Package ‘mda’

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**bruto**

*Fit an Additive Spline Model by Adaptive Backfitting*

**Description**

Fit an additive spline model by adaptive backfitting.

**Usage**

```r
bruto(x, y, w, wp, dfmax, cost, maxit.select, maxit.backfit, thresh = 0.0001, trace.bruto = FALSE, start.linear = TRUE, fit.object, ...)
```

**Arguments**

- `x`  
  a matrix of numeric predictors (does not include the column of 1s).

- `y`  
  a vector or matrix of responses.

- `w`  
  optional observation weight vector.

- `wp`  
  optional weight vector for each column of `y`; the RSS and GCV criteria use a weighted sum of squared residuals.

- `dfmax`  
  a vector of maximum df (degrees of freedom) for each term.

- `cost`  
  cost per degree of freedom; default is 2.

- `maxit.select`  
  maximum number of iterations during the selection stage.

- `maxit.backfit`  
  maximum number of iterations for the final backfit stage (with fixed lambda).

- `thresh`  
  convergence threshold (default is 0.0001); iterations cease when the relative change in GCV is below this threshold.

- `trace.bruto`  
  logical flag. If `TRUE` (default) a progress report is printed during the fitting.

- `start.linear`  
  logical flag. If `TRUE` (default), the model starts with the linear fit.

- `fit.object`  
  This the object returned by `bruto()`; if supplied, the same model is fit to the presumably new `y`.

- `...`  
  further arguments to be passed to or from methods.
Value

A multiresponse additive model fit object of class "bruto" is returned. The model is fit by adaptive backfitting using smoothing splines. If there are np columns in y, then np additive models are fit, but the same amount of smoothing (df) is used for each term. The procedure chooses between df = 0 (term omitted), df = 1 (term linear) or df > 0 (term fitted by smoothing spline). The model selection is based on an approximation to the GCV criterion, which is used at each step of the backfitting procedure. Once the selection process stops, the model is backfit using the chosen amount of smoothing.

A bruto object has the following components of interest:

- **lambda**: a vector of chosen smoothing parameters, one for each column of x.
- **df**: the df chosen for each column of x.
- **type**: a factor with levels "excluded", "linear" or "smooth", indicating the status of each column of x.
- **gcv.select**
- **gcv.backfit**
- **df.select**
- **gcvNselect**
- **gcvNbackfit**
- **dfNselect**
- **nit**: the number of iterations used.
- **fitted.values**: a matrix of fitted values.
- **residuals**: a matrix of residuals.
- **call**: the call that produced this object.

References


Trevor Hastie, Rob Tibshirani and Andreas Buja “Flexible Discriminant Analysis by Optimal Scoring” JASA 1994, 89, 1255-1270.

See Also

- `predict.bruto`

Examples

data(trees)
fit1 <- bruto(trees[-3], trees[3])
fit1$type
fit1$df
## examine the fitted functions
par(mfrow=c(1,2), pty="s")
Xp <- matrix(sapply(trees[1:2], mean), nrow(trees), 2, byrow=TRUE)
for(i in 1:2) {
  xr <- sapply(trees, range)
  Xp1 <- Xp; Xp1[,i] <- seq(xr[1,i], xr[2,i], len=nrow(trees))
  XF <- predict(fit1, Xp1)
  plot(Xp1[,i], XF, xlab=names(trees)[i], ylab="", type="l")
}
coef.fda  

Produce coefficients for an fda or mda object

Description

a method for coef for extracting the canonical coefficients from an fda or mda object

Usage

```r
## S3 method for class 'fda'
coef(object, ...)
```

Arguments

- `object` an fda or mda object.
- `...` not relevant

Details

See the references for details.

Value

A coefficient matrix

Author(s)

Trevor Hastie and Robert Tibshirani

References

“Flexible Discriminant Analysis by Optimal Scoring” by Hastie, Tibshirani and Buja, 1994, JASA, 1255-1270.


