

Package ‘grpCox’

July 22, 2025

Type Package

Title Penalized Cox Model for High-Dimensional Data with Grouped Predictors

Version 1.0.2

Date 2023-12-10

Author Xuan Dang

Maintainer Xuan Dang <xuandang11289@gmail.com>

Description Fit the penalized Cox models with both non-overlapping and overlapping grouped penalties including the group lasso, group smoothly clipped absolute deviation, and group mini-max concave penalty. The algorithms combine the MM approach and group-wise descent with some computational tricks including the screening, active set, and warm-start. Different tuning regularization parameter methods are provided.

License GPL (>= 2)

Encoding UTF-8

Imports Rcpp (>= 1.0.11)

LinkingTo Rcpp, RcppEigen

Depends Matrix (>= 1.6-1.1), MASS, colorspace

RoxygenNote 7.2.3

NeedsCompilation yes

Repository CRAN

Date/Publication 2023-12-11 16:00:11 UTC

Contents

grpCox-package	2
cv.grpCox	3
cv.grpCoxOverlap	5
grpCox	7
grpCoxOverlap	9
plot.Coeff	11
plot.gCoeff	12
plot.lLCV	14

grpCox-package	<i>Penalized Cox Model for High-Dimensional Data with Grouped Predictors</i>
----------------	--

Description

Fit the penalized Cox models with both non-overlapping and overlapping grouped penalties including the group lasso, group smoothly clipped absolute deviation, and group minimax concave penalty. The algorithms combine the MM approach and group-wise descent with some computational tricks including the screening, active set, and warm-start. Different tuning regularization parameter methods are provided.

Package Content

Index of help topics:

cv.grpCox	Cross-validation for grpCox
cv.grpCoxOverlap	Cross-validation for grpCoxOverlap
grpCox	Fit a penalized Cox model.
grpCox-package	Penalized Cox Model for High-Dimensional Data with Grouped Predictors
grpCoxOverlap	Fit a penalized regression path with overlapping grouped covariates.
plot.Coeff	Plots the coefficient paths
plot.gCoeff	Plots the coefficient paths with the same color for the covariates in the same group.
plot.llCV	Plot the cross-validation curve produced by cv.grpCox or cv.grpCoxOverlap

Maintainer

Xuan Dang <xuandang11289@gmail.com>

Author(s)

Xuan Dang

cv.grpCox

Cross-validation for grpCox

Description

Does k-fold cross-validation for grpCox

Usage

```
cv.grpCox(X, y, g, m, penalty=c("lasso", "gSCAD", "gMCP"), lambda=NULL,
nlambda=100, rlambda=NULL, gamma=switch(penalty, SCAD = 3.7, 3),
standardize=TRUE, thresh=1e-3, maxit=1e+4, nfolds=10, foldid=NULL)
```

Arguments

X	The design matrix.
y	The response vector includes time corresponding to failure/censor times, and status indicating failure (1) or censoring (0).
g	A vector indicating the group structure of the covariates. It can be unordered groups.
m	Group multipliers. Default is the square root of group size.
penalty	The penalty to be applied to the model. It is one of glasso, gSCAD, or gMCP.
lambda	A user supplied sequence of lambda values. If it is left unspecified, and the function automatically computes a grid of lambda values.
nlambda	The number of lambda values to use in the regularization path. Default is 100.
rlambda	Smallest value for lambda, as a fraction of the maximum lambda, the data derived entry value, i.e. the smallest value for which all coefficients are zero. The default depends on the sample size relative to the number of covariates. If sample size > #covariates, the default is 0.001, close to zero. If sample size < #covariates, the default is 0.05.
gamma	Tuning parameter of the group SCAD/MCP penalty. Default is 3.7 for SCAD and 3 for MCP.
standardize	Logical flag for variable standardization prior to fitting the model.
thresh	Convergence threshold for one-step coordinate descent. Defaults value is 1E-7.
maxit	Maximum number of passes over the data for all lambda values; default is 1E+5.
nfolds	The number of cross-validation folds. Default is 10.
foldid	An optional vector of values between 1 and nfolds identifying what fold each observation is in.

