Package 'PanCanVarSel'

October 12, 2022

Title Pan-Cancer Variable Selection
Version 0.0.3
Description Provides function for performing Bayesian survival regression using Horseshoe prior in the accelerated failure time model with log normal assumption in order to achieve high dimensional pan-cancer variable selection as developed in Maity et. al. (2019) <doi:10.1111 biom.13132="">.</doi:10.1111>
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R topics documented:
hsaft 2 hsaftallcorr 4 hsaftcovariatecorr 6 hsaftgroupcorr 9
Index 12

2 hsaft

hsaft	Function to implement the horseshoe shrinkage prior in Bayesian survival regression

Description

This function employs the algorithm provided by van der Pas et. al. (2016) for log normal Accelerated Failure Rate (AFT) model to fit survival regression. The censored observations are updated according to the data augmentation of approach of Tanner and Wong (1984).

Usage

```
hsaft(ct, X, method.tau = c("fixed", "truncatedCauchy", "halfCauchy"),
  tau = 1, method.sigma = c("fixed", "Jeffreys"), Sigma2 = 1,
  burn = 1000, nmc = 5000, thin = 1, alpha = 0.05)
```

Arguments

ct	Response, a $n*2$ matrix with first column as response and second column as right censored indicator, 1 is event time and 0 is right censored.
Χ	Matrix of covariates, dimension $n * p$.
method.tau	Method for handling τ . Select "truncatedCauchy" for full Bayes with the Cauchy prior truncated to [1/p, 1], "halfCauchy" for full Bayes with the half-Cauchy prior, or "fixed" to use a fixed value (an empirical Bayes estimate, for example).
tau	Use this argument to pass the (estimated) value of τ in case "fixed" is selected for method.tau. Not necessary when method.tau is equal to "halfCauchy" or "truncatedCauchy". The default (tau = 1) is not suitable for most purposes and should be replaced.
method.sigma	Select "Jeffreys" for full Bayes with Jeffrey's prior on the error variance σ^2 , or "fixed" to use a fixed value (an empirical Bayes estimate, for example).
Sigma2	A fixed value for the error variance σ^2 . Not necessary when method.sigma is equal to "Jeffreys". Use this argument to pass the (estimated) value of Sigma2 in case "fixed" is selected for method.sigma. The default (Sigma2 = 1) is not suitable for most purposes and should be replaced.
burn	Number of burn-in MCMC samples. Default is 1000.
nmc	Number of posterior draws to be saved. Default is 5000.
thin	Thinning parameter of the chain. Default is 1 (no thinning).
alpha	Level for the credible intervals. For example, alpha = 0.05 results in 95% credible intervals.

Details

The model is: t_i is response, c_i is censored time, $t_i^* = \min_{i} t_i$, $t_i^* = \min_{i} t_i$ is observed time, $t_i^* = \min_{i} t_i$ is observed time, $t_i^* = \min_{i} t_i$ is right censored $t_i^* = t_i^* = t_i^* = t_i^*$ if $t_i^* = t_i^* = t$

hsaft 3

Value

SurvivalHat Predictive survival probability.
LogTimeHat Predictive log time.

BetaHat Posterior mean of Beta, a p by 1 vector. LeftCI The left bounds of the credible intervals. RightCI The right bounds of the credible intervals. BetaMedian Posterior median of Beta, a p by 1 vector.

Sigma2Hat Posterior mean of error variance σ^2 . If method.sigma = "fixed" is used, this

value will be equal to the user-selected value of Sigma2 passed to the function.

TauHat Posterior mean of global scale parameter tau, a positive scalar. If method.tau =

"fixed" is used, this value will be equal to the user-selected value of tau passed

to the function.

BetaSamples Posterior samples of Beta.

TauSamples Posterior samples of tau.

Sigma2Samples Posterior samples of Sigma2.

LikelihoodSamples

Posterior Samples of likelihood.