“Elements of Statistical Learning - Data Mining, Inference and Prediction” (2nd edition, Chapter 12) by Hastie, Tibshirani and Friedman, 2009, Springer

See Also

`predict.fda`, `plot.fda`, `mars`, `bruto`, `polyreg`, `softmax`, `confusion`, `coef.fda`
confusion

Examples

```r
data(iris)
irisfit <- fda(Species ~ ., data = iris)
coef(irisfit)
mfit = mda(Species ~ ., data = iris, subclass = 2)
coef(mfit)
```

---

confusion

Confusion Matrices

Description

Compute the confusion matrix between two factors, or for an fda or mda object.

Usage

```r
## Default S3 method:
confusion(object, true, ...)
## S3 method for class 'fda'
confusion(object, data, ...)
```

Arguments

- `object`: the predicted factor, or an fda or mda model object.
- `true`: the true factor.
- `data`: a data frame (list) containing the test data.
- `...`: further arguments to be passed to or from methods.

Details

This is a generic function.

Value

For the default method essentially `table(object, true)`, but with some useful attribute(s).

See Also

`fda`, `predict.fda`
**Examples**

```r
data(iris)
irisfit <- fda(Species ~ ., data = iris)
confusion(predict(irisfit, iris), iris$Species)
##
##       Setosa Versicolor Virginica
## Setosa      50          0         0
## Versicolor   0         48         1
## Virginica    2          2        49
## attr(, "error"): [1] 0.02
```

---

**ESL.mixture**

*Mixture example from “Elements of Statistical Learning”*

---

**Description**

A list with training data and other details for the mixture example.

**Usage**

```r
data(ESL.mixture)
```

**Format**

This list contains the following elements:

- `x` a 200x2 matrix of predictors.
- `y` a 200 vector of y values taking values 0 or 1.
- `xnew` a 6831x2 matrix of prediction points, on a 69x99 grid.
- `prob` a vector of 6831 probabilities - the true probabilities of a 1 at each point in `xnew`.
- `marginal` the marginal distribution of the predictors t each point in `xnew`.
- `px1` grid values for first coordinate in `xnew`.
- `px2` grid values for second coordinate in `xnew`.
- `means` a 20 x 2 matrix of means used in the generation of these data.

**Source**

Flexible Discriminant Analysis

Description

Flexible discriminant analysis.

Usage

```r
fda(formula, data, weights, theta, dimension, eps, method, keep.fitted, ...)
```

Arguments

- **formula**: of the form \( y \sim x \) it describes the response and the predictors. The formula can be more complicated, such as \( y \sim \log(x) + z \) etc (see `formula` for more details). The response should be a factor representing the response variable, or any vector that can be coerced to such (such as a logical variable).
- **data**: data frame containing the variables in the formula (optional).
- **weights**: an optional vector of observation weights.
- **theta**: an optional matrix of class scores, typically with less than \( J-1 \) columns.
- **dimension**: The dimension of the solution, no greater than \( J-1 \), where \( J \) is the number classes. Default is \( J-1 \).
- **eps**: a threshold for small singular values for excluding discriminant variables; default is `.Machine$double.eps`.
- **method**: regression method used in optimal scaling. Default is linear regression via the function `polyreg`, resulting in linear discriminant analysis. Other possibilities are `mars` and `bruto`. For Penalized Discriminant analysis `gen.ridge` is appropriate.
- **keep.fitted**: a logical variable, which determines whether the (sometimes large) component “fitted.values” of the fit component of the returned `fda` object should be kept. The default is `TRUE` if \( n \times \text{dimension} < 5000 \).
- **...**: additional arguments to `method`.

Value

an object of class "fda". Use `predict` to extract discriminant variables, posterior probabilities or predicted class memberships. Other extractor functions are `coef`, `confusion` and `plot`.

