Package ‘SVMMaj’

February 26, 2018

Type Package

Title Implementation of the SVM-Maj Algorithm

Version 0.2-8

Date 2018-01-26

Author Hoksan Yip, Patrick J.F. Groenen, Georgi Nalbantov

Maintainer Hok San Yip <hoksan@gmail.com>

Description Implements the SVM-Maj algorithm to train data with support vector machine as described in Groenen et al. (2008) <doi:10.1007/s11634-008-0020-9>. This algorithm uses two efficient updates, one for linear kernel and one for the nonlinear kernel.

Imports reshape2, scales, gridExtra, dplyr, ggplot2, kernlab

Depends R (>= 2.13.0), stats, graphics

Suggests utils, testthat, magrittr, xtable

License GPL-2

LazyData Yes

RoxygenNote 6.0.1

VignetteBuilder utils

NeedsCompilation no

Repository CRAN

Date/Publication 2018-02-26 00:01:09 UTC

R topics documented:

     auc   AusCredit  classification  diabetes  getHinge  isb  isplinebasis
             2            3               4            4       5           6       7
**Description**

Returns the area under the curve value as a fraction.

**Usage**

```
auc(q, y = attr(q, "y"))
```

**Arguments**

- `q` the predicted values
- `y` a list of the actual classes of `q`

**Value**

the area under the curve value

**Examples**

```r
df <- with(diabetes, cbind(y, X))
lm.y <- glm(y ~ ., data = df, family = binomial())
print(with(lm.y, auc(fitted.values, y)))
```
Description

This file concerns credit card applications of 690 households.

Format

This data set has been split into two components for the convenience of the model training.

data.frame-object \( X \) consists of with 6 numerical and 8 categorical attributes. The labels have been changed for the convenience of the statistical algorithms. For example, attribute 4 originally had 3 labels p,g,gg and these have been changed to labels 1,2,3.

Factor \( y \) indicates whether the application has been Accepted or Rejected.

The training set \( \text{AusCredit.tr} \) contains a randomly selected set of 400 subjects, and \( \text{AusCredit.te} \) contains the remaining 290 subjects. \( \text{AusCredit} \) contains all 690 objects.

Details

All attribute names and values have been changed to meaningless symbols to protect confidentiality of the data.

This dataset is interesting because there is a good mix of attributes – continuous, nominal with small numbers of values, and nominal with larger numbers of values. There are also a few missing values.

Source


Examples

\begin{verbatim}
attach(AusCredit)
summary(X)
summary(y)
detach(AusCredit)
\end{verbatim}
**classification**  
*Show the classification performance*

**Description**

Given the predicted value \( q \) and the observed classes \( y \), it shows an overview of the prediction performances with hit rates, misclassification rates, true positives (TP), false positives (FP) and precision.

**Usage**

```r
classification(q, y, classes = c("-1", "1"), weights = NULL)
```

**Arguments**

- \( q \): the predicted values
- \( y \): a list of the actual classes of \( q \)
- \( classes \): a character vector with the labels of the two classes
- \( weights \): an optional parameter to specify a weighted hit rate and misclassification rate

**Value**

a list with three elements, \( \text{matrix} \) equals the confusion matrix, \( \text{overall} \) equals the overall prediction performance and \( \text{measures} \) the measures per class is stored.

---

**diabetes**  
*Pima Indians Diabetes Data Set*

**Description**

From National Institute of Diabetes and Digestive and Kidney Diseases.

**Format**

\( X \) is a data frame of 768 female patients with 8 attributes.

- no.pregnant: number of pregnancies.
- glucose: plasma glucose concentration in an oral glucose tolerance test
- blood.press: diastolic blood pressure (mm Hg)
- triceps.thick: triceps skin fold thickness (mm)
- insulin: 2-Hour serum insulin (mu U/ml)
- BMI: body mass index (weight in kg/(height in m)^2)
- pedigree: diabetes pedigree function
- age: age in years
getHinge

y contains the class labels: Yes or No, for diabetic according to WHO criteria.
The training set diabetes.tr contains a randomly selected set of 600 subjects, and diabetes.te contains the remaining 168 subjects. diabetes contains all 768 objects.

