Package 'BranchGLM'

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Type Package

Title Efficient Best Subset Selection for GLMs via Branch and Bound Algorithms

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URL https://github.com/JacobSeedorff21/BranchGLM

BugReports https://github.com/JacobSeedorff21/BranchGLM/issues

Description Performs efficient and scalable glm best

subset selection using a novel implementation of a branch and bound algorithm. To speed up the model fitting process, a range of optimization methods are implemented in 'RcppArmadillo'. Parallel computation is available using 'OpenMP'.

License Apache License (>= 2)

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Description

Fits generalized linear models (GLMs) via RcppArmadillo with the ability to perform some computation in parallel with OpenMP.

```
BranchGLM(
  formula,
  data,
  family,
  link,
  offset = NULL,
```

```
method = "Fisher",
 grads = 10,
 parallel = FALSE,
 nthreads = 8,
  tol = 1e-06,
 maxit = NULL,
 init = NULL,
 fit = TRUE,
 contrasts = NULL,
 keepData = TRUE,
 keepY = TRUE
)
BranchGLM.fit(
 Х,
 у,
 family,
 link,
 offset = NULL,
 method = "Fisher",
 grads = 10,
 parallel = FALSE,
 nthreads = 8,
 init = NULL,
 maxit = NULL,
 tol = 1e-06
)
```

Arguments

formula	a formula for the model.
data	a data.frame, list or environment (or object coercible by as.data.frame to a data.frame), containing the variables in formula. Neither a matrix nor an array will be accepted.
family	the distribution used to model the data, one of "gaussian", "gamma", "binomial", or "poisson".
link	the link used to link the mean structure to the linear predictors. One of "identity", "logit", "probit", "cloglog", "sqrt", "inverse", or "log". The accepted links depend on the specified family, see more in details.
offset	the offset vector, by default the zero vector is used.
method	one of "Fisher", "BFGS", or "LBFGS". BFGS and L-BFGS are quasi-newton methods which are typically faster than Fisher's scoring when there are many covariates (at least 50).
grads	a positive integer to denote the number of gradients used to approximate the inverse information with, only for method = "LBFGS".
parallel	a logical value to indicate if parallelization should be used.

nthreads	a positive integer to denote the number of threads used with OpenMP, only used if $parallel = TRUE$.
tol	a positive number to denote the tolerance used to determine model convergence.
maxit	a positive integer to denote the maximum number of iterations performed. The default for Fisher's scoring is 50 and for the other methods the default is 200.
init	a numeric vector of initial values for the betas, if not specified then they are automatically selected via linear regression with the transformation specified by the link function. This is ignored for linear regression models.
fit	a logical value to indicate whether to fit the model or not.
contrasts	see contrasts.arg of model.matrix.default.
keepData	a logical value to indicate whether or not to store a copy of data and the design matrix, the default is TRUE. If this is FALSE, then the results from this cannot be used inside of VariableSelection.
keepY	a logical value to indicate whether or not to store a copy of y, the default is TRUE. If this is FALSE, then the binomial GLM helper functions may not work and this cannot be used inside of VariableSelection.
x	design matrix used for the fit, must be numeric.
у	outcome vector, must be numeric.

Details

Fitting:

Can use BFGS, L-BFGS, or Fisher's scoring to fit the GLM. BFGS and L-BFGS are typically faster than Fisher's scoring when there are at least 50 covariates and Fisher's scoring is typically best when there are fewer than 50 covariates. This function does not currently support the use of weights. In the special case of gaussian regression with identity link the method argument is ignored and the normal equations are solved directly.

The models are fit in C++ by using Rcpp and RcppArmadillo. In order to help convergence, each of the methods makes use of a backtracking line-search using the strong Wolfe conditions to find an adequate step size. There are three conditions used to determine convergence, the first is whether there is a sufficient decrease in the negative log-likelihood, the second is whether the 12-norm of the score is sufficiently small, and the last condition is whether the change in each of the beta coefficients is sufficiently small. The tol argument controls all of these criteria. If the algorithm fails to converge, then iterations will be -1.

All observations with any missing values are removed before model fitting.

BranchGLM.fit can be faster than calling BranchGLM if the x matrix and y vector are already available, but doesn't return as much information. The object returned by BranchGLM.fit is not of class BranchGLM, so all of the methods for BranchGLM objects such as predict or VariableSelection cannot be used.

Dispersion Parameter:

The dispersion parameter for gamma regression is estimated via maximum likelihood, very similar to the gamma.dispersion function from the MASS package. The dispersion parameter for gaussian regression is also estimated via maximum likelihood estimation.

Families and Links:

The binomial family accepts "cloglog", "log", "logit", and "probit" as possible link functions. The gamma and gaussian families accept "identity", "inverse", "log", and "sqrt" as possible link functions. The Poisson family accepts "identity", "log", and "sqrt" as possible link functions.