References

Stephanie van der Pas, James Scott, Antik Chakraborty and Anirban Bhattacharya (2016). horse-shoe: Implementation of the Horseshoe Prior. R package version 0.1.0. https://CRAN.R-project.org/package=horseshoe Arnab Kumar Maity, Anirban Bhattacharya, Bani K. Mallick, and Veerabhadran Baladandayutha-pani (2017). Joint Bayesian Estimation and Variable Selection for TCPA Protein Expression Data

```
burnin <- 500 # number of burnin
       <- 1000 # number of Markov Chain samples
y.sd <- 1
                # standard deviation of the data
       <- 80
                # number of covariates
р
       <- 40
                # number of samples
beta
      <- as.vector(smoothmest::rdoublex(p)) # from double exponential distribution</pre>
       <- mvtnorm::rmvnorm(n, mean = rep(0, p)) # from multivariate normal distribution
y.mu
      <- x %*% beta # mean of the data
      <- as.numeric(stats::rnorm(n, mean = y.mu, sd = y.sd)) # from normal distribution</pre>
У
       <- exp(y) # AFT model
C.
       <- rgamma(n, shape = 1.75, scale = 3) # censoring time</pre>
time
      <- pmin(T, C) # observed time is min of censored and true
status = time == T  # set to 1 if event is observed
       <- as.matrix(cbind(time = time, status = status)) # censored time
posterior.fit <- hsaft(ct, x, method.tau = "truncatedCauchy", method.sigma = "Jeffreys",</pre>
                       burn = burnin, nmc = nmc)
summary(posterior.fit$BetaHat)
```

4 hsaftallcorr

hsaftallcorr	This function extends the main function hsaft to create correlation among covariates.

Description

This function extends the main function hsaft to create correlation among covariates.

Usage

```
hsaftallcorr(ct, X, method.tau = c("fixed", "truncatedCauchy",
   "halfCauchy"), tau = 1, method.sigma = c("fixed", "Jeffreys"),
   Sigma2 = 1, burn = 1000, nmc = 5000, thin = 1, alpha = 0.05, r,
   n.seq, pk)
```

Arguments

ct	Response, a $n*2$ matrix with first column as response and second column as right censored indicator, 1 is event time and 0 is right censored.
Χ	Matrix of covariates, dimension $n * p$.
method.tau	Method for handling τ . Select "truncatedCauchy" for full Bayes with the Cauchy prior truncated to [1/p, 1], "halfCauchy" for full Bayes with the half-Cauchy prior, or "fixed" to use a fixed value (an empirical Bayes estimate, for example).
tau	Use this argument to pass the (estimated) value of τ in case "fixed" is selected for method.tau. Not necessary when method.tau is equal to "halfCauchy" or "truncatedCauchy". The default (tau = 1) is not suitable for most purposes and should be replaced.
method.sigma	Select "Jeffreys" for full Bayes with Jeffrey's prior on the error variance σ^2 , or "fixed" to use a fixed value (an empirical Bayes estimate, for example).
Sigma2	A fixed value for the error variance σ^2 . Not necessary when method.sigma is equal to "Jeffreys". Use this argument to pass the (estimated) value of Sigma2 in case "fixed" is selected for method.sigma. The default (Sigma2 = 1) is not suitable for most purposes and should be replaced.
burn	Number of burn-in MCMC samples. Default is 1000.
nmc	Number of posterior draws to be saved. Default is 5000.
thin	Thinning parameter of the chain. Default is 1 (no thinning).
alpha	Level for the credible intervals. For example, alpha = 0.05 results in 95% credible intervals.
r	number of groups.
n.seq	a vector of sample sizes for all groups.
pk	number of covariates in each group.

hsaftallcorr 5

Value

SurvivalHat

LogTimeHat Predictive log time. BetaHat Posterior mean of Beta, a p by 1 vector. LeftCI The left bounds of the credible intervals. RightCI The right bounds of the credible intervals. BetaMedian Posterior median of Beta, a p by 1 vector. Posterior mean of error variance σ^2 . If method.sigma = "fixed" is used, this Sigma2Hat

Predictive survival probability.

value will be equal to the user-selected value of Sigma2 passed to the function.

