow= 1:2)
plot(leukemia.x[,56], leukemia.y)
plot(leukemia.x[,174],leukemia.z)
par(op)</pre>
```

margin

Classification Margin Between Two Sample Classes

Description

For a set of n observations grouped into two classes (for example n expression values of a gene), the margin function measures the size of the gap between the classes. This is the distance between the observation of response class zero having the lowest value, and the individual of with response one having the highest value.

Usage

margin(x, resp)

Arguments

x	Numeric vector of length n , for example containing gene or cluster expression values of n different cases.
resp	Numeric vector of length n containing the "binary" class labels of the cases. Must be coded by 0 and 1.

Value

A numeric value, the margin. Positive margin indicates perfect separation of the response classes, whereas negative margin means imperfect separation.

Author(s)

Marcel Dettling

References

see those in wilma.

See Also

wilma, score is the second statistic that is used there.

Examples

```
data(leukemia, package="supclust")
op <- par(mfrow=c(1,3))
plot(leukemia.x[,69],leukemia.y)
title(paste("Margin = ", round(margin(leukemia.x[,69], leukemia.y),2)))
## Sign-flipping is very important
plot(leukemia.x[,161],leukemia.y)
title(paste("Margin = ", round(margin(leukemia.x[,161], leukemia.y),2)))
x <- sign.flip(leukemia.x, leukemia.y)$flipped.matrix
plot(x[,161],leukemia.y)
title(paste("Margin = ", round(margin(x[,161], leukemia.y),2)))
par(op)
```

```
pelora
```

Supervised Grouping of Predictor Variables

Description

Performs selection and supervised grouping of predictor variables in large (microarray gene expression) datasets, with an option for simultaneous classification. Works in a greedy forward strategy and optimizes the binomial log-likelihood, based on estimated conditional probabilities from penalized logistic regression analysis.

Usage

Arguments

Х	Numeric matrix of explanatory variables (p variables in columns, n cases in rows). For example, these can be microarray gene expression data which should be grouped.
У	Numeric vector of length n containing the class labels of the individuals. These labels have to be coded by 0 and 1.
u	Numeric matrix of additional (clinical) explanatory variables (m variables in columns, n cases in rows) that are used in the (penalized logistic regression) prediction model, but neither grouped nor averaged. For example, these can be 'traditional' clinical variables.
noc	Integer, the number of clusters that should be searched for on the data.
lambda	Real, defaults to $1/32$. Rescaled penalty parameter that should be in $[0, 1]$.
flip	Character string, describing a method how the x (gene expression) matrix should be sign-flipped. Possible are "pm" (the default) where the sign for each variable is determined upon its entering into the group, "cor" where the sign for each variable is determined a priori as the sign of the empirical correlation of that variable with the y-vector, and "none" where no sign-flipping is carried out.

8

pelora

standardize	Logical, defaults to TRUE. Is indicating whether the predictor variables (genes) should be standardized to zero mean and unit variance.
trace	Integer >= 0; when positive, the output of the internal loops is provided; trace >= 2 provides output even from the internal C routines.

Value

pelora returns an object of class "pelora". The functions print and summary are used to obtain an overview of the variables (genes) that have been selected and the groups that have been formed. The function plot yields a two-dimensional projection into the space of the first two group centroids that pelora found. The generic function fitted returns the fitted values, these are the cluster representatives. coef returns the penalized logistic regression coefficients θ_j for each of the predictors. Finally, predict is used for classifying test data with Pelora's internal penalized logistic regression classifier on the basis of the (gene) groups that have been found.

An object of class "pelora" is a list containing:

genes	A list of length noc, containing integer vectors consisting of the indices (column numbers) of the variables (genes) that have been clustered.
values	A numerical matrix with dimension $n\times \mathrm{noc},$ containing the fitted values, i.e. the group centroids $\tilde{x}_j.$
У	Numeric vector of length \boldsymbol{n} containing the class labels of the individuals. These labels are coded by 0 and 1.
steps	Numerical vector of length noc, showing the number of forward/backward cycles in the fitting process of each cluster.
lambda	The rescaled penalty parameter.
noc	The number of clusters that has been searched for on the data.
рх	The number of columns (genes) in the x-matrix.
flip	The method that has been chosen for sign-flipping the x-matrix.
var.type	A factor with noc entries, describing whether the j th predictor is a group of predictors (genes) or a single (clinical) predictor variable.
crit	A list of length noc, containing numerical vectors that provide information about the development of the grouping criterion during the clustering.
signs	Numerical vector of length p , saying whether the <i>i</i> th variable (gene) should be sign-flipped (-1) or not (+1).
samp.names	The names of the samples (rows) in the x-matrix.
gene.names	The names of the variables (columns) in the x-matrix.
call	The function call.

Author(s)

Marcel Dettling, <dettling@stat.math.ethz.ch>

References

Marcel Dettling (2003) *Extracting Predictive Gene Groups from Microarray Data and Combining them with Clinical Variables* https://stat.ethz.ch/Manuscripts/dettling/presentation3.pdf

Marcel Dettling and Peter Bühlmann (2002). Supervised Clustering of Genes. *Genome Biology*, **3**(12): research0069.1-0069.15, doi:10.1186/gb2002312research0069.

Marcel Dettling and Peter Bühlmann (2004). Finding Predictive Gene Groups from Microarray Data. *Journal of Multivariate Analysis* **90**, 106–131, doi:10.1016/j.jmva.2004.02.012

See Also

wilma for another supervised clustering technique.