The object has the following components:

- **percent.explained**: the percent between-group variance explained by each dimension (relative to the total explained.)
values   optimal scaling regression sum-of-squares for each dimension (see reference). The usual discriminant analysis eigenvalues are given by \( \text{values} / (1-\text{values}) \), which are used to define percent explained.

means   class means in the discriminant space. These are also scaled versions of the final theta’s or class scores, and can be used in a subsequent call to \text{fda} \ ((this only makes sense if some columns of theta are omitted—see the references).

\text{theta.mod} \ (internal) a class scoring matrix which allows predict to work properly.

dimension   dimension of discriminant space.

prior   class proportions for the training data.

\text{fit}   fit object returned by \text{method}.

call   the call that created this object (allowing it to be update-able)

\text{confusion}   confusion matrix when classifying the training data.

The method functions are required to take arguments \( x \) and \( y \) where both can be matrices, and should produce a matrix of fitted values the same size as \( y \). They can take additional arguments \text{weights} and should all have a … for safety sake. Any arguments to \text{method} can be passed on via the … argument of \text{fda}. The default method \text{polyreg} has a degree argument which allows polynomial regression of the required total degree. See the documentation for \text{predict.fda} for further requirements of \text{method}. The package \text{earth} is suggested for this package as well; \text{earth} is a more detailed implementation of the mars model, and works as a \text{method} argument.

Author(s)

Trevor Hastie and Robert Tibshirani

References

“Flexible Discriminant Analysis by Optimal Scoring” by Hastie, Tibshirani and Buja, 1994, JASA, 1255-1270.


“Elements of Statistical Learning - Data Mining, Inference and Prediction” (2nd edition, Chapter 12) by Hastie, Tibshirani and Friedman, 2009, Springer

See Also

\text{predict.fda}, \text{plot.fda}, \text{mars}, \text{bruto}, \text{polyreg}, \text{softmax}, \text{confusion},

Examples

data(iris)
irisfit <- \text{fda}(Species ~ ., data = iris)
irisfit
## \text{fda}(formula = Species ~ ., data = iris)
##
## Dimension: 2
##
## Percent Between-Group Variance Explained:
### gen.ridge

#### Penalized Regression

**Description**

Perform a penalized regression, as used in penalized discriminant analysis.

**Usage**

```r
gen.ridge(x, y, weights, lambda=1, omega, df, ...)
```

**Arguments**

- `x, y, weights` the x and y matrix and possibly a weight vector.
- `lambda` the shrinkage penalty coefficient.
- `omega` a penalty object; omega is the eigendecomposition of the penalty matrix, and need not have full rank. By default, standard ridge is used.
df an alternative way to prescribe lambda, using the notion of equivalent degrees of freedom.

... currently not used.

Value

A generalized ridge regression, where the coefficients are penalized according to omega. See the function definition for further details. No functions are provided for producing one-dimensional penalty objects (omega). `laplacian()` creates a two-dimensional penalty object, suitable for (small) images.

See Also

`laplacian`

---

glass Glass Identification Database

Description

The `glass` data frame has 214 observations and 10 variables, representing glass fragments.

Usage

data(glass)

Format

This data frame contains the following columns:

- **RI** refractive index
- **Na** weight percent in corresponding oxide
- **Mg** weight percent in corresponding oxide
- **Al** weight percent in corresponding oxide
- **Si** weight percent in corresponding oxide
- **K** weight percent in corresponding oxide
- **Ca** weight percent in corresponding oxide
- **Ba** weight percent in corresponding oxide
- **Fe** weight percent in corresponding oxide

**Type** Type of glass:

1. building\_windows\_float\_processed,
2. building\_windows\_non\_float\_processed,
3. vehicle\_windows\_float\_processed,
4. vehicle\_windows\_non\_float\_processed (none in this database),
5. containers,
6. tableware,
7. headlamps
laplacian

create penalty object for two-dimensional smoothing.

Description

Creates a penalty matrix for use by `gen.ridge` for two-dimensional smoothing.

