

Package: changepoints (via r-universe)

November 1, 2024

Type Package

Title A Collection of Change-Point Detection Methods

Version 1.1.0

Date 2022-12-29

Maintainer Haotian Xu <haotian.xu@uclouvain.be>

Description Performs a series of offline and/or online change-point detection algorithms for 1) univariate mean: <doi:10.1214/20-EJS1710>, <arXiv:2006.03283>; 2) univariate polynomials: <doi:10.1214/21-EJS1963>; 3) univariate and multivariate nonparametric settings: <doi:10.1214/21-EJS1809>, <doi:10.1109/TIT.2021.3130330>; 4) high-dimensional covariances: <doi:10.3150/20-BEJ1249>; 5) high-dimensional networks with and without missing values: <doi:10.1214/20-AOS1953>, <arXiv:2101.05477>, <arXiv:2110.06450>; 6) high-dimensional linear regression models: <arXiv:2010.10410>, <arXiv:2207.12453>; 7) high-dimensional vector autoregressive models: <arXiv:1909.06359>; 8) high-dimensional self exciting point processes: <arXiv:2006.03572>; 9) dependent dynamic nonparametric random dot product graphs: <arXiv:1911.07494>; 10) univariate mean against adversarial attacks: <arXiv:2105.10417>.

Depends R (>= 3.5.0)

Imports stats, gglasso, glmnet, MASS, Rcpp, data.tree, cubature

Suggests knitr, abind, rmarkdown

LinkingTo Rcpp, RcppArmadillo

License GPL (>= 3)

RoxygenNote 7.2.3

Encoding UTF-8

URL <https://github.com/HaotianXu/changepoints>

VignetteBuilder knitr

Repository <https://haotianxu.r-universe.dev>
RemoteUrl <https://github.com/haotianxu/changepoints>
RemoteRef HEAD
RemoteSha 002b97ea4e29b38dbd55ccfcbbf24ffecfd9532

Contents

aARC	3
ARC	5
BD_U	6
BS.cov	7
BS.uni.nonpar	8
BS.univar	9
calibrate.online.network.missing	10
changepoints	12
CI.regression	13
CV.search.DP.LR.regression	15
CV.search.DP.poly	16
CV.search.DP.regression	17
CV.search.DP.univar	19
CV.search.DP.VAR1	20
CV.search.DPDU.regression	21
DP.poly	22
DP.regression	23
DP.SEPP	24
DP.univar	26
DP.VAR1	27
DPDU.regression	28
DPDU2.regression	29
gen.cov.mat	30
gen.missing	31
gen.piece.poly	31
gen.piece.poly.noiseless	32
Hausdorff.dist	33
huber_mean	34
kde.biwei.eval	35
kde.epan.eval	35
kde.eval	36
kde.triwei.eval	37
lambda.network.missing	38
local.refine.CV.VAR1	39
local.refine.DPDU.regression	40
local.refine.multi.nonpar.L2	41
local.refine.multi.nonpar.L2.biwei	42
local.refine.multi.nonpar.L2.epan	44
local.refine.multi.nonpar.L2.triwei	45
local.refine.network	47

local.refine.poly	48
local.refine.regression	49
local.refine.univar	50
local.refine.VAR1	51
lowertri2mat	52
LRV.regression	53
online.network	54
online.network.missing	55
online.univar	57
online.univar.multi	58
simu.change.regression	59
simu.RDPG	61
simu.SBM	62
simu.SEPP	63
simu.VAR1	64
softImpute.network.missing	65
thresholdBS	66
trim_interval	67
tuneBSmultinonpar	68
tuneBSnonparRDPG	69
tuneBSnonpar	70
tuneBSunivar	71
WBS.intervals	72
WBS.multi.nonpar	73
WBS.multi.nonpar.L2	74
WBS.multi.nonpar.L2.biwei	76
WBS.multi.nonpar.L2.epan	77
WBS.multi.nonpar.L2.triwei	79
WBS.network	80
WBS.nonpar.RDPG	82
WBS.uni.nonpar	83
WBS.uni.rob	85
WBS.univar	86
WBSIP.cov	87

Index **89**

aARC	<i>Automatic adversarially robust univariate mean change point detection.</i>
------	---

Description

Perform the adversarially robust change point detection method with automatic selection of the contamination proportion epsilon when treating the inliner distributions as Gaussian.