Value

aBetaSTD	A standardized coefficient matrix whose columns correspond to nlambda values of lambda.
aBeta0	A coefficient matrix (without standardization) whose columns correspond to nlambda values of lambda.
mBetaSTD	The coefficient in standardized form gives maximum log-likelihood value using the first cross-validation method.
mBeta0	The coefficient in original form gives maximum log-likelihood value using the first cross-validation method.
pBetaSTD	The coefficient in standardized form gives maximum log-likelihood value using the penalized cross-validation method.
pBeta0	The coefficient in original form gives maximum log-likelihood value using the penalized cross-validation method.
fit	A matrix includes lambda value, the mean cross-validation error.
lambda	The lambda values used.
g	A vector indicating the group structure of the covariates.
cvmax	The maximum value of log likelihood.
lambda.max	The value of lambda corresponds to the maximum value of log likelihood using the first cross-validation method.
lambda.pcv1	The value of lambda corresponds to the maximum value of log likelihood using the penalized cross-validation method.

Author(s)

Xuan Dang <<xuandang11289@gmail.com>>

References

Verweij PJ, Houwelingen HC. Cross-validation in survival analysis. *Statistics in Medicine* 1993; 12(24): 385-395.

Ternes N, Rotolo F, Michiels S. Empirical extensions of the lasso penalty to reduce the false discovery rate in highdimensional Cox regression models. *Statistics in Medicine* 2016; 35(15): 2561-73.

Examples

```
set.seed(200)
N <- 50
p <- 9
x <- matrix(rnorm(N * p), nrow = N)
beta <- c(.65,.65,0,0,.65,.65,0,.65,0)
hx <- exp(x %*% beta)
ty <- rexp(N,hx)
tcens <- 1 - rbinom(n=N, prob = 0.2, size = 1)
y <- data.frame(illt=ty, ills=tcens)
names(y) <- c("time", "status")
```

```

g <- c(1,1,2,2,3,3,2,3,2)
m <- c(sqrt(2),sqrt(4),sqrt(3))

cvfit <- cv.grpCox(x,y,g,m,penalty="glasso")
plot.llCV(cvfit)
plot.gCoef(cvfit$aBeta0, cvfit$g, cvfit$lambda)

```

cv.grpCoxOverlap *Cross-validation for grpCoxOverlap*

Description

Does k-fold cross-validation for grpCoxOverlap

Usage

```

cv.grpCoxOverlap(X0, y, group, penalty=c("glasso", "gSCAD", "gMCP"),
lambda=NULL, nlambda=100, rlambda=NULL, gamma=switch(penalty, SCAD = 3.7, 3),
standardize=TRUE, thresh=1e-3, maxit=1e+4, nfolds=10, foldid=NULL,
returnLatent=TRUE)

```

Arguments

X0	The design matrix.
y	The response vector includes time corresponding to failure/censor times, and status indicating failure (1) or censoring (0).
group	A list of groups, each includes indices of covariates in the group.
penalty	The penalty to be applied to the model. It is one of gLasso, gSCAD, or gMCP.
lambda	A user supplied sequence of lambda values. If it is left unspecified, and the function automatically computes a grid of lambda values.
nlambda	The number of lambda values to use in the regularization path. Default is 100.
rlambda	Smallest value for lambda, as a fraction of the maximum lambda, the data derived entry value, i.e. the smallest value for which all coefficients are zero. The default depends on the sample size relative to the number of covariates. If sample size > #covariates, the default is 0.001, close to zero. If sample size < #covariates, the default is 0.05.
gamma	Tuning parameter of the group SCAD/MCP penalty. Default is 3.7 for SCAD and 3 for MCP.
standardize	Logical flag for variable standardization prior to fitting the model.
thresh	Convergence threshold for one-step coordinate descent. Defaults value is 1E-7.
maxit	Maximum number of passes over the data for all lambda values; default is 1E+5.
nfolds	The number of cross-validation folds. Default is 10.
foldid	An optional vector of values between 1 and nfolds identifying what fold each observation is in.
returnLatent	Return the coefficient matrix in latent space. Default is TRUE.

