Package 'plrs'

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Title Piecewise Linear Regression Splines (PLRS) for the association between DNA copy number and gene expression

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Suggests mytnorm, methods

Description The present package implements a flexible framework for modeling the relationship between DNA copy number and gene expression data using Piecewise Linear Regression Splines (PLRS).

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```
plrs-package
```

Piecewise Linear Regression Splines (PLRS) for the association between DNA copy number and mRNA expression

Description

The present package implements a framework for modeling the relationship between DNA copy number and gene expression data using Piecewise Linear Regression Splines (PLRS). It includes (point and interval) estimation, model selection and testing procedures for such models (possibly under biologically motivated constraints).

Details

The use of the present package can be divided into two approaches:

1. Analysis of a single DNA-mRNA relationship

Main functions are:

plrs: Fit a single plrs model.

plrs.select: Model selection based on AIC, AICC, OSAIC or BIC.

plrs.test: Likelihood ratio test for a given plrs model.

plrs.cb: Confidence bands for a plrs model.

2. Analysis of multiple DNA-mRNA relationships sequentially

Main function is:

plrs.series: point and interval estimation, model selection and testing of DNA-mRNA association for a series of arrays.

Note: This function extend the aforementioned univariate analysis genomewise in the same spirit as some functions of the **limma** package do.

Author(s)

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References

Leday GGR, Van der Vaart AW, Van Wieringen WN, Van de Wiel MA. Modeling association between DNA copy number and gene expression with constrained piecewise linear regression splines. Accepted for publication. *Ann Appl Stat.* (2012).

criteria

Description

Extract AIC, AICC, BIC and OSAIC from an object of class plrs-class.

Usage

```
criteria(obj, crit = "all")
```

Arguments

obj	object of class plrs-class
crit	A character (vector) among "aic", "aicc", "bic", "osaic" or "all".

Value

A list with the following components (if specified):

aic	Akaike's information criterion
aicc	Small sample correction of AIC
bic	Bayesian Information Criterion
osaic	One-Sided AIC. See Hughes and King (2003) for more details.

Author(s)

Gwenael G.R. Leday <g.g.r.leday@vu.nl>

References

Hughes, A. W. and King, M. L. (2003). Model selection using AIC in the presence of one-sided information. *J Stat Plan Infer*, 115(2): 397 411.

Examples

```
# Simulate data
sim <- plrs.sim(n=80, states=4, sigma=0.5)
# Fit
model <- plrs(expr=sim$expr, cghseg=sim$seg, cghcall=sim$cal)
criteria(model)</pre>
```

```
modify.conf
```

Description

This function changes the discrete copy number values for a given gene in order to force a minimum number of observations per state.

Usage

modify.conf(cghcall, min.obs = 3, discard = TRUE)

Arguments

cghcall	Vector of called values
min.obs	Minimum number of observations per state
discard	Logical. Whether discrete states with few observations should be discarded from analysis.

Details

Consider that the number of observations of a given state is lower than min.obs, then:

- if discard = FALSE, observations are not discarded and a rearrangement of called values is carried out as follows. The "normal" copy number state is taken as a reference. If the minimum number of observations is not obtained, "losses" will be merged to "normals", "gains" to "normals" and "amplifications" to "gains". Note that this modifies the configuration of the model. Thus, after fitting a model using plrs, original and modified data are stored in the resulting plrs-class object, respectively under slots data and mdata.

- if discard = TRUE, states for which the number of observations is lower than min.obs are discarded (replaced by NAs).

Value

val

Vector of new called values

Note

This function is implemented within function plrs and plrs.series.

Author(s)

Gwenael G.R. Leday <g.g.r.leday@vu.nl>

Examples