Value

BranchGLM returns a BranchGLM object which is a list with the following components

coefficients a matrix with the coefficient estimates, SEs, Wald test statistics, and p-values iterations number of iterations it took the algorithm to converge, if the algorithm failed to

converge then this is -1

dispersion the value of the dispersion parameter logLik the log-likelihood of the fitted model

vcov the variance-covariance matrix of the fitted model

resDev the residual deviance of the fitted model

AIC the AIC of the fitted model preds predictions from the fitted model

linear predictors from the fitted model

tol tolerance used to fit the model

maxit maximum number of iterations used to fit the model

formula used to fit the model

method iterative method used to fit the model

grads number of gradients used to approximate inverse information for L-BFGS

y vector used in the model, not included if keepY = FALSE

x design matrix used to fit the model, not included if keepData = FALSE

offset offset vector in the model, not included if keepData = FALSE supplied offset vector, not included if keepData = FALSE

data original data argument supplied to the function, not included if keepData =

FALSE

mf the model frame, not included if keepData = FALSE

number of observations in the design matrix

names names of the predictor variables

yname name of y variable

parallel whether parallelization was employed to speed up model fitting process

missing number of missing values removed from the original dataset

link link function used to model the data

family used to model the data

ylevel the levels of y, only included for binomial glms

xlev the levels of the factors in the dataset

terms the terms object used

BranchGLM. fit returns a list with the following components

coefficients a matrix with the coefficients estimates, SEs, Wald test statistics, and p-values iterations number of iterations it took the algorithm to converge, if the algorithm failed to

converge then this is -1

dispersion the value of the dispersion parameter logLik the log-likelihood of the fitted model

vcov the variance-covariance matrix of the fitted model

resDev the residual deviance of the fitted model

AIC the AIC of the fitted model

preds predictions from the fitted model

linpreds linear predictors from the fitted model

tol tolerance used to fit the model

maxit maximum number of iterations used to fit the model

References

McCullagh, P., & Nelder, J. A. (1989). Generalized Linear Models (2nd ed.). Chapman & Hall.

See Also

predict.BranchGLM, coef.BranchGLM, VariableSelection, confint.BranchGLM, logLik.BranchGLM

```
Data <- iris
# Linear regression
## Using BranchGLM
BranchGLM(Sepal.Length ~ ., data = Data, family = "gaussian", link = "identity")
## Using BranchGLM.fit
x <- model.matrix(Sepal.Length ~ ., data = Data)</pre>
y <- Data$Sepal.Length
BranchGLM.fit(x, y, family = "gaussian", link = "identity")
# Gamma regression
## Using BranchGLM
BranchGLM(Sepal.Length ~ ., data = Data, family = "gamma", link = "log")
### init
BranchGLM(Sepal.Length ~ ., data = Data, family = "gamma", link = "log",
init = rep(0, 6), maxit = 50, tol = 1e-6, contrasts = NULL)
### method
BranchGLM(Sepal.Length ~ ., data = Data, family = "gamma", link = "log",
init = rep(0, 6), maxit = 50, tol = 1e-6, contrasts = NULL, method = "LBFGS")
```

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```
### offset
BranchGLM(Sepal.Length ~ ., data = Data, family = "gamma", link = "log",
init = rep(0, 6), maxit = 50, tol = 1e-6, contrasts = NULL,
offset = Data$Sepal.Width)

## Using BranchGLM.fit
x <- model.matrix(Sepal.Length ~ ., data = Data)
y <- Data$Sepal.Length
BranchGLM.fit(x, y, family = "gamma", link = "log", init = rep(0, 6),
maxit = 50, tol = 1e-6, offset = Data$Sepal.Width)</pre>
```

Cindex

Cindex/AUC

Description

Calculates the c-index/AUC.

Usage

```
Cindex(object, ...)

AUC(object, ...)

## S3 method for class 'numeric'
Cindex(object, y, ...)

## S3 method for class 'BranchGLM'
Cindex(object, ...)

## S3 method for class 'BranchGLMROC'
Cindex(object, ...)
```

Arguments

object a BranchGLM object, a BranchGLMROC object, or a numeric vector.

... further arguments passed to other methods.

y Observed values, can be a numeric vector of 0s and 1s, a two-level factor vector, or a logical vector.

Details

Uses trapezoidal rule to calculate AUC when given a BranchGLMROC object and uses Mann-Whitney U to calculate it otherwise. The trapezoidal rule method is less accurate, so the two methods may give different results.

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Value

A number corresponding to the c-index/AUC.

Examples

```
Data <- ToothGrowth
Fit <- BranchGLM(supp ~ ., data = Data, family = "binomial", link = "logit")
Cindex(Fit)
AUC(Fit)</pre>
```

coef.BranchGLM

Extract Coefficients from BranchGLM Objects

Description

Extracts beta coefficients from BranchGLM objects.

Usage

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

Arguments

object a BranchGLM object.

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

Value

A named vector with the corresponding coefficient estimates.

coef.BranchGLMVS

Extract Coefficients from BranchGLMVS or summary.BranchGLMVS Objects

Description

Extracts beta coefficients from BranchGLMVS or summary.BranchGLMVS objects.