Usage

```
laplacian(size, compose)
laplacian(size = 16, compose = FALSE)
```

Arguments

- `size` dimension of the image is `size x size`; default is 16.
- `compose` default is `compose=FALSE`, which means the penalty is returned as an eigen-decomposition. If `compose=TRUE`, a penalty matrix is returned.

Details

Formulas are used to construct a laplacian for smoothing a square image.

Value

If `compose=FALSE`, an eigen-decomposition object is returned. The vectors component is a `size^2 x size^2` orthogonal matrix, and the $values$ component is a `size^2` vector of non-negative eigen-values. If `compose=TRUE`, these are multiplied together to form a single matrix.

Author(s)

Trevor Hastie <hastie@stanford.edu>

References

Here we follow very closely the material on page 635 in JASA 1991 of O’Sullivan’s article on discretized Laplacian Smoothing

See Also

`gen.ridge`, `fda`
Description

Multivariate adaptive regression splines.

Usage

\texttt{mars(x, y, w, wp, degree, nk, penalty, thresh, prune, trace.mars,}
\texttt{forward.step, prevfit, ...)}

Arguments

- \texttt{x} a matrix containing the independent variables.
- \texttt{y} a vector containing the response variable, or in the case of multiple responses, a matrix whose columns are the response values for each variable.
- \texttt{w} an optional vector of observation weights (currently ignored).
- \texttt{wp} an optional vector of response weights.
- \texttt{degree} an optional integer specifying maximum interaction degree (default is 1).
- \texttt{nk} an optional integer specifying the maximum number of model terms.
- \texttt{penalty} an optional value specifying the cost per degree of freedom charge (default is 2).
- \texttt{thresh} an optional value specifying forward stepwise stopping threshold (default is 0.001).
- \texttt{prune} an optional logical value specifying whether the model should be pruned in a backward stepwise fashion (default is \texttt{true}).
- \texttt{trace.mars} an optional logical value specifying whether info should be printed along the way (default is \texttt{false}).
- \texttt{forward.step} an optional logical value specifying whether forward stepwise process should be carried out (default is \texttt{true}).
- \texttt{prevfit} optional data structure from previous fit. To see the effect of changing the penalty parameter, one can use prevfit with \texttt{forward.step = FALSE}.
- \texttt{...} further arguments to be passed to or from methods.

Value

An object of class \texttt{"mars"}, which is a list with the following components:

- \texttt{call} call used to \texttt{mars}.
- \texttt{all.terms} term numbers in full model. 1 is the constant term. Remaining terms are in pairs (2 3, 4 5, and so on). \texttt{all.terms} indicates nonsingular set of terms.
- \texttt{selected.terms} term numbers in selected model.
- \texttt{penalty} the input penalty value.
degree the input degree value.
thresh the input threshold value.
gcv gcv of chosen model.
factor matrix with $ij$-th element equal to 1 if term $i$ has a factor of the form $x_j > c$, equal to $-1$ if term $i$ has a factor of the form $x_j \leq c$, and to 0 if $x_j$ is not in term $i$.
cuts matrix with $ij$-th element equal to the cut point $c$ for variable $j$ in term $i$.
residuals residuals from fit.
fitted fitted values from fit.
lenb length of full model.
coefficients least squares coefficients for final model.
x a matrix of basis functions obtained from the input x matrix.

Note
This function was coded from scratch, and did not use any of Friedman’s mars code. It gives quite similar results to Friedman’s program in our tests, but not exactly the same results. We have not implemented Friedman’s anova decomposition nor are categorical predictors handled properly yet. Our version does handle multiple response variables, however.

Author(s)
Trevor Hastie and Robert Tibshirani

References

See Also
predict.mars, model.matrix.mars.

Package earth also provides multivariate adaptive regression spline models based on the Hastie/Tibshirani mars code in package mda, adding some extra features. It can be used in the method argument of fda or mda.