```
called <- sample(c(rep(-1,5),rep(0,15),rep(1,2),rep(2,1)))
table(called)
table(modify.conf(called, min.obs=3))</pre>
```

neveCN17

Description

Preprocessed copy number data of Neve et al. (2006) for chromosome 17.

Usage

neveCN17

Format

An object of class cghCall

Source

M. Neve et al. in Gray Lab at LBL. Neve2006: expression and CGH data on breast cancer cell lines. R package version 0.1.10.

References

Neve, R.M. et al. (2006). A collection of breast cancer cell lines for the study of functionally distinct cancer subtypes. *Cancer cell*, 10, 515-527.

Examples

data(neveCN17)
dim(neveCN17)
head(fData(neveCN17))

neveGE17

mRNA expression for chromosome 17.

Description

Normalized gene expression data of Neve et al. (2006) for chromosome 17.

Usage

neveGE17

Format

An object of class ExpressionSet

Source

M. Neve et al. in Gray Lab at LBL. Neve2006: expression and CGH data on breast cancer cell lines. R package version 0.1.10.

References

Neve, R.M. et al. (2006). A collection of breast cancer cell lines for the study of functionally distinct cancer subtypes. *Cancer cell*, 10, 515-527.

Examples

```
data(neveGE17)
dim(neveGE17)
head(fData(neveGE17))
```

plot-methods

Plot functions in package 'plrs'

Description

Methods plot in package 'plrs'

Usage

```
## S3 method for class 'plrs'
plot(x, col.line = "black", col.pts = c("red", "blue","green2", "green4"),
col.cb = "yellow", xlim = c(floor(min(x@data$cghseg)),ceiling(max(x@data$cghseg))),
ylim = c(floor(min(x@data$expr)),ceiling(max(x@data$expr))),
pch = 16, lwd=4, cex = 1.2, xlab="", ylab="", main = "",
add = FALSE, lty = 1, lin = FALSE, ...)
```

Arguments

х	An object of class plrs-class or plrs.select-class
col.line	Color of the fitted line
col.pts	Vector of length 4, for colors associated with each state
col.cb	Color for the confidence band
xlim	The x limits of the plot
ylim	The y limits of the plot
pch	See par
lwd	See par
cex	See par
xlab	Title of the x-axis
ylab	Title of the y-axis
main	Main title for the plot
add	If the plot should be added to the current device. Default is FALSE
lty	See par
lin	Logical. Whether the simple linear model should also be plotted
	Other arguments, see par

plrs

Details

plot.plrs plots the observed points, the fitted line and potentially the confidence band.

Methods

signature(x = "plrs") Plot observed points and the fitted line signature(x = "plrs.select") Plot observed points and the fitted line of the selected model.

Author(s)

Gwenael G.R. Leday <g.g.r.leday@vu.nl>

plrs

Fit a (constrained) piecewise linear regression spline

Description

The function fits a piecewise linear regression spline to explain gene expression by the segmented DNA copy number. The called copy number values are used as a template for model building.

Usage

```
plrs(expr, cghseg, cghcall=NULL, probloss = NULL, probnorm = NULL,
probgain = NULL, probamp = NULL, knots = NULL, continuous = FALSE,
constr = TRUE, constr.slopes = 2, constr.intercepts = TRUE,
min.obs = 3, discard.obs = TRUE)
```

Arguments

expr	Vector of gene expression values
cghseg	Vector of segmented copy number values
cghcall	Vector of called copy number values. If not provided, we are reduced to a simple linear model.
probloss	Vector of call probabilities associated with state "loss". Default is NULL.
probnorm	Vector of call probabilities associated with state "normal". Default is NULL.
probgain	Vector of call probabilities associated with state "gain". Default is NULL.
probamp	Vector of call probabilities associated with state "amplification". Default is NULL.
knots	knots or change points. If NULL (default), there are estimated. See details.
continuous	Logical, whether the model is continuous (no jump) or not.
constr	Logical, whether the model is constrained or not. (this has been implemented to turn on and off easily the constraints)
constr.slopes	Type of non-negativity constraints applied on slopes. Either 1 or 2 (default). See details.
constr.interce	
	If TRUE (default) jumps from state to state are also constrained to be non-negative
min.obs	See modify.conf
discard.obs	See modify.conf

Details

If cghcall=NULL, discrete copy number values are omitted, which results in fitting a simple linear model.

If constr.slopes=1, all slopes are constrained to be non-negative. If constr.slopes=2, the slope associated with state "normal" is constrained to be non-negative and all others are forced to be at least equal to the latter.

Two methods are implemented for the estimation of knots. If call probabilities are provided, a knot is determined so that the sum of (the two adjacent) states membership probabilities is maximized. Otherwise, this is defined as the midpoint of the interval between the two consecutive states.

The constrained least squares problem is solved using function solve.QP of package quadprog.

Value

An object of class plrs-class

Author(s)

Gwenael G.R. Leday <g.g.r.leday@vu.nl>

Examples