Value

aBetaLatent	A coefficient matrix whose columns correspond to nlambda values of lambda in latent space.
aBetaOri	A coefficient matrix whose columns correspond to nlambda values of lambda in original space.
mBetaLatent	The coefficient in latent space gives maximum log-likelihood value using the first cross-validation method.
mBetaOri	The coefficient in original space gives maximum log-likelihood value using the first cross-validation method.
pBetaLatent	The coefficient in latent space gives maximum log-likelihood value using the penalized cross-validation method.
pBetaOri	The coefficient in original space gives maximum log-likelihood value using the penalized cross-validation method.
fit	A matrix includes lambda value, the mean cross-validation error.
lambda	The lambda values used.
group	A list of groups, each includes indices of covariates in the group.
glatent	A vector indicating the group structure of the covariates in latent space.
cvmax	The maximum value of log likelihood.
lambda.max	The value of lambda corresponds to the maximum value of log likelihood using the first cross-validation method.
lambda.pcv1	The value of lambda corresponds to the maximum value of log likelihood using the penalized cross-validation method.

Author(s)

Xuan Dang <<xuandang11289@gmail.com>>

References

Verweij PJ, Houwelingen HC. Cross-validation in survival analysis. *Statistics in Medicine* 1993; 12(24): 385-395.

Ternes N, Rotolo F, Michiels S. Empirical extensions of the lasso penalty to reduce the false discovery rate in highdimensional Cox regression models. *Statistics in Medicine* 2016; 35(15): 2561-73.

Examples

```
set.seed(100001)
N <- 50
p <- 6
times <- 1:p
rho <- 0.5
H <- abs(outer(times, times, "-"))
C <- 1 * rho^H
C[cbind(1:p, 1:p)] <- C[cbind(1:p, 1:p)]
sigma <- matrix(C,p,p)
```

```

mu <- rep(0,p)
x <- mvrnorm(n=N, mu, sigma)

beta <- c(0, .8, 1, 2, 1, 0)
hx <- exp(x %*% beta)
ty <- rexp(N,hx)
tcens <- 1 - rbinom(n=N, prob = 0.2, size = 1)
y <- data.frame(illt=ty, ills=tcens)
names(y) <- c("time", "status")

group <- list(g1 = c(1,2,3,4), g2 = c(1,2,6), g3 = c(2,3),
             g4 = c(4,5), g5 = c(5))
cvfit <- cv.grpCoxOverlap(x, y, group, penalty="glasso", nlambda=50)
plot.llCV(cvfit)

```

 grpCox

Fit a penalized Cox model.

Description

Fit the regularization paths for Cox models with grouped covariates.

Usage

```

grpCox(X, y, g, m, penalty=c("glasso", "gSCAD", "gMCP"), lambda=NULL,
nlambda=100, rlambd=NULL, gamma=switch(penalty, gSCAD = 3.7, 3),
standardize=TRUE, thresh=1e-3, maxit=1e+4)

```

Arguments

X	The design matrix.
y	The response vector includes time corresponding to failure/censor times, and status indicating failure (1) or censoring (0).
g	A vector indicating the group structure of the covariates. It can be unordered groups.
m	Group multipliers. Default is the square root of group size.
penalty	The penalty to be applied to the model. It is one of glasso, gSCAD, or gMCP.
lambda	A user supplied sequence of lambda values. If it is left unspecified, and the function automatically computes a grid of lambda values.
nlambda	The number of lambda values to use in the regularization path. Default is 100.
rlambda	Smallest value for lambda, as a fraction of the maximum lambda, the data derived entry value, i.e. the smallest value for which all coefficients are zero. The default depends on the sample size relative to the number of covariates. If sample size > #covariates, the default is 0.001, close to zero. If sample size > #covariates, the default is 0.05.

gamma	Tuning parameter of the group SCAD/MCP penalty. Default is 3.7 for SCAD and 3 for MCP.
standardize	Logical flag for variable standardization prior to fitting the model.
thresh	Convergence threshold for one-step coordinate descent. Defaults value is 1E-7.
maxit	Maximum number of passes over the data for all lambda values; default is 1E+5.