Examples

data(trees)
fit1 <- mars(trees[, -3], trees[3])
showcuts <- function(obj)
{
  tmp <- obj$cuts[obj$sel, ]
  dimnames(tmp) <- list(NULL, names(trees)[-3])
  tmp
}
showcuts(fit1)
## Description

Mixture discriminant analysis.

## Usage

```r
mda(formula, data, subclasses, sub.df, tot.df, dimension, eps, iter, weights, method, keep.fitted, trace, ...)```

## Arguments

- `formula`: of the form `y~x` it describes the response and the predictors. The formula can be more complicated, such as `y~log(x)+z` etc (see `formula` for more details). The response should be a factor representing the response variable, or any vector that can be coerced to such (such as a logical variable).
- `data`: data frame containing the variables in the formula (optional).
- `subclasses`: Number of subclasses per class, default is 3. Can be a vector with a number for each class.
- `sub.df`: If subclass centroid shrinking is performed, what is the effective degrees of freedom of the centroids per class. Can be a scalar, in which case the same number is used for each class, else a vector.
- `tot.df`: The total df for all the centroids can be specified rather than separately per class.
- `dimension`: The dimension of the reduced model. If we know our final model will be confined to a discriminant subspace (of the subclass centroids), we can specify this in advance and have the EM algorithm operate in this subspace.
- `eps`: A numerical threshold for automatically truncating the dimension.
- `iter`: A limit on the total number of iterations, default is 5.
- `weights`: `NOT` observation weights! This is a special weight structure, which for each class assigns a weight (prior probability) to each of the observations in that class of belonging to one of the subclasses. The default is provided by a call to `mda.start(x, g, subclasses, trace, ...)` (by this time x and g are
known). See the help for `mda.start`. Arguments for `mda.start` can be provided via the ... argument to `mda`, and the `weights` argument need never be accessed. A previously fit `mda` object can be supplied, in which case the final subclass responsibility weights are used for `weights`. This allows the iterations from a previous fit to be continued.

**method**

Regression method used in optimal scaling. Default is linear regression via the function `polyreg`, resulting in the usual mixture model. Other possibilities are `mars` and `bruto`. For penalized mixture discriminant models `genridge` is appropriate.

**keep.fitted**

A logical variable, which determines whether the (sometimes large) component "fitted.values" of the fit component of the returned `mda` object should be kept. The default is `TRUE` if `n * dimension < 5000`.

**trace**

If `TRUE`, iteration information is printed. Note that the deviance reported is for the posterior class likelihood, and not the full likelihood, which is used to drive the EM algorithm under `mda`. In general the latter is not available.

... additional arguments to `mda.start` and to `method`.

**Value**

An object of class c("mda", "fda"). The most useful extractor is `predict`, which can make many types of predictions from this object. It can also be plotted, and any functions useful for `fda` objects will work here too, such as `confusion` and `coef`.

The object has the following components:

- **percent.explained**
  
  The percent between-group variance explained by each dimension (relative to the total explained.)

- **values**
  
  Optimal scaling regression sum-of-squares for each dimension (see reference).

- **means**
  
  Subclass means in the discriminant space. These are also scaled versions of the final theta's or class scores, and can be used in a subsequent call to `mda` (this only makes sense if some columns of theta are omitted—see the references)

- **theta.mod**
  
  (Internal) a class scoring matrix which allows `predict` to work properly.

- **dimension**
  
  Dimension of discriminant space.

- **sub.prior**
  
  Subclass membership priors, computed in the fit. No effort is currently spent in trying to keep these above a threshold.

- **prior**
  
  Class proportions for the training data.

- **fit**
  
  Fit object returned by `method`.

- **call**
  
  The call that created this object (allowing it to be update-able).

- **confusion**
  
  Confusion matrix when classifying the training data.

- **weights**
  
  These are the subclass membership probabilities for each member of the training set; see the `weights` argument.

- **assign.theta**
  
  A pointer list which identifies which elements of certain lists belong to individual classes.
The multinomial log-likelihood of the fit. Even though the full log-likelihood drives the iterations, we cannot in general compute it because of the flexibility of the method used. The deviance can increase with the iterations, but generally does not.

The method functions are required to take arguments x and y where both can be matrices, and should produce a matrix of fitted.values the same size as y. They can take additional arguments weights and should all have a ... for safety sake. Any arguments to method() can be passed on via the ... argument of mda. The default method polyreg has a degree argument which allows polynomial regression of the required total degree. See the documentation for predict.fda for further requirements of method. The package earth is suggested for this package as well; earth is a more detailed implementation of the mars model, and works as a method argument.

The function mda.start creates the starting weights; it takes additional arguments which can be passed in via the ... argument to mda. See the documentation for mda.start.

Author(s)
Trevor Hastie and Robert Tibshirani

References
“Flexible Discriminant Analysis by Optimal Scoring” by Hastie, Tibshirani and Buja, 1994, JASA, 1255-1270.
“Elements of Statistical Learning - Data Mining, Inference and Prediction” (2nd edition, Chapter 12) by Hastie, Tibshirani and Friedman, 2009, Springer

See Also
predict.mda, mars, bruto, polyreg, gen.ridge, softmax, confusion

Examples
data(iris)
irisfit <- mda(Species ~ ., data = iris)
irisfit
## Call:
## mda(formula = Species ~ ., data = iris)
## ## Dimension: 4
## ## Percent Between-Group Variance Explained:
## v1 v2 v3 v4
## 96.02 98.55 99.90 100.00
## ## Degrees of Freedom (per dimension): 5
## ## Training Misclassification Error: 0.02 ( N = 150 )
### mda.start

#### Initialization for Mixture Discriminant Analysis

**Description**

Provide starting weights for the mda function which performs discriminant analysis by gaussian mixtures.

**Usage**