```
## Working with a "real" microarray dataset
data(leukemia, package="supclust")
## Generating random test data: 3 observations and 250 variables (genes)
set.seed(724)
xN <- matrix(rnorm(750), nrow = 3, ncol = 250)</pre>
## Fitting Pelora
fit <- pelora(leukemia.x, leukemia.y, noc = 3)</pre>
## Working with the output
fit
summary(fit)
plot(fit)
fitted(fit)
coef(fit)
## Fitted values and class probabilities for the training data
predict(fit, type = "cla")
predict(fit, type = "prob")
## Predicting fitted values and class labels for the random test data
predict(fit, newdata = xN)
predict(fit, newdata = xN, type = "cla", noc = c(1,2,3))
predict(fit, newdata = xN, type = "pro", noc = c(1,3))
## Fitting Pelora such that the first 70 variables (genes) are not grouped
fit <- pelora(leukemia.x[, -(1:70)], leukemia.y, leukemia.x[,1:70])</pre>
## Working with the output
fit
summary(fit)
plot(fit)
fitted(fit)
coef(fit)
```

plot.pelora

```
## Fitted values and class probabilities for the training data
predict(fit, type = "cla")
predict(fit, type = "prob")
## Predicting fitted values and class labels for the random test data
predict(fit, newdata = xN[, -(1:70)], newclin = xN[, 1:70])
predict(fit, newdata = xN[, -(1:70)], newclin = xN[, 1:70], "cla", noc = 1:10)
predict(fit, newdata = xN[, -(1:70)], newclin = xN[, 1:70], type = "pro")
```

plot.pelora

2-Dimensional Visualization of Pelora's Output

Description

Yields a projection of the cases (for example n gene expression profiles) into the space of the first two gene group centroids that were identified by pelora.

Usage

Arguments

х	An R object of class "pelora", typically the result of $pelora()$.
main	A character string, giving the title of the plot.
xlab	A character string, giving the annotation of the x-axis.
ylab	A character string, giving the annotation of the x-axis.
col	A numeric vector of length 2, coding the colors that will be used for plotting the class labels.
	Further arguments passed to and from methods.

Author(s)

Marcel Dettling, <dettling@stat.math.ethz.ch>

See Also

pelora, also for references.