Usage

```
aARC(y, t_dat, guess_true = 0.05, h, block_num = 1)
```

Arguments

y	A numeric vector of observations.
t_dat	A numeric vector of observations that is used to select epsilon in the Huber contamination model.
guess_true	A numeric scalar representing a guess of epsilon value.
h	An integer scalar representing block size.
block_num	An integer scalar representing number of blocks used when searching for local maximiser.

Value

An numeric vector of estimated change point locations.

Author(s)

Mengchu Li

References

Li and Yu (2021) <arXiv:2105.10417>.

Examples

```
##' ### simulate data with contamination
obs_num = 1000
D = 2
noise = 0.1 # proportion of contamination
mu0 = 0
mu1 = 1
sd = 1
idmixture = rbinom(obs_num/D, 1, 1-noise)
dat = NULL
for (j in 1:D){
  for (i in 1:(obs_num/(2*D))){
    if (idmixture[i] == 1){
      dat = c(dat,rnorm(1,mu0,sd))
    }
    else{
      dat = c(dat,rnorm(1,mu1/(2*noise),0))
    }
  }
}
for (i in (obs_num/(2*D)+1):(obs_num/D)){
  if (idmixture[i] == 1){
    dat = c(dat,rnorm(1,mu1,sd))
  }
  else{
```

```
        dat = c(dat,rnorm(1,mu1/(2*noise)-(1-noise)*mu1/noise,0))
      }
    }
  }
plot(dat)
### perform aARC
aARC(dat, dat[1:200], h = 120)
```

ARC

Adversarially robust univariate mean change point detection.

Description

Perform the adversarially robust change point detection method.

Usage

```
ARC(y, h, block_num = 1, epsilon, gaussian = TRUE)
```

Arguments

<code>y</code>	A numeric vector of observations.
<code>h</code>	An integer scalar representing block size.
<code>block_num</code>	An integer scalar representing number of blocks used when searching for local maximiser.
<code>epsilon</code>	A numeric scalar in (0,1) representing contamination proportion.
<code>gaussian</code>	A logical scalar representing whether to treat the inlier distribution (F) as Gaussian distribution.

Value

An numeric vector of estimated change point locations

Author(s)

Mengchu Li

References

Li and Yu (2021) <arXiv:2105.10417>.

Examples

```

### simulate data with contamination
obs_num = 1000
D = 2
noise = 0.1 # proportion of contamination
mu0 = 0
mu1 = 1
sd = 1
idmixture = rbinom(obs_num/D, 1, 1-noise)
dat = NULL
for (j in 1:D){
  for (i in 1:(obs_num/(2*D))){
    if (idmixture[i] == 1){
      dat = c(dat,rnorm(1,mu0,sd))
    }
    else{
      dat = c(dat,rnorm(1,mu1/(2*noise),0))
    }
  }
  for (i in (obs_num/(2*D)+1):(obs_num/D)){
    if (idmixture[i] == 1){
      dat = c(dat,rnorm(1,mu1,sd))
    }
    else{
      dat = c(dat,rnorm(1,mu1/(2*noise)-(1-noise)*mu1/noise,0))
    }
  }
}
plot(dat)
### perform ARC
ARC(dat,h = 120, epsilon = 0.1)

```

BD_U

Backward detection with a robust bootstrap change point test using U-statistics for univariate mean change.

Description

Perform the backward detection method with a robust bootstrap change point test using U-statistics on univariate data.

Usage

```
BD_U(y, M, B = 100, inter = NULL, inter_ini = TRUE)
```

Arguments

y A numeric vector of observations.

M	An integer scalar representing initial block size of the backward detection algorithm.
B	An integer scalar representing the number of bootstrapped samples.
inter	A nuisance parameter.
inter_ini	A nuisance parameter.

Value

An numeric vector of estimated change point locations.

Author(s)

Mengchu Li

References

Yu and Chen (2019) <arXiv:1904.03372>.