Details

The the group SCAD (gSCAD) and group MCP (gMCP) formulations have been presented in Wang et. al 2007, Huang et. al 2012.

Value

aBetaSTD	A standardized coefficient matrix whose columns correspond to nlambda values of lambda.
aBeta0	A coefficient matrix (without standardization) whose columns correspond to nlambda values of lambda.
lambda	The lambda values used.
ll	The log likelihood values.
g	A vector indicating the group structure of the covariates.

Author(s)

Xuan Dang <<xuandang11289@gmail.com>>

References

Wang, L., Chen, G., and Li, H. Group SCAD regression analysis for microarray time course gene expression data. *Bioinformatics* 23.12 (2007), pp. 1486-1494.

Huang, J., Breheny, P., and Ma, S. A selective review of group selection in high-dimensional models. *Statistical Science* 27.4 (2012), pp. 481-499.

Examples

```
set.seed(200)
N <- 50
p <- 9
x <- matrix(rnorm(N * p), nrow = N)
beta <- c(.65,.65,0,0,.65,.65,0,.65,0)
hx <- exp(x %*% beta)
ty <- rexp(N,hx)
tcens <- 1 - rbinom(n=N, prob = 0.2, size = 1)
y <- data.frame(illt=ty, ills=tcens)
names(y) <- c("time", "status")

g <- c(1,1,2,2,3,3,2,3,2)
m <- c(sqrt(2),sqrt(4),sqrt(3))
```



```
fit <- grpCox(x,y,g,m,penalty="glasso")
plot.gCoef(fit$aBeta0, fit$g, fit$lambda)
```

grpCoxOverlap *Fit a penalized regression path with overlapping grouped covariates.*

Description

Fit the regularization paths for Cox's models with overlapping grouped covariates.

Usage

```
grpCoxOverlap(X0, y, group, penalty=c("glasso", "gSCAD", "gMCP"),
lambda=NULL, nlambda=100, rlambda=NULL, gamma=switch(penalty, gSCAD = 3.7, 3),
standardize = TRUE, thresh=1e-3, maxit=1e+4, returnLatent=TRUE)
```

Arguments

X0	The design matrix.
y	The response vector includes time corresponding to failure/censor times, and status indicating failure (1) or censoring (0).
group	A list of groups, each includes indices of covariates in the group.
penalty	The penalty to be applied to the model. It is one of glasso, gSCAD, or gMCP.
lambda	A user supplied sequence of lambda values. If it is left unspecified, and the function automatically computes a grid of lambda values.
nlambda	The number of lambda values to use in the regularization path. Default is 100.
rlambda	Smallest value for lambda, as a fraction of the maximum lambda, the data derived entry value, i.e. the smallest value for which all coefficients are zero. The default depends on the sample size relative to the number of covariates. If sample size > #covariates, the default is 0.001, close to zero. If sample size < #covariates, the default is 0.05.
gamma	Tuning parameter of the group SCAD/MCP penalty. Default is 3.7 for SCAD and 3 for MCP.
standardize	Logical flag for variable standardization prior to fitting the model.
thresh	Convergence threshold for one-step coordinate descent. Defaults value is 1E-7.
maxit	Maximum number of passes over the data for all lambda values; default is 1E+5.
returnLatent	Return the coefficient matrix in latent space. Default is TRUE.

Details

The the group SCAD (gSCAD) and group MCP (gMCP) formulations have been presented in Wang et. al 2007, Huang et. al 2012.

The method based on the latent group approach (Jacob et al. 2009, Obozinski et al. 2011.)

Value

aBetaLatent	A coefficient matrix whose columns correspond to nlambda values of lambda in latent space.
aBetaOri	A coefficient matrix whose columns correspond to nlambda values of lambda in original space.
lambda	The lambda values used.
ll	The log likelihood values.
group	A list of groups, each includes indices of covariates in the group.
glatent	A vector indicating the group structure of the covariates in latent space.