```
## $3 method for class 'BranchGLMVS'
coef(object, which = 1, ...)
## $3 method for class 'summary.BranchGLMVS'
coef(object, which = 1, ...)
```

confint.BranchGLM 9

Arguments

object a BranchGLMVS or summary.BranchGLMVS object.

which a numeric vector of indices or "all" to indicate which models to get coefficients

from, the default is 1 which is used for the best model. For the branch and bound algorithms the number k is used for the kth best model and for the stepwise algorithms the number k is used for the model that is k-1 steps away from the

final model.

... ignored.

Value

A numeric matrix with the corresponding coefficient estimates.

Examples

```
Data <- iris
Fit <- BranchGLM(Sepal.Length ~ ., data = Data,
family = "gaussian", link = "identity")

# Doing branch and bound selection
VS <- VariableSelection(Fit, type = "branch and bound", metric = "BIC",
bestmodels = 10, showprogress = FALSE)

## Getting coefficients from best model
coef(VS, which = 1)

## Getting coefficients from all best models
coef(VS, which = "all")</pre>
```

confint.BranchGLM

Likelihood Ratio Confidence Intervals for Beta Coefficients for BranchGLM Objects

Description

Finds profile likelihood ratio confidence intervals for beta coefficients with the ability to calculate the intervals in parallel.

```
## S3 method for class 'BranchGLM'
confint(object, parm, level = 0.95, parallel = FALSE, nthreads = 8, ...)
```

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Arguments

object a BranchGLM object.

parm a specification of which parameters are to be given confidence intervals, either

a vector of numbers or a vector of names. If missing, all parameters are consid-

ered.

level the confidence level required.

parallel a logical value to indicate if parallelization should be used.

nthreads a positive integer to denote the number of threads used with OpenMP, only used

if parallel = TRUE.

... further arguments passed from other methods.

Details

Endpoints of the confidence intervals that couldn't be found by the algorithm are filled in with NA. When there is a lot of multicollinearity in the data the algorithm may have problems finding many of the intervals.

Value

An object of class BranchGLMCIs which is a list with the following components.

CIs a numeric matrix with the confidence intervals

level the supplied level

MLE a numeric vector of the MLEs of the coefficients

See Also

plot.BranchGLMCIs, plotCI

```
Data <- iris
### Fitting linear regression model
mymodel <- BranchGLM(Sepal.Length ~ ., data = Data, family = "gaussian", link = "identity")
### Getting confidence intervals
CIs <- confint(mymodel, level = 0.95)
CIs
### Plotting CIs
plot(CIs, mary = 7, cex.y = 0.9)</pre>
```

formula.BranchGLM

formula.BranchGLM

Extract Model Formula from BranchGLM Objects

Description

Extracts model formula from BranchGLM objects.

Usage

```
## S3 method for class 'BranchGLM'
formula(x, ...)
```

Arguments

x a BranchGLM object.

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

Value

a formula representing the model used to obtain object.

logLik.BranchGLM

Extract Log-Likelihood from BranchGLM Objects

Description

Extracts log-likelihood from BranchGLM objects.

Usage

```
## S3 method for class 'BranchGLM'
logLik(object, ...)
```

Arguments

object a BranchGLM object.

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

Value

An object of class logLik which is a number corresponding to the log-likelihood with the following attributes: "df" (degrees of freedom) and "nobs" (number of observations).

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MultipleROCCurves

Plotting Multiple ROC Curves

Description

Plotting Multiple ROC Curves

Usage

```
MultipleROCCurves(
    ...,
  legendpos = "bottomright",
  title = "ROC Curves",
  colors = NULL,
  names = NULL,
  lty = 1,
  lwd = 1
)
```

Arguments

... any number of BranchGLMROC objects.

legendpos a keyword to describe where to place the legend, such as "bottomright". The

default is "bottomright"

title title for the plot.

colors vector of colors to be used on the ROC curves.

names vector of names used to create a legend for the ROC curves.

1ty vector of linetypes used to create the ROC curves or a single linetype to be used

for all ROC curves.

lwd vector of linewidths used to create the ROC curves or a single linewidth to be

used for all ROC curves.

Value

This only produces a plot, nothing is returned.