TauHat Posterior mean of global scale parameter tau, a positive scalar. If method.tau =

"fixed" is used, this value will be equal to the user-selected value of tau passed

to the function.

BetaSamples Posterior samples of Beta. TauSamples Posterior samples of tau. Sigma2Samples Posterior samples of Sigma2.

BGHat Posterior samples of b which is a part of the mean of β .

BPHat Posterior samples of b which is the other part of the mean of β .

LikelihoodSamples

Posterior Samples of likelihood.

References

Stephanie van der Pas, James Scott, Antik Chakraborty and Anirban Bhattacharya (2016). horseshoe: Implementation of the Horseshoe Prior. R package version 0.1.0. https://CRAN.R-project.org/package=horseshoe

Arnab Kumar Maity, Anirban Bhattacharya, Bani K. Mallick, and Veerabhadran Baladandayuthapani (2017). Joint Bayesian Estimation and Variable Selection for TCPA Protein Expression Data

```
# Examples for hsaftallcorr function
burnin <- 50 # number of burnin
      <- 100 # number of Markov Chain samples
     <- 1 # standard deviation of the data
      <- 80 # number of covariates
      <- 5
            # number of groups
      <- 80 # number of covariate in each group
      <- 40 # sample size of 1st group
n1
      <- 50
             # sample size of 2nd group
n2
      <- 70
n3
               # sample size of 3rd group
      <- 100
              # sample size of 4th group
      <- 120
              # sample size of 5th group
      < sum(c(n1, n2, n3, n4, n5)) # total sample size
n.seq <- c(n1, n2, n3, n4, n5)
      <- matrix(smoothmest::rdoublex(p * r), nrow = r, ncol = p, byrow = TRUE)</pre>
# from double exponential distribution
```

6 hsaftcovariatecorr

```
<- as.vector(t(Beta)) # vectorize Beta</pre>
beta
x1
       <- mvtnorm::rmvnorm(n1, mean = rep(0, p))
       <- mvtnorm::rmvnorm(n2, mean = rep(0, p))</pre>
x2
х3
       <- mvtnorm::rmvnorm(n3, mean = rep(0, p))
       <- mvtnorm::rmvnorm(n4, mean = rep(0, p))
x4
х5
      <- mvtnorm::rmvnorm(n5, mean = rep(0, p)) # from multivariate normal distribution
y.mu1 <- x1 %*% Beta[1, ]
y.mu2 <- x2 %*% Beta[2, ]
y.mu3 <- x3 %*% Beta[3, ]
y.mu4 <- x4 %*% Beta[4, ]
y.mu5 <- x5 %*% Beta[5, ]
       <- stats::rnorm(n1, mean = y.mu1, sd = y.sd)
у1
y2
       <- stats::rnorm(n2, mean = y.mu2, sd = y.sd)
      <- stats::rnorm(n3, mean = y.mu3, sd = y.sd)
y3
y4
      <- stats::rnorm(n4, mean = y.mu4, sd = y.sd)
y5
      <- stats::rnorm(n5, mean = y.mu5, sd = y.sd)
      <- c(y1, y2, y3, y4, y5)
У
      <- Matrix::bdiag(x1, x2, x3, x4, x5)
Χ
      <- as.matrix(x)
       <- as.numeric(as.matrix(y)) # from normal distribution
       <- exp(y) # AFT model
       <- rgamma(n, shape = 1.75, scale = 3) # censoring time</pre>
time \leftarrow pmin(T, C) # observed time is min of censored and true
status = time == T  # set to 1 if event is observed
       <- as.matrix(cbind(time = time, status = status)) # censored time
posterior.fit <- hsaftallcorr(ct, X, method.tau = "truncatedCauchy", method.sigma = "Jeffreys",</pre>
                              burn = burnin, nmc = nmc,
                              r = r, n.seq = n.seq, pk = p)
summary(posterior.fit$BetaHat)
```

hsaftcovariatecorr

This function extends the main function hsaft to create correlation among covariates.