Author(s)

Xuan Dang <<xuandang11289@gmail.com>>

References

Wang, L., Chen, G., and Li, H. Group SCAD regression analysis for microarray time course gene expression data. *Bioinformatics* 23.12 (2007), pp. 1486-1494.

Huang, J., Breheny, P., and Ma, S. A selective review of group selection in high-dimensional models." *Statistical Science* 27.4 (2012), pp. 481-499.

Jacob, L., Obozinski, G., and Vert, J. P. (2009, June). Group lasso with overlap and graph lasso. In *Proceedings of the 26th annual international conference on machine learning*, ACM: 433-440.

Obozinski, G., Jacob, L., and Vert, J. P. (2011). Group lasso with overlaps: the latent group lasso approach.

Examples

```
set.seed(100001)
N <- 50
p <- 6
times <- 1:p
rho <- 0.5
H <- abs(outer(times, times, "-"))
C <- 1 * rho^H
C[cbind(1:p, 1:p)] <- C[cbind(1:p, 1:p)]
sigma <- matrix(C,p,p)
mu <- rep(0,p)
x <- mvrnorm(n=N, mu, sigma)

beta <- c(0, .8, 1, 2, 1, 0)
hx <- exp(x %*% beta)
ty <- rexp(N,hx)
tcens <- 1 - rbinom(n=N, prob = 0.2, size = 1)
y <- data.frame(illt=ty, ills=tcens)
names(y) <- c("time", "status")

group <- list(g1 = c(1,2,3,4), g2 = c(1,2,6), g3 = c(2,3), g4 = c(4,5), g5 = c(5))
fit <- grpCoxOverlap(x, y, group, penalty="glasso", nlambda=50)
```

```
# plot the coefficient values in latent space
plot.gCoef(fit$aBetaLatent, fit$glatent, fit$lambda)
# plot the coefficient values in original space
plot.Cofef(fit$aBetaOri, fit$lambda)
```

plot.Cofef *Plots the coefficient paths*

Description

Plots the coefficient values as a function of the lambda values used.

Usage

```
## S3 method for class 'Coef'
plot(x, lambda, label=TRUE, xlab="log(Lambda)",
      ylab="Coefficients", title=NULL, ...)
```

Arguments

x	A matrix of coefficients.
lambda	The lambda values used.
label	The indices of covariates. Default is TRUE.
xlab	The name of the x-axis.
ylab	The name of the y-axis.
title	The title of the plot.
...	further arguments to plot

Details

A plot is produced, and nothing is returned.

Value

No return value.

Author(s)

Xuan Dang <<xuandang11289@gmail.com>>

Examples

```

set.seed(100001)
N <- 50
p <- 6
times <- 1:p
rho <- 0.5
H <- abs(outer(times, times, "-"))
C <- 1 * rho^H
C[cbind(1:p, 1:p)] <- C[cbind(1:p, 1:p)]
sigma <- matrix(C,p,p)
mu <- rep(0,p)
x <- mvrnorm(n=N, mu, sigma)

beta <- c(0, .8, 1, 2, 1, 0)
hx <- exp(x %*% beta)
ty <- rexp(N,hx)
tcens <- 1 - rbinom(n=N, prob = 0.2, size = 1)
y <- data.frame(illt=ty, illt=tcens)
names(y) <- c("time", "status")

group <- list(g1 = c(1,2,3,4), g2 = c(1,2,6), g3 = c(2,3), g4 = c(4,5), g5 = c(5))
fit <- grpCoxOverlap(x, y, group, penalty="glasso", nlambda=50)
# plot the coefficient values in latent space
plot.gCoef(fit$betaLatent, fit$glatent, fit$lambda)
# plot the coefficient values in original space
plot.Coeff(fit$betaOri, fit$lambda)

```

plot.gCoef

Plots the coefficient paths with the same color for the covariates in the same group.

Description

Plots the coefficient values as a function of the lambda values used. The covariates in the same group have the same color.