```
Data <- ToothGrowth
### Logistic ROC
LogisticFit <- BranchGLM(supp ~ ., data = Data, family = "binomial", link = "logit")
LogisticROC <- ROC(LogisticFit)
### Probit ROC
ProbitFit <- BranchGLM(supp ~ ., data = Data, family = "binomial", link = "probit")</pre>
```

nobs.BranchGLM

nobs.BranchGLM

Extract Number of Observations from BranchGLM Objects

Description

Extracts number of observations from BranchGLM objects.

Usage

```
## S3 method for class 'BranchGLM'
nobs(object, ...)
```

Arguments

object a BranchGLM object.

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

Value

A single number indicating the number of observations used to fit the model.

plot.BranchGLM

Plot Method for BranchGLM Objects

Description

Creates a plot to help visualize fitted values from BranchGLM objects.

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

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Arguments

x a BranchGLM object.

... further arguments passed to plot.default.

Value

This only produce a plot, nothing is returned.

Examples

```
Data <- iris
Fit <- BranchGLM(Sepal.Length ~ ., data = Data, family = "gaussian", link = "identity")
plot(Fit)</pre>
```

plot.BranchGLMCIs

Plot Method for BranchGLMCIs Objects

Description

Creates a plot to visualize confidence intervals from BranchGLMCIs objects.

Usage

```
## S3 method for class 'BranchGLMCIs'
plot(x, which = "all", mary = 5, ...)
```

Arguments

х	a BranchGLMCIs object.
which	which intervals to plot, can use a numeric vector of indices, a character vector of names of desired variables, or "all" to plot all intervals.
mary	a numeric value used to determine how large to make margin of y-axis. If variable names are cut-off, consider increasing this from the default value of 5.

... further arguments passed to plotCI.

Value

This only produces a plot, nothing is returned.

See Also

plotCI

plot.BranchGLMROC 15

Examples

```
Data <- iris
### Fitting linear regression model
mymodel <- BranchGLM(Sepal.Length ~ ., data = Data, family = "gaussian", link = "identity")
### Getting confidence intervals
CIs <- confint(mymodel, level = 0.95)
CIs
### Plotting CIs
plot(CIs, mary = 7, cex.y = 0.9)</pre>
```

plot.BranchGLMROC

Plot Method for BranchGLMROC Objects

Description

This plots a ROC curve.

Usage

```
## S3 method for class 'BranchGLMROC'
plot(x, xlab = "1 - Specificity", ylab = "Sensitivity", type = "1", ...)
```

Arguments

```
x a BranchGLMROC object.
xlab label for the x-axis.
ylab label for the y-axis.
type what type of plot to draw, see more details at plot.default.
... further arguments passed to plot.default.
```

Value

This only produces a plot, nothing is returned.

```
Data <- ToothGrowth
Fit <- BranchGLM(supp ~ ., data = Data, family = "binomial", link = "logit")
MyROC <- ROC(Fit)
plot(MyROC)</pre>
```

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plot.BranchGLMVS

Plot Method for summary.BranchGLMVS and BranchGLMVS Objects

Description

Creates plots to help visualize variable selection results from BranchGLMVS or summary.BranchGLMVS objects.

Usage

```
## S3 method for class 'BranchGLMVS'
plot(
  Х,
  ptype = "both",
 marnames = 7,
  addLines = TRUE,
  type = "b",
  horiz = FALSE,
  cex.names = 1,
  cex.lab = 1,
  cex.axis = 1,
  cex.legend = 1,
  cols = c("deepskyblue", "indianred", "forestgreen"),
)
## S3 method for class 'summary.BranchGLMVS'
plot(
  Х,
 ptype = "both",
 marnames = 7,
  addLines = TRUE,
  type = "b",
  horiz = FALSE,
  cex.names = 1,
  cex.lab = 1,
  cex.axis = 1,
  cex.legend = 1,
 cols = c("deepskyblue", "indianred", "forestgreen"),
)
```

Arguments

x a summary.BranchGLMVS or BranchGLMVS object.

ptype the type of plot to produce, look at details for more explanation.

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marnames	a numeric value used to determine how large to make margin of axis with variable names, this is only for the "variables" plot. If variable names are cut-off, consider increasing this from the default value of 7.
addLines	a logical value to indicate whether or not to add black lines to separate the models for the "variables" plot. This is typically useful for smaller amounts of models, but can be annoying if there are many models.
type	what type of plot to draw for the "metrics" plot, see more details at plot.default.
horiz	a logical value to indicate whether models should be displayed horizontally for the "variables" plot.
cex.names	how big to make variable names in the "variables" plot.
cex.lab	how big to make axis labels.
cex.axis	how big to make axis annotation.
cex.legend	how big to make legend labels.
cols	the colors used to create the "variables" plot. Should be a character vector of length 3, the first color will be used for included variables, the second color will be used for excluded variables, and the third color will be used for kept variables.
•••	further arguments passed to plot.default for the "metrics" plot and image.default for the "variables" plot.

Details

The different values for ptype are as follows

- "metrics" for a plot that displays the metric values ordered by rank for the branch and bound algorithms or a plot which displays the metric values in the path taken by the stepwise algorithms
- "variables" for a plot that displays which variables are in each of the top models for the branch and bound algorithms or a plot which displays the path taken by the stepwise algorithms
- "both" for both plots

If there are so many models that the "variables" plot appears to be entirely black, then set addLines to FALSE.

Value

This only produces plots, nothing is returned.