Details
Several constraints were placed on the selection of these instances from a larger database. In particular, all patients here are females at least 21 years old of Pima Indian heritage.

Source

References

Examples

attach(diabetes)
summary(x)
summary(y)

gethinge

Hinge error function of SVM-Maj

Description
This function creates a function to compute the hinge error, given its predicted value q and its class y, according to the loss term of the Support Vector machine loss function.

Usage

getHinge(hinge = "quadratic", delta = 3, eps = 1e-08)

Arguments

hinge  Hinge error function to be used, possible values are 'absolute', 'quadratic' and 'huber'
delta  The parameter of the huber hinge (only if hinge = 'huber').
eps  Specifies the maximum steepness of the quadratic majorization function
      m(q) = a * q ^ 2 - 2 * b * q + c, where a <= .25 * eps ^ -1.
Value

The hinge error function with arguments q and y to compute the hinge error. The function returns a list with the parameters of the majorization function SVM-Maj (a, b and c) and the loss error of each object (loss).

References


See Also

svmmaj

Examples

hingefunction <- getHinge()
## plot hinge function value and, if specified, 
## the majorization function at z
## plot(hingefunction, z = 3)
## generate loss function value
loss <- hingefunction(q = -10:10, y = 1)$loss
print(loss)
plot(hingefunction, z = 3)

---

**ish**

*I-spline basis of each column of a given matrix*

Description

Create a I-spline basis for an array. isb will equally distribute the knots over the value range using quantiles.

Usage

ish(x, spline.knots = 0, knots = NULL, spline.degree = 1)

Arguments

- **x**  The predictor variable, which will be transformed into I-spline basis.
- **spline.knots**  Number of inner knots to use. isb will equally distribute the knots over the value range using quantiles. spline.knots will only be used if knots is not given.
- **knots**  An array consisting all knots (boundary knots as well as the interior knots) to be used to create the spline basis.
- **spline.degree**  The polynomial degree of the spline basis.
Value

The I-spline with the used spline settings as attribute. The spline settings attribute can transform the same attribute of any other objects using the same knots.

Author(s)

Hok San Yip, Patrick J.F. Groenen, Georgi Nalbantov

References


See Also

svmmaj

Examples

```r
## plot the spline transformation given a monotone sequence
B0 <- isb(0:100, spline.knots = 2, spline.degree = 3)
plot(NULL, xlim = c(0, 140), ylim = c(0, 1), xlab = 'x', ylab = 'I-spline')
for(i in 1:ncol(B0))
  lines(B0[,i], col = i, lwd = 3)
legend('bottomright', legend = 1:ncol(B0), col = 1:ncol(B0),
       lty = 1, lwd = 3, title = 'Spline Columns')

## create I-spline basis for the first 50 observations
x <- iris$Sepal.Length
B1 <- isb(x[1:50], spline.knots = 4, spline.degree = 3)

## extracting the spline transformation settings
spline.param <- attr(B1, 'splineInterval')

## use the same settings to apply to the next 50 observations
B2 <- isb(x[-(1:50)], spline.degree = 3, knots = spline.param)
```

```r

isplinebasis(x, knots, d)

```

Description

Inner function call to create I-splines based on the user defined knots and polynomial degree d of the splines

Usage

```r
isplinebasis(x, knots, d)
```
Arguments

- **x**  
  a scalar or vector of values which will be transformed into splines
- **knots**  
  a vector of knot values of the splines
- **d**  
  the polynomial degree of the splines

Value

a matrix with for each value of x the corresponding spline values.

---

normalize  
Normalize/standardize the columns of a matrix

Description

Standardize the columns of an attribute matrix x to zscores, to the range [0, 1] or a prespecified scale.

Usage

```r
normalize(x, standardize = "zscore")
```

Arguments

- **x**  
  An attribute variable which will be scaled.
- **standardize**  
  Either a string value denoting a predefined scaling, or a list with values `a` and `b` corresponding with the numeric centering and scaling, that is, using the function `x * standardize$b - standardize$a`.