Usage

```

## S3 method for class 'gCoef'
plot(x,g,lambda,label=TRUE,xlab="log(Lambda)",
      ylab="Coefficients", title=NULL,...)

```

Arguments

x	A matrix of coefficients.
g	A vector indicating the group structure of the covariates.
lambda	The lambda values used.
label	The indices of covariates. Default is TRUE.

xlab	The name of the x-axis.
ylab	The name of the y-axis.
title	The title of the plot.
...	further arguments to plot

Details

A plot is produced, and nothing is returned.

Value

No return value.

Author(s)

Xuan Dang <<xuandang11289@gmail.com>>

Examples

```

set.seed(100001)
N <- 50
p <- 6
times <- 1:p
rho <- 0.5
H <- abs(outer(times, times, "-"))
C <- 1 * rho^H
C[cbind(1:p, 1:p)] <- C[cbind(1:p, 1:p)]
sigma <- matrix(C,p,p)
mu <- rep(0,p)
x <- mvrnorm(n=N, mu, sigma)

beta <- c(0, .8, 1, 2, 1, 0)
hx <- exp(x %*% beta)
ty <- rexp(N,hx)
tcens <- 1 - rbinom(n=N, prob = 0.2, size = 1)
y <- data.frame(illt=ty, ills=tcens)
names(y) <- c("time", "status")

group <- list(g1 = c(1,2,3,4), g2 = c(1,2,6), g3 = c(2,3), g4 = c(4,5), g5 = c(5))
fit <- grpCoxOverlap(x, y, group, penalty="glasso", nlambda=50)
# plot the coefficient values in latent space
plot.gCoef(fit$BetaLatent, fit$glatent, fit$lambda)
# plot the coefficient values in original space
plot.Coef(fit$BetaOri, fit$lambda)

```

plot.l1CV	<i>Plot the cross-validation curve produced by cv.grpCox or cv.grpCoxOverlap</i>
-----------	--

Description

Plots the cross-validation curve, and upper and lower standard deviation curves, as a function of the lambda values used.

Usage

```
## S3 method for class 'l1CV'  
plot(x, ...)
```

Arguments

x	fitted cv.grpCox or cv.grpCoxOverlap object
...	further arguments to plot

Details

A plot is produced, and nothing is returned.

Value

No return value.

Author(s)

Xuan Dang <<xuandang11289@gmail.com>>

Examples

```
set.seed(200)  
N <- 50  
p <- 9  
x <- matrix(rnorm(N * p), nrow = N)  
beta <- c(.65,.65,0,0,.65,.65,0,.65,0)  
hx <- exp(x %*% beta)  
ty <- rexp(N,hx)  
tcens <- 1 - rbinom(n=N, prob = 0.2, size = 1)  
y <- data.frame(illt=ty, ills=tcens)  
names(y) <- c("time", "status")  
  
g <- c(1,1,2,2,3,3,2,3,2)  
m <- c(sqrt(2),sqrt(4),sqrt(3))  
  
cvfit <- cv.grpCox(x,y,g,m,penalty="glasso")  
plot.l1CV(cvfit)
```

Index

* Cox models

- cv.grpCox, 3
- cv.grpCoxOverlap, 5
- grpCox, 7
- grpCoxOverlap, 9
- plot.Coeff, 11
- plot.gCoeF, 12
- plot.l1CV, 14

* group regularization

- cv.grpCox, 3
- cv.grpCoxOverlap, 5
- grpCox, 7
- grpCoxOverlap, 9
- plot.Coeff, 11
- plot.gCoeF, 12
- plot.l1CV, 14

* overlapping group

- cv.grpCoxOverlap, 5
- grpCoxOverlap, 9
- plot.Coeff, 11
- plot.gCoeF, 12
- plot.l1CV, 14

* package

- grpCox-package, 2

cv.grpCox, 3

cv.grpCoxOverlap, 5

grpCox, 7

grpCox-package, 2

grpCoxOverlap, 9

plot.Coeff, 11

plot.gCoeF, 12

plot.l1CV, 14