```
## Running the examples of Pelora's help page
example(pelora, echo = FALSE)
plot(fit)
```

plot.wilma

Description

Yields a projection of the cases (for example n gene expression profiles) into the space of the first two gene group centroids that were identified by wilma.

Usage

Arguments

х	an R object of class "wilma", typically the result of wilma().
xlab	character string, giving the annotation of the x-axis.
ylab	character string, giving the annotation of the x-axis.
col	a numeric vector of length 2, coding the colors that will be used for plotting the class labels.
main	a character string, giving the title of the plot.
	Further arguments passed to and from methods.

Author(s)

Marcel Dettling, <dettling@stat.math.ethz.ch>

See Also

wilma, also for references.

```
## Running the examples of Wilma's help page
example(wilma, echo = FALSE)
plot(fit)
```

predict.pelora

Description

Yields fitted values, predicted class labels and conditional probability estimates for training and test data, which are based on the gene groups pelora found, and on its internal penalized logistic regression classifier.

Usage

Arguments

object	An R object of class "pelora", typically the result of pelora().
newdata	Numeric matrix with the same number of explanatory variables as the original x-matrix (p variables in columns, r cases in rows). For example, these can be additional microarray gene expression data which should be predicted.
newclin	Numeric matrix with the same number of additional (clinical) explanatory variables as the original u-matrix (m variables in columns, r cases in rows) that are used in the (penalized logistic regression) prediction model, but neither grouped nor averaged. Only needs to be given, if the model fit included an u-matrix. For example, these can be 'traditional' clinical variables.
type	Character string, describing whether fitted values "fitted", estimated condi- tional probabilites "probs" or class labels "class" should be returned.
noc	Integer, saying with how many clusters the fitted values, probability estimates or class labels should be determined. Also numeric vectors are allowed as an argument. The output is then a numeric matrix with fitted values, probability estimates or class labels for a multiple number of clusters.
	Further arguments passed to and from methods.

Details

If newdata = NULL, then the in-sample fitted values, probability estimates and class label predictions are returned.

Value

Depending on whether noc is a single number or a numeric vector. In the first case, a numeric vector of length r is returned, which contains fitted values for noc clusters, or probability estimates/class label predictions with noc clusters.

In the latter case, a numeric matrix with length(noc) columns, each containing fitted values for noc clusters, or probability estimates/class label predictions with noc clusters, is returned.

Author(s)

Marcel Dettling, <dettling@stat.math.ethz.ch>

See Also

pelora, also for references.

Examples

```
## Working with a "real" microarray dataset
data(leukemia, package="supclust")
## Generating random test data: 3 observations and 250 variables (genes)
set.seed(724)
xN <- matrix(rnorm(750), nrow = 3, ncol = 250)</pre>
## Fitting Pelora
fit <- pelora(leukemia.x, leukemia.y, noc = 3)</pre>
## Fitted values and class probabilities for the training data
predict(fit, type = "cla")
predict(fit, type = "prob")
## Predicting fitted values and class labels for the random test data
predict(fit, newdata = xN)
predict(fit, newdata = xN, type = "cla", noc = c(1,2,3))
predict(fit, newdata = xN, type = "pro", noc = c(1,3))
## Fitting Pelora such that the first 70 variables (genes) are not grouped
fit <- pelora(leukemia.x[, -(1:70)], leukemia.y, leukemia.x[,1:70])</pre>
## Fitted values and class probabilities for the training data
predict(fit, type = "cla")
predict(fit, type = "prob")
## Predicting fitted values and class labels for the random test data
predict(fit, newdata = xN[, -(1:70)], newclin = xN[, 1:70])
predict(fit, newdata = xN[, -(1:70)], newclin = xN[, 1:70], "cla", noc = 1:10)
predict(fit, newdata = xN[, -(1:70)], newclin = xN[, 1:70], type = "pro")
```

predict.wilma Predict Method for Wilma

Description

Yields fitted values or predicted class labels for training and test data, which are based on the supervised gene clusters wilma found, and on a choice of four different classifiers: the nearest-neighbor rule, diagonal linear discriminant analysis, logistic regression and aggregated trees.

predict.wilma

Usage

Arguments

object	an R object of class "wilma", typically the result of wilma().
newdata	numeric matrix with the same number of explanatory variables as the original x-matrix (p variables in columns, r cases in rows). For example, these can be additional microarray gene expression data which should be predicted.
type	character string describing whether fitted values "fitted" or predicted class labels "class" should be returned.
classifier	character string specifying which classifier should be used. Choices are "nnr", the 1-nearest-neighbor-rule; "dlda", diagonal linear discriminant analysis; "logreg", logistic regression; "aggtrees" aggregated trees.
noc	integer specifying how many clusters the fitted values or class label predictions should be determined. Also numeric vectors are allowed as an argument. The output is then a numeric matrix with fitted values or class label predictions for a multiple number of clusters.
	further arguments passed to and from methods.

Details

If newdata = NULL, then the in-sample fitted values or class label predictions are returned.

Value

Depending on whether noc is a single number or a numeric vector. In the first case, a numeric vector of length r is returned, which contains fitted values for noc clusters, or class label predictions with noc clusters.

In the latter case, a numeric matrix with length(noc) columns, each containing fitted values for noc clusters, or class label predictions with noc clusters, is returned.

Author(s)

Marcel Dettling, <dettling@stat.math.ethz.ch>

See Also

wilma also for *references*, and for the four classifiers nnr, dlda, logreg, aggtrees.

Examples