```
Data <- iris
Fit <- BranchGLM(Sepal.Length ~ ., data = Data, family = "gaussian", link = "identity")
# Doing branch and bound selection
VS <- VariableSelection(Fit, type = "branch and bound", metric = "BIC", bestmodels = 10, showprogress = FALSE)
VS</pre>
```

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```
## Getting summary of the process
Summ <- summary(VS)
Summ

## Plotting the BIC of best models
plot(Summ, type = "b", ptype = "metrics")

## Plotting the BIC of the best models
plot(Summ, ptype = "variables")

### Alternative colors
plot(Summ, ptype = "variables",
cols = c("yellowgreen", "purple1", "grey50"))

### Smaller text size for names
plot(Summ, ptype = "variables", cex.names = 0.75)</pre>
```

plotCI

Plot Confidence Intervals

Description

Creates a plot to display confidence intervals.

Usage

```
plotCI(
   CIs,
   points = NULL,
   ylab = "",
   ylas = 2,
   cex.y = 1,
   decreasing = FALSE,
   ...
)
```

Arguments

CIs	a numeric matrix of confidence intervals, must have exactly 2 columns. The variable names displayed in the plot are taken from the column names.
points	points to be plotted in the middle of the CIs, typically means or medians. The default is to plot the midpoints of the intervals.
ylab	a label for the y-axis.
ylas	the style of the y-axis label, see more about this at las in par.
cex.y	font size used for variable names on y-axis.

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decreasing a logical value indicating if confidence intervals should be displayed in decreasing or increasing order according to points. Can use NA if no ordering is desired.

... further arguments passed to plot.default.

Value

This only produces a plot, nothing is returned.

Examples

```
Data <- iris
### Fitting linear regression model
mymodel <- BranchGLM(Sepal.Length ~ ., data = Data, family = "gaussian", link = "identity")
### Getting confidence intervals
CIs <- confint.default(mymodel, level = 0.95)
xlim <- c(min(CIs), max(CIs))
### Plotting CIs
par(mar = c(5, 7, 3, 1) + 0.1)
plotCI(CIs, main = "95% Confidence Intervals", xlim = xlim, cex.y = 0.9,
xlab = "Beta Coefficients")
abline(v = 0)</pre>
```

predict.BranchGLM

Predict Method for BranchGLM Objects

Description

Obtains predictions from BranchGLM objects.

```
## S3 method for class 'BranchGLM'
predict(
  object,
  newdata = NULL,
  offset = NULL,
  type = "response",
  na.action = na.pass,
  ...
)
```

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Arguments

object a BranchGLM object.

newdata a data.frame, if not specified then the data the model was fit on is used.

offset a numeric vector containing the offset variable, this is ignored if newdata is not supplied.

type one of "linpreds" which is on the scale of the linear predictors or "response" which is on the scale of the response variable. If not specified, then "response" is used.

na.action a function which indicates what should happen when the data contains NAs.

The default is na.pass. This is ignored if newdata is not supplied and data isn't

included in the supplied BranchGLM object.

further arguments passed to or from other methods.

Value

A numeric vector of predictions.

```
Data <- airquality
# Example without offset
Fit <- BranchGLM(Temp ~ ., data = Data, family = "gaussian", link = "identity")
## Using default na.action
predict(Fit)
## Using na.omit
predict(Fit, na.action = na.omit)
## Using new data
predict(Fit, newdata = Data[1:20, ], na.action = na.pass)
# Using offset
FitOffset <- BranchGLM(Temp ~ . - Day, data = Data, family = "gaussian",
link = "identity", offset = Data$Day * -0.1)
## Getting predictions for data used to fit model
### Don't need to supply offset vector
predict(FitOffset)
## Getting predictions for new dataset
### Need to include new offset vector since we are
### getting predictions for new dataset
predict(FitOffset, newdata = Data[1:20, ], offset = Data$Day[1:20] * -0.1)
```

predict.BranchGLMVS

 $\begin{tabular}{ll} predict. Branch GLMVS & Predict Method for Branch GLMVS or summary. Branch GLMVS Objects \\ \end{tabular}$

Description

Obtains predictions from BranchGLMVS or summary.BranchGLMVS objects.