Value

The standardized matrix. The numeric centering and scalings used are returned as attribute "standardize".

Author(s)

Hok San Yip, Patrick J.F. Groenen, Georgi Nalbantov

References


See Also

`svmmaj`
plot.hinge

Examples

```r
## # standardize the first 50 objects to zscores
x <- iris$Sepal.Length
x1 <- normalize(x[1:50], standardize = 'zscore')
## # use the same settings to apply to the next 100 observations
x2 <- normalize(x[-(1:50)], standardize = attr(x1, 'standardization'))
```

Description

This function plots the hinge object created by `getHinge`.

Usage

```r
## S3 method for class 'hinge'
plot(x, y = 1, z = NULL, ...)
```

Arguments

- `x`: The hinge object returned from `getHinge`.
- `y`: Specifies the class (-1 or 1) to be plotted for the hinge error.
- `z`: If specified, the majorization function with the supporting point z will also be plotted.
- `...`: Other arguments passed to plot method.

Examples

```r
hingefunction <- getHinge()
## plot hinge function value
plot(hingefunction, z = 3)
```

plot.svmmajcrossval

Plot the cross validation output

Description

Shows the results of the cross validation graphically. Possible graphics are among others the distribution of the predicted values per class per lambda value and the misclassification rate per lambda.
Usage

```r
## S3 method for class 'svmmajcrossval'
plot(x, type = "grid", ...)
```

Arguments

- `x`: the `svmmajcrossval` object
- `type`: the type of graph being shown, possible values are 'grid' for the missclassification rate per lambda value, 'profile' the distribution of predicted values of the classes per lambda value
- `...`: Further arguments passed to or from other methods.

**plotWeights**

*Plot the weights of all attributes from the trained SVM model*

Description

Shows, one graph per attribute, the weights of all attributes. The type of graph depends on the type of the attribute: the spline line of the corresponding attribute in case a spline has been used, a bar plot for categorical and logical values, and a linear line for all other type of the attribute values. This function cannot be used in a model with a non-linear kernel.

Usage

```r
plotWeights(object, plotdim = c(3, 3), ...)
```

Arguments

- `object`: The model returned from `svmmaj`.
- `plotdim`: A vector of the form `c(nr, nc)`. Subsequent figures will be drawn in an `nr`-by-`nc` array on the device.
- `...`: other parameters given to the `plot` function

**predict.svmmaj**

*Out-of-Sample Prediction from Unseen Data.*

Description

This function predicts the predicted value (including intercept), given a previous trained model which has been returned by `svmmaj`. 
**Usage**

```r
## S3 method for class 'svmmaj'
predict(object, X.new, y = NULL, weights = NULL,
       show.plot = FALSE, ...)
```

**Arguments**

- `object`: Model which has been trained beforehand using `svmmaj`.
- `X.new`: Attribute matrix of the objects to be predicted, which has the same number of attributes as the untransformed attribute matrix in `model`.
- `y`: The actual class labels (only if `show.plot`==TRUE).
- `weights`: The weight of observation as the relative importance of the prediction error of the observation.
- `show.plot`: If `show.plot`=TRUE, it plots the density of the predicted value for both class labels, if `y` is not specified, the density of all objects will be plotted.
- `...`: Arguments to be passed to methods.

**Value**

The predicted value (including intercept) of class q.svmmaj, with attributes:

- `y`: The observed class labels of each object.
- `yhat`: The predicted class labels of each object.
- `classes`: The class labels.

**Author(s)**

Hok San Yip, Patrick J.F. Groenen, Georgi Nalbantov

**References**


**See Also**

- `svmmaj`

**Examples**

```r
attach(AusCredit)

## model training
model <- svmmaj(X[1:400, ], y[1:400], hinge = 'quadratic', lambda = 1)
## model prediction
q4 <- predict(model, X[-(1:400), ], y[-(1:400)], show.plot = TRUE)
q4
```
predict.transDat  *Perform the transformation based on predefined settings*

**Description**

Given the input parameters, which are generated from `transformdata`, it performs the same transformation with the same settings to the given input.