Description

This function extends the main function hsaft to create correlation among covariates.

Usage

```
hsaftcovariatecorr(ct, X, method.tau = c("fixed", "truncatedCauchy",
   "halfCauchy"), tau = 1, method.sigma = c("fixed", "Jeffreys"),
   Sigma2 = 1, burn = 100, nmc = 500, thin = 1, alpha = 0.05, r,
   n.seq, pk)
```

hsaftcovariatecorr 7

Arguments

ct Response, a n * 2 matrix with first column as response and second column as

right censored indicator, 1 is event time and 0 is right censored.

X Matrix of covariates, dimension n * p.

method. tau Method for handling τ . Select "truncatedCauchy" for full Bayes with the Cauchy

prior truncated to [1/p, 1], "halfCauchy" for full Bayes with the half-Cauchy prior, or "fixed" to use a fixed value (an empirical Bayes estimate, for example).

tau Use this argument to pass the (estimated) value of τ in case "fixed" is selected for

method.tau. Not necessary when method.tau is equal to "half Cauchy" or "truncated Cauchy". The default (tau = 1) is not suitable for most purposes and should

be replaced.

method.sigma Select "Jeffreys" for full Bayes with Jeffrey's prior on the error variance σ^2 , or

"fixed" to use a fixed value (an empirical Bayes estimate, for example).

Sigma2 A fixed value for the error variance σ^2 . Not necessary when method sigma is

equal to "Jeffreys". Use this argument to pass the (estimated) value of Sigma2 in case "fixed" is selected for method.sigma. The default (Sigma2 = 1) is not

suitable for most purposes and should be replaced.

burn Number of burn-in MCMC samples. Default is 1000.

nmc Number of posterior draws to be saved. Default is 5000.

thin Thinning parameter of the chain. Default is 1 (no thinning).

alpha Level for the credible intervals. For example, alpha = 0.05 results in 95% credi-

ble intervals.

r number of groups.

n. seq a vector of sample sizes for all groups.pk number of covariates in each group.

Value

SurvivalHat Predictive survival probability.

LogTimeHat Predictive log time.

BetaHat Posterior mean of Beta, a p by 1 vector. LeftCI The left bounds of the credible intervals. RightCI The right bounds of the credible intervals. BetaMedian Posterior median of Beta, a p by 1 vector.

Sigma2Hat Posterior mean of error variance σ^2 . If method.sigma = "fixed" is used, this

value will be equal to the user-selected value of Sigma2 passed to the function.

TauHat Posterior mean of global scale parameter tau, a positive scalar. If method.tau =

"fixed" is used, this value will be equal to the user-selected value of tau passed

to the function.

BetaSamples Posterior samples of Beta.

TauSamples Posterior samples of tau.

8 hsaftcovariatecorr

```
Sigma2Samples Posterior samples of Sigma2. BHat Posterior samples of b which is the mean of \beta. LikelihoodSamples Posterior Samples of likelihood.
```

References

Stephanie van der Pas, James Scott, Antik Chakraborty and Anirban Bhattacharya (2016). horseshoe: Implementation of the Horseshoe Prior. R package version 0.1.0. https://CRAN.R-project.org/package=horseshoe