Usage

```
## $3 method for class 'BranchGLMVS'
predict(object, which = 1, ...)
## $3 method for class 'summary.BranchGLMVS'
predict(object, which = 1, ...)
```

Arguments

object a BranchGLMVS or summary.BranchGLMVS object.

which a positive integer to indicate which model to get predictions from, the default

is 1 which is used for the best model. For the branch and bound algorithms the number k is used for the kth best model and for the stepwise algorithms the number k is used for the model that is k-1 steps away from the final model.

... further arguments passed to predict.BranchGLM.

Value

A numeric vector of predictions.

See Also

```
predict.BranchGLM
```

```
Data <- iris
Fit <- BranchGLM(Sepal.Length ~ ., data = Data,
family = "gamma", link = "log")

# Doing branch and bound selection
VS <- VariableSelection(Fit, type = "branch and bound", metric = "BIC",
bestmodels = 10, showprogress = FALSE)

## Getting predictions from best model
predict(VS, which = 1)

## Getting linear predictors from 5th best model
predict(VS, which = 5, type = "linpreds")</pre>
```

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print.BranchGLM

Print Method for BranchGLM Objects

Description

Print method for BranchGLM objects.

Usage

```
## S3 method for class 'BranchGLM'
print(x, coefdigits = 4, digits = 2, ...)
```

Arguments

x a BranchGLM object.

coefdigits number of digits to display for coefficients table.

digits number of digits to display for information after table.

further arguments passed to or from other methods.

Value

The supplied BranchGLM object.

print.BranchGLMCIs

Print Method for BranchGLMCIs Objects

Description

Print method for BranchGLMCIs objects.

Usage

```
## S3 method for class 'BranchGLMCIs'
print(x, digits = 4, ...)
```

Arguments

x a BranchGLMCIs object.

digits number of significant digits to display.

... further arguments passed from other methods.

Value

The supplied BranchGLMCIs object.

print.BranchGLMROC

print.BranchGLMROC Print Method for BranchGLMROC Objects

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Description

Print method for BranchGLMROC objects.

Usage

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

Arguments

x a BranchGLMROC object.

. . . further arguments passed to other methods.

Value

The supplied BranchGLMROC object.

Description

Print method for BranchGLMTable objects.

Usage

```
## S3 method for class 'BranchGLMTable'
print(x, digits = 4, ...)
```

Arguments

x a BranchGLMTable object.
digits number of digits to display.

. . . further arguments passed to other methods.

Value

The supplied BranchGLMTable object.

print.BranchGLMVS

Print Method for BranchGLMVS Objects

Description

Print method for BranchGLMVS objects.

Usage

```
## S3 method for class 'BranchGLMVS'
print(x, digits = 2, ...)
```

Arguments

```
x a BranchGLMVS object.digits number of digits to display.further arguments passed to other methods.
```

Value

The supplied BranchGLMVS object.

```
print.summary.BranchGLMVS
```

Print Method for summary.BranchGLMVS Objects

Description

Print method for summary.BranchGLMVS objects.

Usage

```
## S3 method for class 'summary.BranchGLMVS'
print(x, digits = 2, ...)
```

Arguments

```
x a summary.BranchGLMVS object.digits number of digits to display.... further arguments passed to other methods.
```

Value

The supplied summary.BranchGLMVS object.

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ROC

ROC Curve

Description

Creates an ROC curve.

Usage

```
ROC(object, ...)
## S3 method for class 'numeric'
ROC(object, y, ...)
## S3 method for class 'BranchGLM'
ROC(object, ...)
```

Arguments

object a BranchGLM object or a numeric vector.

... further arguments passed to other methods.

y observed values, can be a numeric vector of 0s and 1s, a two-level factor vector, or a logical vector.

Value

A BranchGLMROC object which can be plotted with plot(). The AUC can also be calculated using AUC().

Examples

```
Data <- ToothGrowth
Fit <- BranchGLM(supp ~ ., data = Data, family = "binomial", link = "logit")
MyROC <- ROC(Fit)
plot(MyROC)</pre>
```

summary.BranchGLMVS

Summary Method for BranchGLMVS Objects

Description

Summary method for BranchGLMVS objects.

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

Arguments

object a BranchGLMVS object.

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

Value

An object of class summary. BranchGLMVS which is a list with the following components

results a data.frame which has the metric values for the best models along with the sets

of variables included in each model

VS the supplied BranchGLMVS object

formulas a list containing the formulas of the best models

metric the metric used to perform variable selection

See Also

plot.summary.BranchGLMVS, coef.summary.BranchGLMVS, predict.summary.BranchGLMVS

```
Data <- iris
Fit <- BranchGLM(Sepal.Length ~ ., data = Data, family = "gaussian", link = "identity")
# Doing branch and bound selection
VS <- VariableSelection(Fit, type = "branch and bound", metric = "BIC",
bestmodels = 10, showprogress = FALSE)
VS
## Getting summary of the process
Summ <- summary(VS)
Summ
## Plotting the BIC of the best models
plot(Summ, type = "b")
## Plotting the variables in the best models
plot(Summ, ptype = "variables")
## Getting coefficients
coef(Summ)</pre>
```

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Description

Creates a confusion matrix and calculates related measures.