BS.cov	<i>Binary Segmentation for covariance change points detection through Operator Norm.</i>
--------	--

Description

Perform binary segmentation for covariance change points detection through Operator Norm.

Usage

```
BS.cov(X, s, e, level = 0)
```

Arguments

X	A numeric matrix of observations with with horizontal axis being time, and vertical axis being dimensions.
s	A integer scalar of starting index.
e	A integer scalar of ending index.
level	A parameter for tracking the level at which a change point is detected. Should be fixed as 0.

Value

An object of `class` "BS", which is a list with the structure:

- S: A vector of estimated changepoints (sorted in strictly increasing order).
- Dval: A vector of values of CUSUM statistic based on KS distance.
- Level: A vector representing the levels at which each change point is detected.
- Parent: A matrix with the starting indices on the first row and the ending indices on the second row.

Author(s)

Haotian Xu

References

Wang, Yu and Rinaldo (2021) <doi:10.3150/20-BEJ1249>.

See Also[thresholdBS](#) for obtain change points estimation.**Examples**

```

p = 10
A1 = gen.cov.mat(p, 1, "equal")
A2 = gen.cov.mat(p, 2, "diagonal")
A3 = gen.cov.mat(p, 3, "power")
X = cbind(t(MASS::mvrnorm(100, mu = rep(0, p), A1)),
          t(MASS::mvrnorm(150, mu = rep(0, p), A2)),
          t(MASS::mvrnorm(200, mu = rep(0, p), A3)))
temp = BS.cov(X, 1, 450)
thresholdBS(temp, 10)

```

BS.uni.nonpar

Standard binary segmentation for univariate nonparametric change points detection.

Description

Perform standard binary segmentation for univariate nonparametric change points detection.

Usage

BS.uni.nonpar(Y, s, e, N, delta, level = 0)

Arguments

Y	A numeric matrix of observations with horizontal axis being time, and vertical axis being multiple observations on each time point.
s	A integer scalar of starting index.
e	A integer scalar of ending index.
N	A integer vector representing number of multiple observations on each time point.
delta	A positive integer scalar of minimum spacing.
level	Should be fixed as 0.

Value

An object of class "BS", which is a list with the following structure:

S	A vector of estimated change points (sorted in strictly increasing order).
Dval	A vector of values of CUSUM statistic based on KS distance.
Level	A vector representing the levels at which each change point is detected.
Parent	A matrix with the starting indices on the first row and the ending indices on the second row.

Author(s)

Oscar Hernan Madrid Padilla & Haotian Xu

References

Padilla, Yu, Wang and Rinaldo (2021) <doi:10.1214/21-EJS1809>.

See Also

[thresholdBS](#) for obtaining change points estimation, [tuneBSuninonpar](#) for a tuning version.

Examples

```
Y = t(as.matrix(c(rnorm(100, 0, 1), rnorm(100, 0, 10), rnorm(100, 0, 40))))
N = rep(1, 300)
temp = BS.uni.nonpar(Y, 1, 300, N, 5)
plot.ts(t(Y))
points(x = tail(temp$S[order(temp$Dval)],4), y = Y[,tail(temp$S[order(temp$Dval)],4)], col = "red")
```

BS.univar	<i>Standard binary segmentation for univariate mean change points detection.</i>
-----------	--

Description

Perform standard binary segmentation for univariate mean change points detection.

Usage

```
BS.univar(y, s, e, delta = 2, level = 0)
```

Arguments

y	A numeric vector of observations.
s	A integer scalar of starting index.
e	A integer scalar of ending index.
delta	A positive numeric scalar of minimum spacing.
level	Should be fixed as 0.

Value

An object of class "BS", which is a list with the following structure:

S	A vector of estimated change point locations (sorted in strictly increasing order).
Dval	A vector of values of CUSUM statistic.
Level	A vector representing the levels at which each change point is detected.
Parent	A matrix with the starting indices on the first row and the ending indices on the second row.

Author(s)

Haotian Xu

References

Wang, Yu and Rinaldo (2020) <doi:10.1214/20-EJS1710>.