```r
mda.start(x, g, subclasses = 3, trace.mda.start = FALSE, 
          start.method = c("kmeans", "lvq"), tries = 5, 
          criterion = c("misclassification", "deviance"), ...)
```

**Arguments**

- `x` The x data, or an mda object.
- `g` The response vector g.
- `subclasses` number of subclasses per class, as in mda.
- `trace.mda.start` Show results of each iteration.
- `start.method` Either "kmeans" or "lvq". The latter requires package class (from the VR package bundle).
- `tries` Number of random starts.
- `criterion` By default, classification errors on the training data. Posterior deviance is also an option.
- `...` arguments to be passed to the mda fitter when using posterior deviance.
Value

A list of weight matrices, one for each class.

---

**model.matrix.mars**  
*Produce a Design Matrix from a ‘mars’ Object*

Description

Produce a design matrix from a ‘mars’ object.

Usage

```r
## S3 method for class 'mars'
model.matrix(object, x, which, full = FALSE, ...)
```

Arguments

- `object`: a mars object.
- `x`: optional argument; if supplied, the mars basis functions are evaluated at these new observations.
- `which`: which columns should be used. The default is to use the columns described by the component `selected.terms` on `object`.
- `full`: if TRUE the entire set of columns are selected, even redundant ones. This is used for updating a mars fit.
- `...`: further arguments to be passed from or to methods.

Value

A model matrix corresponding to the selected columns.

See Also

`mars`, `predict.mars`
mspline

Vector Smoothing Spline

Description

Fit a smoothing spline to a matrix of responses, single x.

Usage

mspline(x, y, w, df = 5, lambda, thresh = 1e-04, ...)

Arguments

x  x variable (numeric vector).
y  response matrix.
w  optional weight vector, defaults to a vector of ones.
df  requested degrees of freedom, as in smooth.spline.
lambda  can provide penalty instead of df.
thresh  convergence threshold for df inversion (to lambda).
...  holdall for other arguments.

Details

This function is based on the ingredients of smooth.spline, and allows for simultaneous smoothing of multiple responses

Value

A list is returned, with a number of components, only some of which are of interest. These are

lambda  The value of lambda used (in case df was supplied)
df  The df used (in case lambda was supplied)
s  A matrix like y of smoothed responses
lev  Self influences (diagonal of smoother matrix)

Author(s)

Trevor Hastie

See Also

smooth.spline

Examples

x=rnorm(100)
y=matrix(rnorm(100*10),100,10)
fit=mspline(x,y,df=5)
package: plot.fda

Plot for Flexible Discriminant Analysis

Description

Plot in discriminant (canonical) coordinates a fda or (by inheritance) a mda object.

Usage

## S3 method for class 'fda'
plot(x, data, coords, group, colors, pch, mcolors, mpch, pcex, mcex, ...)

Arguments

x an object of class "fda".

data the data to plot in the discriminant coordinates. If group="true", then data should be a data frame with the same variables that were used in the fit. If group="predicted", data need not contain the response variable, and can in fact be the correctly-sized "x" matrix.

coords vector of coordinates to plot, with default coords=c(1,2). All pairs of plots are produced.

group if group="true" (the default), each point is color and symbol coded according to the response in data. If group="predicted", the class of each point is predicted from the model, and used instead.

colors a vector of colors to be used in the plotting.

pch a vector of plotting characters.

mcolors a vector of colors for the class centroids; default is colors.

mpch a vector of plotting characters for the centroids.

pcex character expansion factor for the points; default is pcex="0.5".

mcex character expansion factor for the centroids; default is pcex="2.5".

... further arguments to be passed to or from methods.

See Also

fda, mda, predict.fda

Examples

data(iris)