**Usage**

```r
## S3 method for class 'transDat'
predict(x, attrib = NULL, values = NULL,
    standardization = NULL, splineInterval = NULL, splineDegree = NULL)
```

**Arguments**

- `x`  
  a (new) vector of numerics to be transformed
- `attrib`  
  either a list of settings, or NULL in case the attributes are given as separate input
- `values`  
  a vector of levels in case `x` is a factor
- `standardization`  
  the standardization rules from `normalize`
- `splineInterval`  
  the knots to be used for spline basis
- `splineDegree`  
  the polynomial degree of the splines

**Value**

a transformed data based on the user defined settings

---

**print.q.svmmaj**  *SVM-Maj Algorithm*

**Description**

SVM-Maj is an algorithm to compute a support vector machine (SVM) solution. In its most simple form, it aims at finding hyperplane that optimally separates two given classes. This objective is equivalent to finding a linear combination of `k` predictor variables to predict the two classes for `n` observations. SVM-Maj minimizes the standard support vector machine (SVM) loss function. The algorithm uses three efficient updates for three different situations: primal method which is efficient in the case of `n > k`, the decomposition method, used when the matrix of predictor variables is not of full rank, and a dual method, that is efficient when `n < k`. Apart from the standard absolute hinge error, SVM-Maj can also handle the quadratic and the Huber hinge.
print.q.svmmaj

Usage

## S3 method for class 'q.svmmaj'
print(x, ...)

svmmaj(x, y, lambda = 1, weights.obs = 1, weights.var = 1,
        scale = c("interval", "zscore", "none"), spline.knots = 0,
        spline.degree = 1L, kernel = vanilladot, kernel.sigma = 1,
        kernel.scale = 1, kernel.degree = 1, kernel.offset = 1,
        hinge = c("absolute", "quadratic", "huber", "logitistic"),
        hinge.delta = 1e-08, options = setSVMoptions(), initial.point = NULL,
        verbose = FALSE, na.action = na.omit, ...)

## Default S3 method:
svmmaj(x, y, lambda = 1, weights.obs = 1,
        weights.var = 1, scale = c("interval", "zscore", "none"),
        spline.knots = 0, spline.degree = 1L, kernel = vanilladot,
        kernel.sigma = 1, kernel.scale = 1, kernel.degree = 1,
        kernel.offset = 1, hinge = c("absolute", "quadratic", "huber",
        "logitistic"), hinge.delta = 1e-08, options = setSVMoptions(),
        initial.point = NULL, verbose = FALSE, na.action = na.omit, ...)

Arguments

- **x**
  - the svmmaj object as result of `svmmaj`

- **X**
  - A data frame (or object coercible by `as.data.frame` to a data frame) consisting
    the attributes, the class of each attribute can be either numeric, logical or
    factor.

- **y**
  - A factor (or object coercible by `factor` to a factor) consisting the class labels.

- **lambda**
  - Regularization parameter of the penalty term.

- **weights.obs**
  - a vector of length \(n\) with the nonnegative weight for the residual of each object
    (with length \(n\)). If the length is 2, then it specifies the weight per class.

- **weights.var**
  - a vector of length \(k\) with weights for each attribute.

- **scale**
  - Specifies whether the columns of attribute matrix \(X\) needs to be standardized
    into zscores or to the interval \([0, 1]\). Possible values are: none, zscore and
    interval. Moreover, the standardization parameters can be given instead.

- **spline.knots**
  - equals the number of internal knots of the spline basis. When the number
    of knots exceeds the number of (categorical) values of an explanatory vari-
    able, the duplicate knots will be removed using `unique`. For no splines, use
    `spline.knots = 0`.

- **spline.degree**
  - equals the polynomial degree of the splines, for no splines: `spline.degree = 1`.

- **kernel**
  - Specifies which kernel function to be used (see `dots` of package `kernlab`). De-
    fault kernel is the linear kernel.