See Also

[thresholdBS](#) for obtaining change points estimation, [tuneBSunivar](#) for a tuning version.

Examples

```
set.seed(0)
cpt_true = c(20, 50, 170)
y = rnorm(300) + c(rep(0,20),rep(2,30),rep(0,120),rep(2,130))
temp = BS.univar(y, 1, length(y), delta = 5)
plot.ts(y)
points(x = tail(temp$S[order(temp$Dval)],4),
       y = y[tail(temp$S[order(temp$Dval)],4)], col = "red")
BS_result = thresholdBS(temp, tau = 4)
BS_result
print(BS_result$BS_tree, "value")
print(BS_result$BS_tree_trimmed, "value")
cpt_hat = sort(BS_result$cpt_hat[,1]) # the threshold tau is specified to be 4
Hausdorff.dist(cpt_hat, cpt_true)
cpt_LR = local.refine.univar(cpt_hat, y)
Hausdorff.dist(cpt_LR, cpt_true)
```

calibrate.online.network.missing

Calibrate step for online change point detection for network data with missing values.

Description

Calibrate step for online change point detection for network data by controlling the false alarm rate at level alpha.

Usage

```
calibrate.online.network.missing(
  train_miss_list,
  train_eta_list,
  threshold_len,
  alpha_grid,
  permu_num,
  pi_lb_hat,
  pi_ub_hat,
  rho_hat,
  rank_hat,
  C_lambda = 2/3,
  delta = 5
)
```

Arguments

<code>train_miss_list</code>	A list of adjacency matrices (with entries being 0 or 1) with missing values being coercing to 0.
<code>train_eta_list</code>	A list of matrices associated with <code>data_incomplete_list</code> , each matrix indicates the missing entries in corresponding adjacency matrix.
<code>threshold_len</code>	An integer scalar of the length of tuned thresholds.
<code>alpha_grid</code>	A numeric vector in (0,1) representing the desired false alarm rate.
<code>permu_num</code>	An integer scalar of number of random permutation for calibration.
<code>pi_lb_hat</code>	A numeric scalar of the lower bound of the missing probability.
<code>pi_ub_hat</code>	A numeric scalar of the upper bound of the missing probability.
<code>rho_hat</code>	A numeric scalar of the sparsity parameter.
<code>rank_hat</code>	An integer scalar of the rank of the underlying graphon matrix.
<code>C_lambda</code>	A numeric scalar of an absolute constant, which is set to be 2/3 by default.
<code>delta</code>	An integer scalar of minimum spacing.

Value

A list with the following structure:

<code>C_lambda</code>	The absolute constant
<code>rho_hat</code>	the (estimated) sparsity parameter
<code>rank_hat</code>	the (estimated) rank of underlying graphon matrix
<code>pi_lb_hat</code>	the (estimated) lower bound of the missing probability
<code>pi_ub_hat</code>	the (estimated) upper bound of the missing probability
<code>thresholds_array</code>	A numeric array of calibrated threshold

Author(s)