Usage

```
Table(object, ...)
## S3 method for class 'numeric'
Table(object, y, cutoff = 0.5, ...)
## S3 method for class 'BranchGLM'
Table(object, cutoff = 0.5, ...)
```

Arguments

object	a BranchGLM object or a numeric vector.
	further arguments passed to other methods.
У	observed values, can be a numeric vector of 0s and 1s, a two-level factor vector, or a logical vector.
cutoff	cutoff for predicted values, the default is 0.5.

Value

A BranchGLMTable object which is a list with the following components

```
table a matrix corresponding to the confusion matrix
accuracy a number corresponding to the accuracy
sensitivity a number corresponding to the sensitivity
specificity a number corresponding to the specificity
PPV a number corresponding to the positive predictive value
levels a vector corresponding to the levels of the response variable
```

```
Data <- ToothGrowth
Fit <- BranchGLM(supp ~ ., data = Data, family = "binomial", link = "logit")
Table(Fit)</pre>
```

VariableSelection

Variable Selection for GLMs

Description

Performs forward selection, backward elimination, and efficient best subset variable selection with information criterion for generalized linear models (GLMs). Best subset selection is performed with branch and bound algorithms to greatly speed up the process.

```
VariableSelection(object, ...)
## S3 method for class 'formula'
VariableSelection(
  object,
  data,
  family,
  link,
  offset = NULL,
  method = "Fisher",
  type = "switch branch and bound",
  metric = "AIC",
  bestmodels = NULL,
  cutoff = NULL,
  keep = NULL,
  keepintercept = TRUE,
  maxsize = NULL,
  grads = 10,
  parallel = FALSE,
  nthreads = 8,
  tol = 1e-06,
  maxit = NULL,
  contrasts = NULL,
  showprogress = TRUE,
)
## S3 method for class 'BranchGLM'
VariableSelection(
  object,
  type = "switch branch and bound",
  metric = "AIC",
  bestmodels = NULL,
  cutoff = NULL,
  keep = NULL,
  keepintercept = TRUE,
```

```
maxsize = NULL,
parallel = FALSE,
nthreads = 8,
showprogress = TRUE,
...
)
```

Arguments

object a formula or a BranchGLM object.

... further arguments.

data a data frame, list or environment (or object coercible by as data frame to a data frame),

containing the variables in formula. Neither a matrix nor an array will be ac-

cepted.

family the distribution used to model the data, one of "gaussian", "gamma", "binomial",

or "poisson".

link the link used to link the mean structure to the linear predictors. One of "identity",

"logit", "probit", "cloglog", "sqrt", "inverse", or "log".

offset the offset vector, by default the zero vector is used.

method one of "Fisher", "BFGS", or "LBFGS". Fisher's scoring is recommended for

forward selection and the branch and bound algorithms since they will typically

fit many models with a small number of covariates.

type one of "forward", "backward", "branch and bound", "backward branch and bound",

or "switch branch and bound" to indicate the type of variable selection to perform. The default value is "switch branch and bound". See more about these

algorithms in details

metric the metric used to choose the best models, the default is "AIC", but "BIC" and

"HQIC" are also available. AIC is the Akaike information criterion, BIC is the Bayesian information criterion, and HQIC is the Hannan-Quinn information

criterion.

bestmodels a positive integer to indicate the number of the best models to find according to

the chosen metric or NULL. If this is NULL, then cutoff is used instead. This is

only used for the branch and bound algorithms.

cutoff a non-negative number which indicates that the function should return all models

that have a metric value within cutoff of the best metric value or NULL. Only one of this or bestmodels should be specified and when both are NULL a cutoff

of 0 is used. This is only used for the branch and bound algorithms.

keep a character vector of names to denote variables that must be in the models.

keepintercept a logical value to indicate whether to keep the intercept in all models, only used

if an intercept is included in the formula.

maxsize a positive integer to denote the maximum number of variables to consider in a

single model, the default is the total number of variables. This number adds onto any variables specified in keep. This argument only works for type = "forward" and type = "branch and bound". This argument is now deprecated.

grads a positive integer to denote the number of gradients used to approximate the

inverse information with, only for method = "LBFGS".

parallel a logical value to indicate if parallelization should be used.

nthreads a positive integer to denote the number of threads used with OpenMP, only used

if parallel = TRUE.

tol a positive number to denote the tolerance used to determine model convergence.

maxit a positive integer to denote the maximum number of iterations performed. The

default for Fisher's scoring is 50 and for the other methods the default is 200.

contrasts see contrasts.arg of model.matrix.default.

showprogress a logical value to indicate whether to show progress updates for branch and

bound algorithms.