- **kernel.sigma**
  - additional parameters used for the kernel function (see `dots`)

- **kernel.scale**
  - additional parameters used for the kernel function (see `dots`)
kernel.degree  additional parameters used for the kernel function (see dots)
kernel.offset  additional parameters used for the kernel function (see dots)
hinge  Specifies with hinge function from getHinge should be used.
hinge.delta  The parameter of the huber hinge (only if hinge = 'huber').
options  additional settings used in the svmmaj algorithm
initial.point  Initial solution.
verbose  TRUE shows the progress of the iteration.
na.action  Generic function for handling NA values.

Details

The following settings can be added as element in the options parameter: decomposition Specifies whether the QR decomposition should be used for efficient updates. Possible values are 'svd' for Singular value decomposition (Eigenvalue decomposition for non-linear kernel) or 'chol' for Cholesky (or QR decomposition in case of linear kernel)
convergence Specifies the convergence criterion of the algorithm. Default is 1e-08. increase.step The iteration number from which relaxed update will be used. eps The relaxation of the majorization function for absolute hinge: .25 * eps^-1 is the maximum steepness of the majorization function.
check.positive Specifies whether a check has to be made for positive input values. max.iter maximum number of iterations to use

Value

Returns a svmmaj-class object, of which the methods plot, plotWeights, summary and predict can be applied. (see also predict.svmmaj and print.svmmaj)

Author(s)

Hok San Yip, Patrick J.F. Groenen, Georgi Nalbantov

References


See Also

dots for the computations of the kernels. predict.svmmaj normalize isb getHinge

Examples

```r
# using default settings
model1 <- svmmaj(
  diabetes$x, diabetes$y, hinge = 'quadratic', lambda = 1)
summary(model1)
```
weights.obs = list(positive = 2, negative = 1)
## using radial basis kernel
library(kernlab)
model2 <- svmmaj(
    diabetes$x, diabetes$y, hinge = 'quadratic', lambda = 1,
    weights.obs = weights.obs, scale = 'interval',
    kernel = rbfdot,
    kernel.sigma = 1
)
summary(model2)
## I-spline basis
library(ggplot2)
model3 <- svmmaj(
    diabetes$x, diabetes$y, weight.obs = weight.obs,
    spline.knots = 3, spline.degree = 2
)
plotWeights(model3, plotdim = c(2, 4))

---

**Description**

Trained SVM model as output from `svmmaj`. The returning object consist of the following values:

- **call** The function specifications which has been called.
- **lambda** The regularization parameter of the penalty term which has been used.
- **loss** The corresponding loss function value of the final solution.
- **iteration** Number of iterations needed to evaluate the algorithm.
- **X** The attribute matrix of $\dim(X) = c(n,k)$.
- **y** The vector of length $n$ with the actual class labels. These labels can be numeric $[0, 1]$ or two strings.
- **classes** A vector of length $n$ with the predicted class labels of each object, derived from $q.tilde$
- **Xtrans** The attribute matrix $X$ after standardization and (if specified) spline transformation.
- **norm.param** The applied normalization parameters (see `normalize`).
- **splineInterval** The spline knots which has been used (see `isb`).
- **splineLength** Denotes the number of spline basis of each explanatory variable in $X$.
- **method** The decomposition matrices used in estimating the model.
- **hinge** The hinge function which has been used (see `getHinge`).
- **beta** If identified, the beta parameters for the linear combination (only available for linear kernel).
- **q** A vector of length $n$ with predicted values of each object including the intercept.
- **nSV** Number of support vectors.
Usage

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

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

## S3 method for class 'summary.svmmaj'
print(x, ...)

## S3 method for class 'svmmaj'
plot(x, ...)

Arguments

x the svmmaj object as result of svmmaj

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

object the svmmaj object as result of svmmaj

---

print.svmmajcrossval  Print SVMaj cross validation results

Description

Prints the result from the cross validation procedure in svmmajcrossval.

Usage

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

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

Arguments

x the cross-validation output from svmmajcrossval

... ignored

object the output object from svmmajcrossval
roccurve

Plot the ROC curve of the predicted values

Description

Given the predicted values q and its corresponding observed classes y, it shows its separation performances by showing the roc-curve.