Arnab Kumar Maity, Anirban Bhattacharya, Bani K. Mallick, and Veerabhadran Baladandayuthapani (2017). Joint Bayesian Estimation and Variable Selection for TCPA Protein Expression Data

```
# Examples for hsaftcovariatecorr function
burnin <- 50 # number of burnin
      <- 100 # number of Markov Chain samples
y.sd <- 1
               # standard deviation of the data
       <- 80 # number of covariates
       <- 5
               # number of groups
             # number of covariate in each group
       <- 80
      <- 40 # sample size of 1st group
n1
      <- 50 # sample size of 2nd group
      <- 70 # sample size of 3rd group
      <- 100 # sample size of 4th group
       <- 120 # sample size of 5th group
       \leftarrow sum(c(n1, n2, n3, n4, n5)) # total sample size
n.seq <- c(n1, n2, n3, n4, n5)
      <- matrix(smoothmest::rdoublex(p * r), nrow = r, ncol = p, byrow = TRUE)</pre>
# from double exponential distribution
beta <- as.vector(t(Beta)) # vectorize Beta</pre>
       <- mvtnorm::rmvnorm(n1, mean = rep(0, p))
x1
       <- mvtnorm::rmvnorm(n2, mean = rep(0, p))
x3
      <- mvtnorm::rmvnorm(n3, mean = rep(0, p))</pre>
      <- mvtnorm::rmvnorm(n4, mean = rep(0, p))</pre>
x4
      <- mvtnorm::rmvnorm(n5, mean = rep(0, p)) # from multivariate normal distribution
x5
y.mu1 <- x1 %*% Beta[1, ]
y.mu2 <- x2 %*% Beta[2, ]
y.mu3 <- x3 %*% Beta[3, ]
y.mu4 <- x4 %*% Beta[4, ]
y.mu5 <- x5 %*% Beta[5, ]
      <- stats::rnorm(n1, mean = y.mu1, sd = y.sd)
у1
      <- stats::rnorm(n2, mean = y.mu2, sd = y.sd)
y2
      <- stats::rnorm(n3, mean = y.mu3, sd = y.sd)
у3
y4
       <- stats::rnorm(n4, mean = y.mu4, sd = y.sd)
у5
      <- stats::rnorm(n5, mean = y.mu5, sd = y.sd)
      <-c(y1, y2, y3, y4, y5)
      <- Matrix::bdiag(x1, x2, x3, x4, x5)
Χ
      <- as.matrix(x)
      <- as.numeric(as.matrix(y)) # from normal distribution
       <- exp(y) # AFT model
```

hsaftgroupcorr 9

hsaftgroupcorr

This function extends the main function hsaft to create correlation among groups.

Description

This function extends the main function hsaft to create correlation among groups.

Usage

```
hsaftgroupcorr(ct, X, method.tau = c("fixed", "truncatedCauchy", "halfCauchy"), tau = 1, method.sigma = c("fixed", "Jeffreys"), Sigma2 = 1, burn = 1000, nmc = 5000, thin = 1, alpha = 0.05, r, n.seq, pk)
```

Arguments

ct	Response, a $n * 2$ matrix with first column as response and second column as
	right censored indicator, 1 is event time and 0 is right censored.

X Matrix of covariates, dimension n * p.

method. tau Method for handling τ . Select "truncatedCauchy" for full Bayes with the Cauchy

prior truncated to [1/p, 1], "halfCauchy" for full Bayes with the half-Cauchy prior, or "fixed" to use a fixed value (an empirical Bayes estimate, for example).

tau Use this argument to pass the (estimated) value of τ in case "fixed" is selected for

method.tau. Not necessary when method.tau is equal to "halfCauchy" or "truncatedCauchy". The default (tau = 1) is not suitable for most purposes and should

be replaced.

method.sigma Select "Jeffreys" for full Bayes with Jeffrey's prior on the error variance σ^2 , or

"fixed" to use a fixed value (an empirical Bayes estimate, for example).

Sigma2 A fixed value for the error variance σ^2 . Not necessary when method sigma is

equal to "Jeffreys". Use this argument to pass the (estimated) value of Sigma2 in case "fixed" is selected for method.sigma. The default (Sigma2 = 1) is not

suitable for most purposes and should be replaced.

10 hsaftgroupcorr

burn Number of burn-in MCMC samples. Default is 1000.

nmc Number of posterior draws to be saved. Default is 5000.

thin Thinning parameter of the chain. Default is 1 (no thinning).

alpha Level for the credible intervals. For example, alpha = 0.05 results in 95% credi-

ble intervals.

r number of groups.

n. seq a vector of sample sizes for all groups.pk number of covariates in each group.

Value

SurvivalHat Predictive survival probability.

LogTimeHat Predictive log time.

BetaHat Posterior mean of Beta, a p by 1 vector. LeftCI The left bounds of the credible intervals. RightCI The right bounds of the credible intervals. BetaMedian Posterior median of Beta, a p by 1 vector.