```
## Working with a "real" microarray dataset
data(leukemia, package="supclust")
## Generating random test data: 3 observations and 250 variables (genes)
set.seed(724)
xN <- matrix(rnorm(750), nrow = 3, ncol = 250)</pre>
## Fitting Wilma
fit <- wilma(leukemia.x, leukemia.y, noc = 3, trace = 1)</pre>
## Fitted values and class predictions for the training data
predict(fit, type = "cla")
predict(fit, type = "fitt")
## Predicting fitted values and class labels for test data
predict(fit, newdata = xN)
predict(fit, newdata = xN, type = "cla", classifier = "nnr", noc = c(1,2,3))
predict(fit, newdata = xN, type = "cla", classifier = "dlda", noc = c(1,3))
predict(fit, newdata = xN, type = "cla", classifier = "logreg")
predict(fit, newdata = xN, type = "cla", classifier = "aggtrees")
```

print.pelora Print Method for Pelora Objects

Description

Yields an overview about the type, size and final criterion value of the predictor variables that were selected by pelora.

Usage

```
## S3 method for class 'pelora'
print(x, digits = getOption("digits"), details = FALSE, ...)
```

Arguments

х	an R object of class "pelora", typically the result of pelora().
digits	the number of digits that should be printed.
details	logical, defaults to FALSE. If set to TRUE, the output corresponds to summary.pelora.
	Further arguments passed to and from methods.

Author(s)

Marcel Dettling, <dettling@stat.math.ethz.ch>

See Also

pelora, also for references.

16

print.wilma

Examples

```
## Running the examples of Pelora's help page
example(pelora, echo = FALSE)
print(fit)
```

print.wilma

Print Method for Wilma Objects

Description

Yields an overview about the size and the final criterion values of the clusters that were selected by wilma.

Usage

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

Arguments

х	An R object of class "wilma", typically the result of wilma().
	Further arguments passed to and from methods.

Author(s)

Marcel Dettling, <dettling@stat.math.ethz.ch>

See Also

wilma, also for references.

