Package 'HCTR'

October 12, 2022

Title Higher Criticism Tuned Regression

Version 0.1.1

Description A novel searching scheme for tuning parameter in high-dimensional
penalized regression. We propose a new estimate of the regularization
parameter based on an estimated lower bound of the proportion of false
null hypotheses (Meinshausen and Rice (2006) <doi:10.1214 009053605000000741="">).</doi:10.1214>
The bound is estimated by applying the empirical null distribution of the higher
criticism statistic, a second-level significance testing, which is constructed
by dependent p-values from a multi-split regression and aggregation method
(Jeng, Zhang and Tzeng (2019) <doi:10.1080 01621459.2018.1518236="">). An estimate</doi:10.1080>
of tuning parameter in penalized regression is decided corresponding to the lower
bound of the proportion of false null hypotheses. Different penalized
regression methods are provided in the multi-split algorithm.
Depends R (>= $3.4.0$)
Imports glmnet (>= 2.0-18), harmonicmeanp (>= 3.0), MASS, nevreg (>=
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bounding.seq

Bounding Sequence

Description

Calculates bounding sequence of higher crticism for proportion estimator using p-values

Usage

```
bounding.seq(p.value, alpha)
```

Arguments

p.value A matrix of p-values from permutation: row is from each permutation; column

is from each variable.

alpha Probability of Type I error for bounding sequence, the default value is 1 / sqrt(log(p)),

where p is number of p-values in each permutation.

Value

A bounding value of higher criticism with (1 - alpha) confidence.

References

Jeng XJ, Zhang T, Tzeng J (2019). "Efficient Signal Inclusion With Genomic Applications." *Journal of the American Statistical Association*, 1–23.

Examples

```
set.seed(10)
X <- matrix(runif(n = 10000, min = 0, max = 1), nrow = 100)
result <- bounding.seq(p.value = X)</pre>
```

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est.lambda	Estimated Lambda
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Description

Estimate upper and lower bound of new tuning region of regularization parameter Lambda.

Usage

```
est.lambda(cv.fit, pihat, p, cov.num = 0)
```

Arguments

cv.fit	An object of either class "cv.glmnet" from glmnet::cv.glmnet() or class "cv.ncvreg" from ncvreg::cv.ncvreg(), which is a list generated by a cross-validation fit.
pihat	eatimated proprtion from HCTR::est.prop().
р	Total number of variables, except for covariates.
cov.num	Number of covariates in model, default is 0. Covariate matrix, W, is assumed on the left side of variable matrix, X. The column index of covariates are before those of variables.

Value

A list of (1) lambda.max, upper bound of new tuning region; (2) lambda.min, lower bound of new tuning region.

Examples

```
set.seed(10)
X <- matrix(rnorm(20000), nrow = 100)
beta <- rep(0, 200)
beta[1:100] <- 5
Y <- MASS::mvrnorm(n = 1, mu = X%*%beta, Sigma = diag(100))
fit <- glmnet::cv.glmnet(x = X, y = Y)
pihat <- 0.01
result <- est.lambda(cv.fit = fit, pihat = pihat, p = ncol(X))</pre>
```

est.prop

Proportion Estimation

Description

Estimates false null hypothesis Proportion from multiple p-values using higher criticism test estimator.

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Usage

```
est.prop(p.value, cn, adj = TRUE)
```

Arguments

p. value A sequence of p-values from test data, not including p-values from covariates.
 cn A value of bounding sequence generated by HCTR::bounding.seq().
 adj A boolean algebra to decide whether to use adjusted Higher Criticism test statistic, the default value is TRUE.

Value

An estimated proportion of false null hypothesis.

References

Meinshausen N, Rice J (2006). "Estimating the proportion of false null hypotheses among a large number of independently tested hypotheses." *The Annals of Statistics*, **34**(1), 373–393.

Examples

```
set.seed(10)
X <- matrix(runif(n = 10000, min = 0, max = 1), nrow = 100)
result <- bounding.seq(p.value = X)
Y <- matrix(runif(n = 100, min = 0, max = 1), nrow = 100)
test <- est.prop(p.value = Y, cn = result)</pre>
```

final.selection

Final Selection

Description

Returns the index of final selected variables in the final chosen model.

Usage

```
final.selection(cv.fit, pihat, p, cov.num = 0)
```

Arguments

cv.fit	An object of either class "cv.glmnet" from glmnet::cv.glmnet() or class "cv.ncvreg"
	from nevreg::ev.nevreg(), which is a list generated by a cross-validation fit.
pihat	eatimated proprtion from HCTR::est.prop().
р	Total number of variables, except for covariates.
cov.num	Number of covariates in model, default is 0. Covariate matrix, W, is assumed on the left side of variable matrix, X. The column index of covariates are before those of variables.

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Value

A sequence of index of final selected variables in the final chosen model.