Sigma2Hat Posterior mean of error variance σ^2 . If method.sigma = "fixed" is used, this

value will be equal to the user-selected value of Sigma2 passed to the function.

TauHat Posterior mean of global scale parameter tau, a positive scalar. If method.tau =

"fixed" is used, this value will be equal to the user-selected value of tau passed

to the function.

BetaSamples Posterior samples of Beta.

TauSamples Posterior samples of tau.

Sigma2Samples Posterior samples of Sigma2.

BHat Posterior samples of b which is the mean of β .

LikelihoodSamples

Posterior Samples of likelihood.

References

Stephanie van der Pas, James Scott, Antik Chakraborty and Anirban Bhattacharya (2016). horseshoe: Implementation of the Horseshoe Prior. R package version 0.1.0. https://CRAN.R-project.org/package=horseshoe

Arnab Kumar Maity, Anirban Bhattacharya, Bani K. Mallick, and Veerabhadran Baladandayutha-pani (2017). Joint Bayesian Estimation and Variable Selection for TCPA Protein Expression Data

hsaftgroupcorr 11

```
<- 80
              # number of covariate in each group
              # sample size of 1st group
       <- 40
      <- 50 # sample size of 2nd group
      <- 70 # sample size of 3rd group
      <- 100 # sample size of 4th group
       <- 120 # sample size of 5th group
       < sum(c(n1, n2, n3, n4, n5)) # total sample size
n.seq <- c(n1, n2, n3, n4, n5)
Beta <- matrix(smoothmest::rdoublex(p * r), nrow = r, ncol = p, byrow = TRUE)</pre>
# from double exponential distribution
beta <- as.vector(t(Beta)) # vectorize Beta</pre>
       <- mvtnorm::rmvnorm(n1, mean = rep(0, p))</pre>
x1
x2
       <- mvtnorm::rmvnorm(n2, mean = rep(0, p))
      <- mvtnorm::rmvnorm(n3, mean = rep(0, p))</pre>
х3
x4
      <- mvtnorm::rmvnorm(n4, mean = rep(0, p))</pre>
x5
      <- mvtnorm::rmvnorm(n5, mean = rep(0, p)) # from multivariate normal distribution
y.mu1 <- x1 %*% Beta[1, ]
y.mu2 <- x2 %*% Beta[2, ]
y.mu3 <- x3 %*% Beta[3, ]
y.mu4 <- x4 %*% Beta[4, ]
y.mu5 <- x5 %*% Beta[5, ]
y1
      <- stats::rnorm(n1, mean = y.mu1, sd = y.sd)
      <- stats::rnorm(n2, mean = y.mu2, sd = y.sd)</pre>
y2
      <- stats::rnorm(n3, mean = y.mu3, sd = y.sd)
y3
      <- stats::rnorm(n4, mean = y.mu4, sd = y.sd)</pre>
y4
      <- stats::rnorm(n5, mean = y.mu5, sd = y.sd)
у5
      <- c(y1, y2, y3, y4, y5)
У
      <- Matrix::bdiag(x1, x2, x3, x4, x5)
Χ
      <- as.matrix(x)
      <- as.numeric(as.matrix(y)) # from normal distribution
      <- exp(y) # AFT model
C
       <- rgamma(n, shape = 1.75, scale = 3) # censoring time</pre>
time \leftarrow pmin(T, C) # observed time is min of censored and true
status = time == T  # set to 1 if event is observed
       <- as.matrix(cbind(time = time, status = status)) # censored time
posterior.fit <- hsaftgroupcorr(ct, X, method.tau = "truncatedCauchy", method.sigma = "Jeffreys",</pre>
                                burn = burnin, nmc = nmc,
                                r = r, n.seq = n.seq, pk = p)
summary(posterior.fit$BetaHat)
```

Index

```
hsaft, 2, 4, 6, 9
hsaftallcorr, 4
hsaftcovariatecorr, 6
hsaftgroupcorr, 9
```