irisfit <- fda(Species ~ ., data = iris)
plot(irisfit)
data(ESL.mixture)
## Not a data frame
mixture.train=ESL.mixture[,c("x","y")]
mixfit=mda(y~x, data=mixture.train)
plot(mixfit, mixture.train)
plot(mixfit, data=ESL.mixture$xnew, group="pred")
**polyreg**

---

**Polynomial Regression**

**Description**

Simple minded polynomial regression.

**Usage**

polyreg(x, y, w, degree = 1, monomial = FALSE, ...)

**Arguments**

- **x**: predictor matrix.
- **y**: response matrix.
- **w**: optional (positive) weights.
- **degree**: total degree of polynomial basis (default is 1).
- **monomial**: If TRUE a monomial basis is used (no cross terms). Default is FALSE.
- **...**: currently not used.

**Value**

A polynomial regression fit, containing the essential ingredients for its predict method.

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**predict.bruto**

---

**Predict method for BRUTO Objects**

**Description**

Predicted values based on ‘bruto’ additive spline models which are fit by adaptive backfitting.

**Usage**

## S3 method for class ‘bruto’
predict(object, newdata, type=c("fitted", "terms"), ...)

**Arguments**

- **object**: a fitted bruto object
- **newdata**: values at which predictions are to be made.
- **type**: if type is "fitted", the fitted values are returned. If type is "terms", a list of fitted terms is returned, each with an x and y component. These can be used to show the fitted functions.
- **...**: further arguments to be passed to or from methods.
Value

Either a fit matrix or a list of fitted terms.

See Also

bruto, predict

Examples

data(trees)
fit1 <- bruto(trees[,3], trees[3])
fitted.terms <- predict(fit1, as.matrix(trees[,3]), type = "terms")
par(mfrow=c(1,2), pty="s")
for(tt in fitted.terms) plot(tt, type="l")

predict.fda  

Classify by Flexible Discriminant Analysis

Description

Classify observations in conjunction with fda.

Usage

## S3 method for class 'fda'
predict(object, newdata, type, prior, dimension, ...)

Arguments

object  
an object of class "fda".
newdata  
new data at which to make predictions. If missing, the training data is used.
type  
kind of predictions: type = "class" (default) produces a fitted factor, type = "variates" produces a matrix of discriminant (canonical) variables, type = "posterior" produces a matrix of posterior probabilities (based on a gaussian assumption), and type = "hierarchical" produces the predicted class in sequence for models of all dimensions.
prior  
the prior probability vector for each class; the default is the training sample proportions.
dimension  
the dimension of the space to be used, no larger than the dimension component of object.
...  
further arguments to be passed to or from methods.

Value

An appropriate object depending on type. object has a component fit which is regression fit produced by the method argument to fda. There should be a predict method for this object which is invoked. This method should itself take as input object and optionally newdata.
predict.mars

See Also

fda, mars, bruto, polyreg, softmax, confusion

Examples

data(iris)
irisfit <- fda(Species ~ ., data = iris)
irisfit
## Call:
## fda(x = iris$x, g = iris$g)
## ## Dimension: 2
## ## Percent Between-Group Variance Explained:
## ## v1 v2
## ## 99.12 100
confusion(predict(irisfit, iris), iris$Species)
## ## Setosa Versicolor Virginica
## ## Versicolor 0 48 1
## ## Virginica 0 2 49
## attr(, "error"): 
## [1] 0.02

predict.mars

Predict method for MARS Objects

Description

Predicted values based on ‘mars’ multivariate adaptive regression spline models.

Usage

## S3 method for class 'mars'
predict(object, newdata, ...)

Arguments

object an object of class "mars".
newdata values at which predictions are to be made.
... further arguments to be passed to or from methods.

Value

the fitted values.

See Also

mars, predict, model.matrix.mars
Classify by Mixture Discriminant Analysis

Description

Classify observations in conjunction with mda.

Usage