Haotian Xu

References

Dubey, Xu and Yu (2021) <arxiv:2110.06450>

See Also[online.network.missing](#) for detecting online change point.**Examples**

```

p = 6 # number of nodes
rho = 0.5 # sparsity parameter
block_num = 3 # number of groups for SBM
train_obs_num = 150 # sample size for each segment
conn1_mat = rho * matrix(c(0.6,1,0.6,1,0.6,0.5,0.6,0.5,0.6), nrow = 3) # connectivity matrix
set.seed(1)
can_vec = sample(1:p, replace = FALSE) # randomly assign nodes into groups
sbm = simu.SBM(conn1_mat, can_vec, train_obs_num, symm = TRUE, self = TRUE)
train_mat = sbm$obs_mat
train_list = lapply(1:ncol(train_mat), function(t) lowertri2mat(train_mat[,t], p, diag = TRUE))
pi_mat = matrix(0.9, p, p)
train_eta_list = lapply(1:length(train_list), function(t) gen.missing(pi_mat, symm = TRUE))
train_miss_list = lapply(1:length(train_list), function(t) train_eta_list[[t]] * train_list[[t]])
pi_lb_hat = quantile(Reduce("+", train_eta_list)/train_obs_num, 0.05) # estimator of pi_lb
pi_ub_hat = quantile(Reduce("+", train_eta_list)/train_obs_num, 0.95) # estimator of pi_ub
C_lambda = 2/3
lambda = lambda.network.missing(1, length(train_miss_list), length(train_miss_list), 0.05,
                                rho = 0.509, pi_ub = pi_ub_hat, p, C_lambda)
graphon_miss_impute = softImpute.network.missing(train_miss_list, train_eta_list, lambda, 1)
graphon_miss_hat = graphon_miss_impute$u %*% diag(as.numeric(graphon_miss_impute$d)) %*%
                    t(graphon_miss_impute$v)
rho_hat = quantile(graphon_miss_hat, 0.95)
rank_hat = sum(graphon_miss_impute$d != 0)
alpha_grid = c(0.05)
permu_num = 10
threshold_len = 30
temp = calibrate.online.network.missing(train_miss_list, train_eta_list, threshold_len, alpha_grid,
                                       permu_num, pi_lb_hat, pi_ub_hat, rho_hat, rank_hat, C_lambda, delta = 5)

```

Description

Performs a series of offline and/or online change-point detection algorithms for 1) univariate mean; 2) univariate polynomials; 3) univariate and multivariate nonparametric settings; 4) high-dimensional covariances; 5) high-dimensional networks with and without missing values; 6) high-dimensional linear regression models; 7) high-dimensional vector autoregressive models; 8) high-dimensional self exciting point processes; 9) dependent dynamic nonparametric random dot product graphs; 10) univariate mean against adversarial attacks. For more informations, see Wang et al. (2020) <arXiv:1810.09498>; Yu et al. (2020) <arXiv:2006.03283>; Yu and Chatterjee (2020) <arXiv:2007.09910>; Padilla et al. (2021) <arXiv:1905.10019>; Padilla et al. (2019) <arXiv:1910.13289>; Wang et al. (2021) <arXiv:1712.09912>; Wang et al. (2018) <arXiv:1809.09602>; Padilla et al. (2019) <arXiv:1911.07494>; Yu et al. (2021) <arXiv:2101.05477>; Rinaldo et al. (2020) <arXiv:2010.10410>; Wang et al. (2019) <arXiv:1909.06359>; Wang et al. (2020) <arXiv:2006.03572>; Dubey et al. (2021) <arXiv:2110.06450>; Li and Yu (2021) <arXiv:2105.10417>.

CI.regression	<i>Confidence interval construction of change points for regression settings with change points.</i>
---------------	--

Description

Construct element-wise confidence interval for change points.

Usage

```
CI.regression(
  cpt_init,
  cpt_LR,
  beta_hat,
  y,
  X,
  w = 0.9,
  B = 1000,
  M,
  alpha_vec,
  rounding = TRUE
)
```

Arguments

<code>cpt_init</code>	An integer vector of initial changepoints estimation (sorted in strictly increasing order).
<code>cpt_LR</code>	An integer vector of refined changepoints estimation (sorted in strictly increasing order).
<code>beta_hat</code>	A numeric ($p \times (K_{\text{hat}}+1)$) matrix of estimated regression coefficients.
<code>y</code>	A numeric vector of response variable.

X	A numeric matrix of covariates with vertical axis being time.
w	A numeric scalar in (0,1) representing the weight for interval truncation.
B	An integer scalar corresponding to the number of simulated two-sided Brownian motion with drift.
M	An integer scalar corresponding to the length for each side of the limiting distribution, i.e. the two-sided Brownian motion with drift.
alpha_vec	An numeric vector in (0,1) representing the vector of significance levels.
rounding	A boolean scalar representing if the confidence intervals need to be rounded into integer intervals.

Value

An $\text{length}(\text{cpt_init}) - 2 - \text{length}(\text{alpha_vec})$ array of confidence intervals.

Author(s)

Haotian Xu

References

Xu, Wang, Zhao and Yu (2022) <arXiv:2207.12453>.