Usage

roccurve(q, y = attr(q, "y"), class = 1, ...)

Arguments

q the predicted values
y a list of the actual classes of q
class the base class to show the roc-curve
... additional parameters given as input to the plot function

Examples

model <- svm(diabetes$X, diabetes$y)
roccurve(model$q)

Description

Supermarket data 1996

Format

This dataframe contains the following columns

• STORE: Identifier of the store
• CITY: The city of the store
• ZIP: The zip code of the store
• GROCERY_sum
• GROCCOUP_sum
• AGE9
• AGE60
• ETHNIC
• EDUC
• NOCAR
• INCOME
• INCSIGMA
• HSIZEAVG
• HSIZE1
• HSIZE2
• HSIZE34
• HSIZE567
• HH3PLUS
• HH4PLUS
• HHSINGLE
• HHLARGE
• WORKWOM
• SINHOUSE
• DENSITY
• HVAL150
• HVAL200
• HVALMEAN
• SINGLE
• RETIRED
• UNEMP
• WRKCH5
• WRKCH17
• NWRKCH5
• NWRKCH17
• WRKCH
• NWRKCH
• WRKWCH
• WRKWNCH
• TELEPHN
• MORTGAGE
• NWHITE
• POVERTY
• SHPCONS
• SHPHURR
• SHPAVID
• SHPKSTR
• SHPUNFT
• SHPBIRD
• SHOPINDX
• SHPINDX
**Examples**

head(supermarket1996, 3)

---

**svmmajcrossval**

---

### k-fold Cross-Validation of SVM-Maj

**Description**

This function performs a gridsearch of k-fold cross-validations using SVM-Maj and returns the combination of input values which has the best forecasting performance.

**Usage**

```r
svmmajcrossval(x, y, search.grid = list(lambda = 2^seq(-5, 5, length.out = 19)), ..., convergence = 1e-04, weights.obs = 1, check.positive = TRUE, mc.cores = getOption("mc.cores"), options = NULL, verbose = FALSE, ngroup = 5, groups = NULL, return.model = FALSE)
```

**Arguments**

- **x**: A data frame (or object coercible by `as.data.frame` to a data frame) consisting of the attributes.
- **y**: A factor (or object coercible by `factor` to a factor) consisting of the class labels.
- **search.grid**: A list with for each factor the range of values to search for.
- **...**: Other arguments to be passed through `svmmaj`.
- **convergence**: Specifies the convergence criterion for `svmmaj`. Default is `1e-08`.
- **weights.obs**: Weights for the classes.
- **check.positive**: Specifies whether a check should be performed for positive `lambda` and `weights.obs`.
- **mc.cores**: the number of cores to be used (for parallel computing).
- **options**: additional settings used in the `svmmaj` algorithm.
- **verbose**: =TRUE shows the progress of the cross-validation.
- **ngroup**: The number of groups to be divided into.
- **groups**: A predetermined group division for performing the cross validation.
- **return.model**: =TRUE estimates the model with the optimal parameters.

**Value**

- **loss.opt**: The minimum (weighted) missclassification rate found in out-of-sample training along the search grid.
- **param.opt**: The level of the factors which gives the minimum loss term value.
- **loss.grp**: A list of missclassification rates per hold-out sample.
- **groups**: A vector defining the cross-validation groups which has been used.
qhat  The estimated out-of-sample predicted values in the cross-validation.
qhat.in The trained predicted values
param.grid The matrix of all gridpoints which has been performed during the cross-validation, with its corresponding weighted out-of-sample misclassification rate.
model  The svmmaj-object with the estimated model using the optimal parameters found in the cross-validation.