Details

Variable Selection Details:

The supplied formula or the formula from the fitted model is treated as the upper model. The variables specified in keep along with an intercept (if included in formula and keepintercept = TRUE) is the lower model. Factor variables are either kept in their entirety or entirely removed and interaction terms are properly handled. All observations that have any missing values in the upper model are removed.

Branch and Bound Algorithms:

The branch and bound algorithm is an efficient algorithm used to find the optimal models. The backward branch and bound algorithm is very similar to the branch and bound algorithm, except it tends to be faster when the best models contain most of the variables. The switch branch and bound algorithm is a combination of the two algorithms and is typically the fastest of the 3 branch and bound algorithms. All of the branch and bound algorithms are guaranteed to find the optimal models (up to numerical precision).

GLM Fitting:

Fisher's scoring is recommended for branch and bound selection and forward selection. L-BFGS may be faster for backward elimination especially when there are many variables.

Value

A BranchGLMVS object which is a list with the following components

initmodel the BranchGLM object corresponding to the upper model

numchecked number of models fit

names character vector of the names of the predictor variables

order the order the variables were added to the model or removed from the model, this

is only included for the stepwise algorithms

type type of variable selection employed

optType whether the type specified used a heuristic or exact algorithm

metric metric used to select best models

bestmodels numeric matrix used to describe the best models for the branch and bound al-

gorithms or a numeric matrix describing the models along the path taken for

stepwise algorithms

bestmetrics numeric vector with the best metrics found in the search for the branch and

bound algorithms or a numeric vector with the metric values along the path

taken for stepwise algorithms

beta numeric matrix of beta coefficients for the models in bestmodels

cutoff the cutoff that was used, this is set to -1 if bestmodels was used instead or if a

stepwise algorithm was used

keep vector of which variables were kept through the selection process

keepintercept a boolean value denoting whether to keep the intercept through the selection

process or not

See Also

plot.BranchGLMVS, coef.BranchGLMVS, predict.BranchGLMVS, summary.BranchGLMVS

```
Data <- iris
Fit <- BranchGLM(Sepal.Length ~ ., data = Data, family = "gaussian",
link = "identity")
# Doing branch and bound selection
VS <- VariableSelection(Fit, type = "branch and bound", metric = "BIC",
bestmodels = 10, showprogress = FALSE)
## Plotting the BIC of the best models
plot(VS, type = "b")
## Getting the coefficients of the best model according to BIC
FinalModel <- coef(VS, which = 1)
FinalModel
# Now doing it in parallel (although it isn't necessary for this dataset)
parVS <- VariableSelection(Fit, type = "branch and bound", parallel = TRUE,</pre>
metric = "BIC", bestmodels = 10, showprogress = FALSE)
## Getting the coefficients of the best model according to BIC
FinalModel <- coef(parVS, which = 1)</pre>
FinalModel
# Using a formula
formVS <- VariableSelection(Sepal.Length ~ ., data = Data, family = "gaussian",
link = "identity", metric = "BIC", type = "branch and bound", bestmodels = 10,
showprogress = FALSE)
## Getting the coefficients of the best model according to BIC
FinalModel <- coef(formVS, which = 1)
```

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```
FinalModel
# Using the keep argument
keepVS <- VariableSelection(Fit, type = "branch and bound",</pre>
keep = c("Species", "Petal.Width"), metric = "BIC", bestmodels = 4,
showprogress = FALSE)
keepVS
## Getting the coefficients from the fourth best model according to BIC when
## keeping Petal.Width and Species in every model
FinalModel <- coef(keepVS, which = 4)</pre>
FinalModel
# Treating categorical variable beta parameters separately
## This function automatically groups together parameters from a categorical variable
## to avoid this, you need to create the indicator variables yourself
x \leftarrow model.matrix(Sepal.Length \sim ., data = iris)
Sepal.Length <- iris$Sepal.Length</pre>
Data <- cbind.data.frame(Sepal.Length, x[, -1])
VSCat <- VariableSelection(Sepal.Length ~ ., data = Data, family = "gaussian",
link = "identity", metric = "BIC", bestmodels = 10, showprogress = FALSE)
VSCat
## Plotting results
plot(VSCat, cex.names = 0.75)
```

vcov.BranchGLM

Extract covariance matrix from BranchGLM Objects

Description

Extracts covariance matrix of beta coefficients from BranchGLM objects.

Usage

```
## S3 method for class 'BranchGLM'
vcov(object, ...)
```

Arguments

```
object a BranchGLM object.... further arguments passed to or from other methods.
```

Value

A numeric matrix which is the covariance matrix of the beta coefficients.

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