Xu, Wang, Zhao and Yu (2022) <arXiv:2207.12453>.

Examples

```
d0 = 5
p = 10
n = 200
cpt_true = c(70, 140)
data = simu.change.regression(d0, cpt_true, p, n, sigma = 1, kappa = 9)
lambda_set = c(0.1, 0.5, 1, 2)
zeta_set = c(10, 15, 20)
temp = CV.search.DPDU.regression(y = data$y, X = data$X, lambda_set, zeta_set)
temp$test_error # test error result
# find the indices of lambda_set and zeta_set which minimizes the test error
min_idx = as.vector(arrayInd(which.min(temp$test_error), dim(temp$test_error)))
lambda_set[min_idx[2]]
zeta_set[min_idx[1]]
cpt_init = unlist(temp$cpt_hat[min_idx[1], min_idx[2]])
beta_hat = matrix(unlist(temp$beta_hat[min_idx[1], min_idx[2]]), ncol = length(cpt_init)+1)
cpt_LR = local.refine.DPDU.regression(cpt_init, beta_hat, data$y, data$X, w = 0.9)
alpha_vec = c(0.01, 0.05, 0.1)
CI.regression(cpt_init, cpt_LR, beta_hat, data$y, data$X, w = 0.9, B = 1000, M = n, alpha_vec)
```

 CV.search.DP.LR.regression

Grid search based on Cross-Validation of all tuning parameters (gamma, lambda and zeta) for regression.

Description

Perform grid search based on Cross-Validation of all tuning parameters (gamma, lambda and zeta)

Usage

```
CV.search.DP.LR.regression(
  y,
  X,
  gamma_set,
  lambda_set,
  zeta_set,
  delta,
  eps = 0.001
)
```

Arguments

y	A numeric vector of response variable.
X	A numeric matrix of covariates with vertical axis being time.
gamma_set	A numeric vector of candidate tuning parameter associated with the l0 penalty.
lambda_set	A numeric vector of candidate tuning parameter for the lasso penalty.
zeta_set	A numeric vector of candidate tuning parameter for the group lasso.
delta	A strictly positive integer scalar of minimum spacing.
eps	A numeric scalar of precision level for convergence of lasso.

Value

A list with the following structure:

cpt_hat	A list of vector of estimated changepoints (sorted in strictly increasing order)
K_hat	A list of scalar of number of estimated changepoints
test_error	A list of vector of testing errors (each row corresponding to each gamma, and each column corresponding to each lambda)
train_error	A list of vector of training errors

Author(s)

Daren Wang & Haotian Xu

References

Rinaldo, Wang, Wen, Willett and Yu (2020) <arxiv:2010.10410>

Examples

```
set.seed(123)
d0 = 8
p = 15
n = 100
cpt_true = c(30, 70)
data = simu.change.regression(d0, cpt_true, p, n, sigma = 1, kappa = 9)
gamma_set = c(0.01, 0.1)
lambda_set = c(0.01, 0.1)
temp = CV.search.DP.regression(y = data$y, X = data$X, gamma_set, lambda_set, delta = 2)
temp$test_error # test error result
# find the indices of gamma_set and lambda_set which minimizes the test error
min_idx = as.vector(arrayInd(which.min(temp$test_error), dim(temp$test_error)))
gamma_set[min_idx[1]]
lambda_set[min_idx[2]]
cpt_init = unlist(temp$cpt_hat[min_idx[1], min_idx[2]])
zeta_set = c(0.1, 1)
temp_zeta = CV.search.DP.LR.regression(data$y, data$X, gamma_set[min_idx[1]],
                                     lambda_set[min_idx[2]], zeta_set, delta = 2, eps = 0.001)
min_zeta_idx = which.min(unlist(temp_zeta$test_error))
cpt_LR = local.refine.regression(cpt_init, data$y, X = data$X, zeta = zeta_set[min_zeta_idx])
Hausdorff.dist(cpt_init, cpt_true)
Hausdorff.dist(cpt_LR, cpt_true)
```

CV.search.DP.poly	<i>Grid search for dynamic programming to select the tuning parameter through Cross-Validation.</i>
-------------------	---

Description

Perform grid search for dynamic programming to select the tuning parameter through Cross-Validation.