```
## Running the examples of Wilma's help page
example(wilma, echo = FALSE)
print(fit)
```

score

Description

For a set of n observations grouped into two classes (for example n expression values of a gene), the score function measures the separation of the classes. It can be interpreted as counting for each observation having response zero, the number of individuals of response class one that are smaller, and summing up these quantities.

Usage

score(x, resp)

Arguments

х	Numeric vector of length n , for example containing gene or cluster expression values of n different cases.
resp	Numeric vector of length n containing the "binary" class labels of the cases. Must be coded by 0 and 1.

Value

A numeric value, the score. The minimal score is zero, the maximal score is the product of the number of samples in class 0 and class 1. Values near the minimal or maximal score indicate good separation, whereas intermediate score means poor separation.

Author(s)

Marcel Dettling, <dettling@stat.math.ethz.ch>

See Also

wilma also for *references*; margin is the second statistic that is used there.

```
data(leukemia, package="supclust")
op <- par(mfrow=c(1,3))
plot(leukemia.x[,69],leukemia.y)
title(paste("Score = ", score(leukemia.x[,69], leukemia.y)))</pre>
```

```
## Sign-flipping is very important
plot(leukemia.x[,161],leukemia.y)
title(paste("Score = ", score(leukemia.x[,161], leukemia.y),2))
x <- sign.flip(leukemia.x, leukemia.y)$flipped.matrix
plot(x[,161],leukemia.y)
title(paste("Score = ", score(x[,161], leukemia.y),2))
par(op)</pre>
```

sign.change

Description

Computes the empirical correlation for each predictor variable (gene) in the x-Matrix with the response y, and multiplies its values with (-1) if the empirical correlation has a negative sign. For gene expression data, this amounts to treating under- and overexpression symmetrically. After the sign.change, low (expression) values point towards response class 0 and high (expression) values point towards class 1.

Usage

sign.change(x, y)

Arguments

Х	Numeric matrix of explanatory variables (p variables in columns, n cases in rows). For example, these can be microarray gene expression data which should be sign-flipped and then grouped.
У	Numeric vector of length n containing the class labels of the individuals. These labels have to be coded by 0 and 1.

Value

Returns a list containing:

x.new	The sign-flipped x-matrix.
signs	Numeric vector of length p , which for each predictor variable indicates whether it was sign-flipped (coded by -1) or not (coded by +1).

Author(s)

Marcel Dettling, <dettling@stat.math.ethz.ch>

See Also

pelora also for *references*, as well as for older methodology, wilma and sign.flip.

```
data(leukemia, package="supclust")
op <- par(mfrow=c(1,3))
plot(leukemia.x[,69],leukemia.y)
title(paste("Margin = ", round(margin(leukemia.x[,69], leukemia.y),2)))
## Sign-flipping is very important</pre>
```

sign.flip

```
plot(leukemia.x[,161],leukemia.y)
title(paste("Margin = ", round(margin(leukemia.x[,161], leukemia.y),2)))
x <- sign.change(leukemia.x, leukemia.y)$x.new
plot(x[,161],leukemia.y)
title(paste("Margin = ", round(margin(x[,161], leukemia.y),2)))
par(op)</pre>
```

sign.flip

Sign-flipping of Predictor Variables to Obtain Equal Polarity

Description

Computes the score for each predictor variable (gene) in the x-Matrix, and multiplies its values with (-1) if its score is greater or equal than half of the maximal score. For gene expression data, this amounts to treating under- and overexpression symmetrically. After the sign-flip procedure, low (expression) values point towards response class 0 and high (expression) values point towards class 1.

Usage

sign.flip(x, y)

Arguments

x	Numeric matrix of explanatory variables (p variables in columns, n cases in rows). For example, these can be microarray gene expression data which should be sign-flipped and then clustered.
У	Numeric vector of length n containing the class labels of the individuals. These labels have to be coded by 0 and 1.

Value

Returns a list containing:

flipped.matrix	The sign-flipped x-matrix.
signs	Numeric vector of length p , which for each predictor variable indicates whether it was sign-flipped (coded by -1) or not (coded by +1).

Author(s)

Marcel Dettling, <dettling@stat.math.ethz.ch>

See Also

wilma also for the *references* and score, as well as for a newer methodology, pelora and sign.change.

20

standardize.genes

Examples

data(leukemia, package="supclust")