```
# Simulate data
sim <- plrs.sim(n=80, states=4, sigma=0.5)
# Fit a model
model <- plrs(expr=sim$expr, cghseg=sim$seg, cghcall=sim$cal)
model
# Methods
coef(model)
effects(model)
fitted(model)
knots(model)
model.matrix(model)
plot(model)
plot(model)
predict(model, newcghseg=seq(0,5, length.out=100))
residuals(model)
summary(model)
```

plrs-class

Class plrs

Description

An S4 class representing the output of the plrs function.

plrs-class

Slots

coefficients: Object of class numeric containing spline coefficients

fitted.values: Object of class numeric containing the fitted values

residuals: Object of class numeric containing the residuals

X: Object of class matrix containing the design matrix

data: Object of class list containing input data

- mdata: Object of class list containing (possibly modified) data used to fit the model (See modify.conf).
- QP: Object of class list containing input elements used for quadratic programming. If the model is unconstrained this contains a light version of an 1m object.

test: Object of class list containing results from testing.

cb: Object of class list containing lower and upper bounds for predicted values.

selected: Object of class logical indicating whether the model results from a selection procedure.

type: Object of class character giving the type of model

call.arg: Object of class list containing the input arguments (for reproducibility)

Methods

coef Returns the coefficients

criteria See criteria

effects Returns matrix of effects

fitted Returns the fitted values

knots Returns the knots

model.matrix Returns the design matrix

plot See plot.plrs

predict See predict.plrs

print Print the object information

residuals Returns the residuals

show Print the object information

summary Print a summary of the object information

Author(s)

Gwenael G.R. Leday <g.g.r.leday@vu.nl>

plrs.cb

Description

Determine uniform confidence intervals for predicted values of a 'plrs' model.

Usage

```
plrs.cb(object, alpha=0.05, newcgh=NULL)
```

Arguments

object	An object of class plrs-class.
alpha	Significance level
newcgh	Vector of segmented values. Support for building CB.

Details

The input object of class plrs-class has to result from function plrs.test.

The problem of finding (at a given x) a confidence interval for the mean response is expressed as a semi-definite optimization problem and solved using function csdp of package **Rcsdp**.

Value

An object of class plrs-class that contains CB information.

Author(s)

Gwenael G.R. Leday <g.g.r.leday@vu.nl>

References

Leday GGR, Van der Vaart AW, Van Wieringen WN, Van de Wiel MA. Modeling association between DNA copy number and gene expression with constrained piecewise linear regression splines. Accepted for publication. *Ann Appl Stat.* (2012).

See Also

plrs.test

Examples

```
# Simulate data
sim <- plrs.sim(n=80, states=4, sigma=0.5)
# Fit a model
model <- plrs(expr=sim$expr, cghseg=sim$seg, cghcall=sim$cal)</pre>
```

Confidence bands

plrs.select

```
model <- plrs.test(model)
model <- plrs.cb(model, alpha=0.05)
plot(model)</pre>
```

plrs.select Model selection

Description

Selection of a model based on an information criterion (AIC, AICC, BIC or OSAIC).

Usage

```
plrs.select(object, crit = ifelse(object@call.arg$constr,"osaic","aic"))
```

Arguments

object	An object of class plrs-class
crit	Character corresponding to the criterion to use. See criteria.

Value

An object of class plrs.select-class

Author(s)

Gwenael G.R. Leday <g.g.r.leday@vu.nl>

plrs.select-class Class plrs.select

Description

An S4 class representing the output of the plrs.select function.

Slots

table: Object of class matrix containing the criterion value for all models
model: Object of class plrs containing the selected model
crit: Object of class character containing the criterion used for model selection

Methods

plot See plot.plrs
print Print the object information
show Print the object information
summary Print a summary of the object information

Author(s)

Gwenael G.R. Leday <g.g.r.leday@vu.nl>