```r
## S3 method for class 'mda'
predict(object, newdata, type, prior, dimension, g, ...)
```

Arguments

- `object`: a fitted mda object.
- `newdata`: new data at which to make predictions. If missing, the training data is used.
- `type`: kind of predictions: `type = "class"` (default) produces a fitted factor, `type = "variates"` produces a matrix of discriminant variables (note that the maximal dimension is determined by the number of subclasses), `type = "posterior"` produces a matrix of posterior probabilities (based on a gaussian assumption), `type = "hierarchical"` produces the predicted class in sequence for models of dimensions specified by `dimension` argument.
- `prior`: the prior probability vector for each class; the default is the training sample proportions.
- `dimension`: the dimension of the space to be used, no larger than the dimension component of `object`, and in general less than the number of subclasses. `dimension` can be a vector for use with `type = "hierarchical"`.
- `g`: `???
- `...`: further arguments to be passed to or from methods.

Value

An appropriate object depending on `type`. `object` has a component `fit` which is regression fit produced by the method argument to `mda`. There should be a `predict` method for this object which is invoked. This method should itself take as input `object` and optionally `newdata`.

See Also

`mda`, `fda`, `mars`, `bruto`, `polyreg`, `softmax`, `confusion`
softmax

Examples

data(glass)
samp <- sample(1:nrow(glass), 100)
glass.train <- glass[samp,]
glass.test <- glass[-samp,]
glass.mda <- mda(Type ~ ., data = glass.train)
predict(glass.mda, glass.test, type = "post") # abbreviations are allowed
confusion(glass.mda, glass.test)

softmax

Find the Maximum in Each Row of a Matrix

Description

Find the maximum in each row of a matrix.

Usage

softmax(x, gap = FALSE)

Arguments

x

a numeric matrix.

gap

if TRUE, the difference between the largest and next largest column is returned.

Value

A factor with levels the column labels of x and values the columns corresponding to the maximum column. If gap = TRUE a list is returned, the second component of which is the difference between the largest and next largest column of x.

See Also

predict.fda, confusion, fda.mda

Examples

data(iris)
irisfit <- fda(Species ~ ., data = iris)
posterior <- predict(irisfit, type = "post")
confusion(softmax(posterior), iris[, "Species"])
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