```
op <- par(mfrow=c(1,3))
plot(leukemia.x[,69],leukemia.y)
title(paste("Margin = ", round(margin(leukemia.x[,69], leukemia.y),2)))
## Sign-flipping is very important
plot(leukemia.x[,161],leukemia.y)
title(paste("Margin = ", round(margin(leukemia.x[,161], leukemia.y),2)))
x <- sign.flip(leukemia.x, leukemia.y)$flipped.matrix
plot(x[,161],leukemia.y)
title(paste("Margin = ", round(margin(x[,161], leukemia.y),2)))
par(op)# reset</pre>
```

standardize.genes Standardization of Predictor Variables

Description

Standardizes each column (gene) of the x-matrix to zero mean and unit variance. This function is not to be called by the user, the standardization is handled internally in pelora.

Usage

```
standardize.genes(exmat)
```

Arguments

exmat	Numeric matrix of explanatory variables (p variables in columns, n cases in
	rows). For example, these can be microarray gene expression data which should
	be standardized and then grouped.

Value

Returns a list containing:

x	The standardized x-matrix
means	Numeric vector of length p , containing the initial mean of each column (gene) of the x-matrix.
sdevs	Numeric vector of length p , containing the initial standard deviation of each column (gene) of the x-matrix.

Author(s)

Marcel Dettling, <dettling@stat.math.ethz.ch>

See Also

pelora also for the *references*.

summary.pelora Summary Method for Pelora Objects

Description

Yields detailed information about the variables (genes) that have been selected, and how they were grouped.

Usage

S3 method for class 'pelora'
summary(object, digits, ...)

Arguments

object	an R object of class "pelora", typically the result of pelora()
digits	The number of digits that should be printed.
	Further arguments passed to and from methods.

Author(s)

Marcel Dettling, <dettling@stat.math.ethz.ch>

See Also

pelora, also for references.

```
## Running the examples of Pelora's help page
example(pelora, echo = FALSE)
summary(fit)
```

summary.wilma

Description

Yields detailed information about the variables (genes) that have been selected, and how they were clustered.

Usage

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

Arguments

object	An R object of class "wilma", typically the result of wilma().
	Further arguments passed to and from methods.

Author(s)

Marcel Dettling, <dettling@stat.math.ethz.ch>

See Also

wilma, also for references.

Examples

```
## Running the examples of Wilma's help page
example(wilma, echo = FALSE)
summary(fit)
```

wilma

Supervised Clustering of Predictor Variables

Description

Performs supervised clustering of predictor variables for large (microarray gene expression) datasets. Works in a greedy forward strategy and optimizes a combination of the Wilcoxon and Margin statistics for finding the clusters.

Usage

```
wilma(x, y, noc, genes = NULL, flip = TRUE, once.per.clust = FALSE, trace = 0)
```

Arguments

х	Numeric matrix of explanatory variables (p variables in columns, n cases in rows). For example, these can be microarray gene expression data which should be clustered.
У	Numeric vector of length n containing the class labels of the individuals. These labels have to be coded by 0 and 1.
noc	Integer, the number of clusters that should be searched for on the data.
genes	Defaults to NULL. An optional list (of length noc) of vectors containing the in- dices (column numbers) of the previously known initial clusters.
flip	Logical, defaults to TRUE. Is indicating whether the clustering should be done with or without sign-flipping.
once.per.clust	Logical, defaults to FALSE. Is indicating if each variable (gene) should only be allowed to enter into each cluster once; equivalently, the cluster mean profile has only weights ± 1 for each variable.
trace	Integer >= 0; when positive, the output of the internal loops is provided; trace >= 2 provides output even from the internal C routines.

Value

wilma returns an object of class "wilma". The functions print and summary are used to obtain an overview of the clusters that have been found. The function plot yields a two-dimensional projection into the space of the first two clusters that wilma found. The generic function fitted returns the fitted values, these are the cluster representatives. Finally, predict is used for classifying test data on the basis of Wilma's cluster with either the nearest-neighbor-rule, diagonal linear discriminant analysis, logistic regression or aggregated trees.

An object of class "wilma" is a list containing:

clist	A list of length noc, containing integer vectors consisting of the indices (column numbers) of the variables (genes) that have been clustered.
steps	Numerical vector of length noc, showing the number of forward/backward cycles in the fitting process of each cluster.
у	Numeric vector of length n containing the class labels of the individuals. These labels have to be coded by 0 and 1.
x.means	A list of length noc, containing numerical matrices consisting of the cluster representatives after insertion of each variable.
noc	Integer, the number of clusters that has been searched for on the data.
signs	Numerical vector of length p , saying whether the <i>i</i> th variable (gene) should be sign-flipped (-1) or not (+1).

Author(s)

Marcel Dettling, <dettling@stat.math.ethz.ch>

References

Marcel Dettling and Peter Bühlmann (2002). Supervised Clustering of Genes. *Genome Biology*, **3**(12): research0069.1-0069.15, doi:10.1186/gb2002312research0069.

wilma

See Also

score, margin, and for a newer methodology, pelora.