```
plrs.series
```

Description

The function fits plrs models for a series of arrays. Model selection and testing procedures may be applied.

Usage

```
plrs.series(expr, cghseg, cghcall=NULL,
probloss = NULL, probnorm = NULL, probgain = NULL, probamp = NULL,
control.model = list(continuous = FALSE,
                      constr = TRUE,
                      constr.slopes = 2,
                      constr.intercepts = TRUE,
                      min.obs = 3,
                      discard.obs = TRUE),
control.select = list(crit = ifelse(control.model$constr, "osaic", "aic")),
control.test = list(testing = TRUE,
                      cb = FALSE,
                      alpha = 0.05),
control.output = list(save.models = FALSE,
                      save.plots = FALSE,
                      plot.lin = FALSE,
                      type = "jpeg"))
```

Arguments

expr	Either a matrix of expression profiles or an ExpressionSet object.
cghseg	Either a matrix of segmented copy number values or objects of class cghSeg or cghCall
cghcall	Matrix of called copy number
probloss	Matrix of call probabilities associated with state "loss". Default is NULL.
probnorm	Matrix of call probabilities associated with state "normal". Default is NULL.
probgain	Matrix of call probabilities associated with state "gain". Default is NULL.
probamp	Matrix of call probabilities associated with state "amplification". Default is NULL.
control.model	See details
<pre>control.select</pre>	See details
control.test	See details
control.output	See details

plrs.series

Details

If DNA and mRNA input data are matrices, rows should correspond to genes and columns to arrays. Alternatively, expression data may be provided as an ExpressionSet object and aCGH data as cghSeg or cghCall objects. A cghCall object contain all data from the calling step, thus arguments probloss, probnorm, probnorm and probamp can be omitted. An object of class cghSeg does not contain such data so only simple linear models will be fitted.

control.model allows the user to specify the type of model that has to be fitted. This must be a list with one or more of the following components: constr.slopes, constr.intercepts, min.obs and discard.obs. See functions plrs and modify.conf for more details.

control.select allows the user to specify whether model selection should be done and how. This must be a list with a component named crit. See function plrs.select for more details. If control.select = NULL then no model selection is done.

control.output allows the user to plot and save each plrs model. This must be a list with components:

save.models, a logical. This will create within the work directory a new directory named "plrsSeriesObjects" that will contain all objects.

save.plots, a logical. This will create within the work directory a new directory named "plrsSeries-Plots" that will contains all saved plots.

plot.lin, a logical. Whether the simple linear model should aslo be plotted.

type, a character. Format of file. To pass through function savePlot.

Value

An object of class plrs.series-class

Author(s)

Gwenael G.R. Leday <g.g.r.leday@vu.nl>

Examples

```
# Simulate data
ngenes <- 10
narray <- 48
rna <- dnaseg <- dnacal <- matrix(NA, ngenes, narray)
idx <- sample(1:4, ngenes, replace=TRUE, prob=rep(1/4,4))
for(i in 1:ngenes){
Sim <- plrs.sim(n=narray, states=idx[i], sigma=0.5)
rna[i,] <- Sim$expr
dnaseg[i,] <- Sim$seg
dnacal[i,] <- Sim$cal
}
# Screening procedure with linear model
series <- plrs.series(expr = rna, cghseg = dnaseg, cghcall = NULL, control.select = NULL)
# Screening procedure with full plrs model
series <- plrs.series(expr = rna, cghseg = dnaseg, cghcall = dnacal, control.select = NULL)</pre>
```