Examples

```
set.seed(10)
X <- matrix(rnorm(20000), nrow = 100)
beta <- rep(0, 200)
beta[1:100] <- 5
Y <- MASS::mvrnorm(n = 1, mu = X%*%beta, Sigma = diag(100))
fit <- glmnet::cv.glmnet(x = X, y = Y)
pihat <- 0.01
result <- est.lambda(cv.fit = fit, pihat = pihat, p = ncol(X))
lambda.seq <- seq(from = result$lambda.min, to = result$lambda.max, length.out = 100)
# Note: The lambda sequences in glmnet and ncvreg are diffrent.
fit2 <- glmnet::cv.glmnet(x = X, y = Y, lambda = lambda.seq)
result2 <- final.selection(cv.fit = fit2, pihat = 0.01, p = ncol(X))</pre>
```

highdim.p

p-values in high-dimensional linear model

Description

Calculates p-values in high-dimentional linear models using multi-split method

Usage

```
highdim.p(Y, X, W = NULL, type, B = 100, fold.num)
```

Arguments

Υ	A numeric response vector, containing nobs variables.
Χ	An input matrix, of dimension nobs x nvars.
W	A covariate matrix, of dimension nobs x ncors, default is NULL.
type	Penalized regression type, valid parameters include "Lasso", "AdaLasso", "SCAD", and "MCP".
В	Multi-split times, default is 100.
fold.num	The number of cross validation folds.

Value

A list of objects containing: (1) harmonic mean p-values; (2) original p-values; (3) index of selected samples; (4) index of selected variables

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Examples

```
set.seed(10)
X <- matrix(rnorm(20000), nrow = 100)
beta <- rep(0, 200)
beta[1:100] <- 5
Y <- MASS::mvrnorm(n = 1, mu = X%*%beta, Sigma = diag(100))
result <- highdim.p(Y=Y, X=X, type = "Lasso", B = 2, fold.num = 10)</pre>
```

multi.adlasso

Multi-split Adaptive Lasso

Description

Multi-splitted variable selection using Adaptive Lasso

Usage

```
multi.adlasso(X, Y, covar.num = NULL, fold.num)
```

Arguments

X An input matrix, of dimension nobs x nvars.

Y A numeric response vector, containing nobs variables.

covar.num Number of covariates in model, default is NULL.

fold.num The number of cross validation folds.

Value

A list of two numeric objects of index of (1) selected and (2) unselected variables.

multi.lasso

Multi-split Lasso

Description

Multi-splitted variable selection using Lasso

Usage

```
multi.lasso(X, Y, p.fac = NULL, fold.num)
```

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Arguments

Y A numeric response vector, containing nobs variables.

p.fac A sequence of penalty factor applied on each variable.

fold.num The number of cross validation folds.

Value

A list of two numeric objects of index of (1) selected and (2) unselected variables.

multi.mcp

Multi-split MCP

Description

Multi-splitted variable selection using MCP

Usage

```
multi.mcp(X, Y, p.fac = NULL, fold.num)
```

Arguments

X An input matrix, of dimension nobs x nvars.

Y A numeric response vector, containing nobs variables.

p. fac A sequence of penalty factor applied on each variable.

fold.num The number of cross validation folds.

Value

A list of two numeric objects of index of (1) selected and (2) unselected variables.

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Description

Multi-splitted variable selection using SCAD

Usage

```
multi.scad(X, Y, p.fac = NULL, fold.num)
```

Arguments

Χ	An input matrix,	of dimension	nobs x nvars.

Y A numeric response vector, containing nobs variables.
p.fac A sequence of penalty factor applied on each variable.

fold.num The number of cross validation folds.

Value

A list of two numeric objects of index of (1) selected and (2) unselected variables.

pmpv	Permutation p-values	

Description

Calculates

Usage

```
pmpv(Y, X, W = NULL, type, B = 100, fold.num = 10, perm.num = 1000)
```

Arguments

Υ	A numeric response vector, containing nobs variables.
Χ	An input matrix, of dimension nobs x nvars.
W	A covariate matrix, of dimension nobs x ncors, default is NULL.
type	Penalized regression type, valid parameters include "Lasso", "AdaLasso", "SCAD", and "MCP".
В	Multi-split times, default is 100.
fold.num	The number of cross validation folds, default is 10.
perm.num	Permutation times, default is 1000.

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Value

A matrix containing harmonic mean p-values from permutation.

Examples

```
set.seed(10)
X <- matrix(rnorm(20000), nrow = 100)
beta <- rep(0, 200)
beta[1:100] <- 5
Y <- MASS::mvrnorm(n = 1, mu = X%*%beta, Sigma = diag(100))
result <- pmpv(Y=Y, X=X, type = "Lasso", B = 2, fold.num = 10, perm.num = 10)</pre>
```

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