```
## Working with a "real" microarray dataset
data(leukemia, package="supclust")
## Generating random test data: 3 observations and 250 variables (genes)
set.seed(724)
xN <- matrix(rnorm(750), nrow = 3, ncol = 250)</pre>
## Fitting Wilma
fit <- wilma(leukemia.x, leukemia.y, noc = 3, trace = 1)</pre>
## Working with the output
fit
summary(fit)
plot(fit)
fitted(fit)
## Fitted values and class predictions for the training data
predict(fit, type = "cla")
predict(fit, type = "fitt")
## Predicting fitted values and class labels for test data
predict(fit, newdata = xN)
predict(fit, newdata = xN, type = "cla", classifier = "nnr", noc = c(1,2,3))
predict(fit, newdata = xN, type = "cla", classifier = "dlda", noc = c(1,3))
predict(fit, newdata = xN, type = "cla", classifier = "logreg")
predict(fit, newdata = xN, type = "cla", classifier = "aggtrees")
```

Index

* classif back.search, 2 coef.pelora, 2 dlda, 3 fitted.pelora, 4 fitted.wilma, 5 pelora, 8 plot.pelora, 11 plot.wilma, 12 predict.pelora, 13 predict.wilma, 14 print.pelora, 16 print.wilma, 17 summary.pelora, 22 summary.wilma, 23 * cluster back.search, 2 coef.pelora, 2 fitted.pelora, 4 fitted.wilma, 5 pelora, 8 plot.pelora, 11 plot.wilma, 12 predict.pelora, 13 predict.wilma, 14 print.pelora, 16 print.wilma, 17 summary.pelora, 22 summary.wilma, 23 wilma, 23 * datasets leukemia. 6 * htest margin,7 score, 18 * manip sign.change, 19 sign.flip, 20 standardize.genes, 21 aggtrees, 15 aggtrees (dlda), 3 back.search, 2 class, 2, 5, 11-13, 15-17, 22, 23 coef.pelora, 2 dlda, 3, 15 fitted.pelora, 4 fitted.wilma, 5 leukemia, 6 logreg, 15 logreg (dlda), 3 margin, 7, 18, 25 nnr, 15 nnr (dlda), 3 p.1clust (back.search), 2 pelora, 2-5, 8, 11, 13, 14, 16, 19, 20, 22, 25 plot.pelora, 11 plot.wilma, 12 predict.pelora, 13 predict.wilma, 14 print.pelora, 16 print.wilma, 17 printClist (back.search), 2 ridge.coef (back.search), 2 score, 7, 18, 20, 25 sign.change, 19, 20 sign.flip, 19, 20 standardize.genes, 21 summary.pelora, 16, 22 summary.wilma, 23

wilma, 2, 4-7, 10, 12, 15, 17-20, 23, 23