```
# Model selection
series <- plrs.series(expr = rna, cghseg = dnaseg, cghcall = dnacal)</pre>
```

plrs.series-class Class plrs.series

Description

An S4 class representing the output of the plrs.series function.

Slots

coefficients: Matrix containing coefficients of models

effects: List containing effects

test: Matrix containing results from testing.

general: Matrix providing the distribution of the number genes and arrays regarding the copy number states

modelsType: List providing models' type

call.arg: List providing details on the type of models that have been fitted.

Methods

print Print the object information**show** Print the object information

summary Print a summary of the object information

Author(s)

Gwenael G.R. Leday <g.g.r.leday@vu.nl>

plrs.sim Simulation of a plrs model

Description

Simulation of a piecewise relationship.

The function has been only implemented for convenience of simulations and R examples.

Usage

plrs.sim(n = 80, states = 4, sigma = 01, x = NULL)

plrs.test

Arguments

n	Number of simulated data points
states	Number of states for the model
sigma	Noise
х	Segmented values.

Details

To be written...

Author(s)

Gwenael G.R. Leday <g.g.r.leday@vu.nl>

Examples

```
# Simulate 1-state model
sim <- plrs.sim(n=80, states=1, sigma=0.5)
model <- plrs(expr=sim$expr, cghseg=sim$seg, cghcall=sim$cal)
plot(model)</pre>
```

```
# Simulate 2-state model
sim <- plrs.sim(n=80, states=2, sigma=0.5)
model <- plrs(expr=sim$expr, cghseg=sim$seg, cghcall=sim$cal)
plot(model)</pre>
```

```
# Simulate 3-state model
sim <- plrs.sim(n=90, states=3, sigma=0.5)
model <- plrs(expr=sim$expr, cghseg=sim$seg, cghcall=sim$cal)
plot(model)
# Simulate 4-state model</pre>
```

```
sim <- plrs.sim(n=80, states=4, sigma=0.5)
model <- plrs(expr=sim$expr, cghseg=sim$seg, cghcall=sim$cal)
plot(model)</pre>
```

plrs.test

Likelihood ratio test for a plrs model

Description

Test whether copy number has an effect on mRNA expression.

Usage

plrs.test(object, alpha=0.05)

Arguments

object	An object of class plrs-class
alpha	Significance level

Details

Two cases present themselves:

1. The model is unconstrained. Thus, the model under the null hypothesis is the intercept and an F-test is performed.

2. The model is constrained and the following hypothesis are tested:

H0: All constraints are actives (=)

H1: At least one constraint is strict (>)

Under H0, we always have the intercept model. Indeed, if constr.slopes = 1 (or 2) and constr.intercepts = T, then the only parameter free of inequality constraint is the overall intercept. If constr.intercepts = F, the local intercepts are additionally constrained to be 0 in order to obtain the intercept model under the null. The likelihood ratio statistic (unknown variance) is asymptotically distributed as a weighted mixture of Beta distribution (cf Gromping (2010)). Calculation of p-values is based on functions ic.weights and pbetabar of package **ic.infer**. The package **mvtnorm** is also involved.

In both cases the input model is taken as the model under the alternative.

Value

A list object with the following components:

stat	Test statistic
pvalue	Calculated pvalue
wt.bar	Weights (if the model is constrained)
df.bar	Degrees of freedom.
unconstr	Unconstrained model of class plrs-class
qbetabar	(1-alpha) quantile of the beta mixture distribution
alpha	Significance level

Author(s)

Gwenael G.R. Leday <g.g.r.leday@vu.nl>

References

Gromping, U. (2010). Inference with linear equality and inequality constraints using R: The package ic.infer. *J Stat Softw*, 33(i10).

Examples

```
# Simulate data
sim <- plrs.sim(n=80, states=2, sigma=0.5)
# Fit a model
model <- plrs(expr=sim$expr, cghseg=sim$seg, cghcall=sim$cal)
# Testing
model <- plrs.test(model)
model</pre>
```

predict.plrs

Description

Determine predicted values based on a given plrs model

Usage

```
## S3 method for class 'plrs'
predict(object, newcghseg, ...)
```

Arguments

object	An object of class plrs-class
newcghseg	A vector of new segmented CGH values
	further arguments

Value

A vector containing the fitted values.

Author(s)

Gwenael G.R. Leday <g.g.r.leday@vu.nl>

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*Topic copy number, gene expression, regression splines, model selection, constrained inference.

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