Author(s)

Hok San Yip, Patrick J.F. Groenen, Georgi Nalbantov

References


See Also

svmmaj

Examples

Xt <- diabetes$X
yt <- diabetes$y

## performing gridsearch with k-fold cross-validation
results <- svmmajcrossval(
  Xt, yt,
  scale = 'interval',
  mc.cores = 2,
  ngroup = 5,
  return.model = TRUE
)

summary(results$model)
results
plot(results)
plot(results, 'profile')

---

**transformdata**

*Transform the data with normalization and/or spline basis*

Description

Performs subsequently a normalization of the input data and creating spline basis based on the user defined input.
Usage

transformdata(x, standardize = c("interval", "zscore", "none"),
  spline.knots = 0, spline.degree = 1)

Arguments

x a single column of values as input for the data transformation
standardize Either a string value denoting a predefined scaling, or a list with values a and b corresponding with the numeric centering and scaling, that is, using the function 
  x * standardize$db - standardize$da.
spline.knots Number of inner knots to use. isb will equally distribute the knots over the value range using quantiles. spline.knots will only be used if knots is not given.
spline.degree The polynomial degree of the spline basis.

Value

transformed data in spline basis or (in case of no spline) a normalized vector

voting  

Congressional Voting Records Data Set

Description

1984 United Stated Congressional Voting Records; Classify as Republican or Democrat.

Format

X is a data frame with 434 congress members and 16 attributes: 16 key votes identified by the Congressional Quarterly Almanac (CQA). All attributes are binary values, with 1= yes and 0= no.

X1  handicapped-infants
X2  water-project-cost-sharing
X3  adoption-of-the-budget-resolution
X4  physician-fee-freeze
X5  el-salvador-aid
X6  religious-groups-in-schools
X7  anti-satellite-test-ban
X8  aid-to-nicaraguian-contras
X9  mx-missile
X10 immigration
X11 synfuels-corporation-cutback
X12 education-spending
X13 superfund-right-to-sue
X14 crime
X15 duty-free-exports
X16 export-administration-act-south-africe
y consists factors which denotes whether the congress member is a Republican or a Democrat. The training set voting.tr contains a randomly selected set of 300 subjects, and voting.te contains the remaining 134 subjects. voting contains all 434 objects.

Details

This data set includes votes for each of the U.S. House of Representatives Congressmen on the 16 key votes identified by the CQA. The CQA lists nine different types of votes: voted for, paired for, and announced for (these three simplified to yea), voted against, paired against, and announced against (these three simplified to nay), voted present, voted present to avoid conflict of interest, and did not vote or otherwise make a position known (these three simplified to an unknown disposition).

Source


Examples

attach(voting)
summary(X)
summary(y)

X.svmmaj

Returns transformed attributes

Description

For efficiency use in svmmajcrossval

Usage

X.svmmaj(object, X.new, weights = NULL)

Arguments

object Model which has been trained beforehand using svmmaj.
X.new Attribute matrix of the objects to be predicted, which has the same number of attributes as the untransformed attribute matrix in model.
weights The weight of observation as the relative importance of the prediction error of the observation.
Index

*Topic **datasets**
  - AusCredit, 3
  - diabetes, 4
  - supermarket1996, 17
  - voting, 21

  as.data.frame, 13, 19
  auc, 2
  AusCredit, 3

  classification, 4

  diabetes, 4
  dots, 13, 14

  factor, 13, 19

  getHinge, 5, 14, 15

  isb, 6, 14, 15
  isplinebasis, 7

  normalize, 8, 14, 15

  plot.hinge, 9
  plot.svmmaj(print.svmmaj), 15
  plot.svmmajcrossval, 9
  plotWeights, 10
  predict.svmmaj, 10, 14
  predict.transDat, 12
  print.hinge (getHinge), 5
  print.q.svmmaj, 12
  print.summary.svmmaj (print.svmmaj), 15
  print.svmmaj, 14, 15
  print.svmmajcrossval, 16

  roccurve, 17

  summary.svmmaj (print.svmmaj), 15
  summary.svmmajcrossval
    (print.svmmajcrossval), 16

  supermarket1996, 17
  svmmaj, 6–8, 10, 11, 13, 15, 16, 20, 22
  svmmaj (print.q.svmmaj), 12
  svmmajcrossval, 16, 19

  transformdata, 20

  unique, 13

  voting, 21

  X.svmmaj, 22