Usage

```
CV.search.DP.poly(y, r, gamma_set, delta)
```

Arguments

y	A numeric vector of observations.
r	An integer scalar order of polynomials.
gamma_set	A numeric vector of candidate tuning parameter associated with the l0 penalty.
delta	A positive integer scalar of minimum spacing.

Value

A list with the following structure:

cpt_hat	A list of vector of estimated change points locations (sorted in strictly increasing order)
K_hat	A list of scalar of number of estimated change points
test_error	A list of vector of testing errors
train_error	A list of vector of training errors

Author(s)

Haotian Xu

References

Yu and Chatterjee (2020) <arXiv:2007.09910>

Examples

```
set.seed(0)
cpt_true = c(20, 50, 170)
y = rnorm(300) + c(rep(0,20),rep(2,30),rep(0,120),rep(2,130))
plot.ts(y)
gamma_set = 3:9
DP_result = CV.search.DP.poly(y, r = 2, gamma_set, delta = 5)
min_idx = which.min(DP_result$test_error)
cpt_init = unlist(DP_result$cpt_hat[min_idx])
local.refine.poly(cpt_init, y, r = 2, delta_lr = 5)
```

CV.search.DP.regression

Grid search based on cross-validation of dynamic programming for regression change points localisation with l_0 penalisation.

Description

Perform grid search to select tuning parameters gamma (for l_0 penalty of DP) and lambda (for lasso penalty) based on cross-validation.

Usage

```
CV.search.DP.regression(y, X, gamma_set, lambda_set, delta, eps = 0.001)
```

Arguments

y	A numeric vector of response variable.
X	A numeric matrix of covariates with vertical axis being time.
gamma_set	A numeric vector of candidate tuning parameters associated with l_0 penalty of DP.
lambda_set	A numeric vector of candidate tuning parameters for lasso penalty.
delta	A strictly positive integer scalar of minimum spacing.
eps	A numeric scalar of precision level for convergence of lasso.

Value

A list with the following structure:

cpt_hat	A list of vector of estimated change points
K_hat	A list of scalar of number of estimated change points
test_error	A list of vector of testing errors (each row corresponding to each gamma, and each column corresponding to each lambda)
train_error	A list of vector of training errors

Author(s)

Daren Wang

References

Rinaldo, Wang, Wen, Willett and Yu (2020) <arxiv:2010.10410>

Examples

```
d0 = 10
p = 20
n = 100
cpt_true = c(30, 70)
data = simu.change.regression(d0, cpt_true, p, n, sigma = 1, kappa = 9)
gamma_set = c(0.01, 0.1, 1)
lambda_set = c(0.01, 0.1, 1, 3)
temp = CV.search.DP.regression(y = data$y, X = data$X, gamma_set, lambda_set, delta = 2)
temp$test_error # test error result
# find the indices of gamma_set and lambda_set which minimizes the test error
min_idx = as.vector(arrayInd(which.min(temp$test_error), dim(temp$test_error)))
gamma_set[min_idx[1]]
lambda_set[min_idx[2]]
cpt_init = unlist(temp$cpt_hat[min_idx[1], min_idx[2]])
```

CV.search.DP.univar *Grid search for dynamic programming to select the tuning parameter through Cross-Validation.*

Description

Perform grid search for dynamic programming to select the tuning parameter through Cross-Validation.

Usage

```
CV.search.DP.univar(y, gamma_set, delta)
```

Arguments

y	A numeric vector of observations.
gamma_set	A numeric vector of candidate tuning parameter associated with the l_0 penalty.
delta	A positive integer scalar of minimum spacing.

Value

A list with the following structure:

cpt_hat	A list of vector of estimated change points (sorted in strictly increasing order).
K_hat	A list of scalar of number of estimated change points.
test_error	A list of vector of testing errors.
train_error	A list of vector of training errors.

Author(s)

Daren Wang & Haotian Xu

References

Wang, Yu and Rinaldo (2020) <doi:10.1214/20-EJS1710>

Examples

```
set.seed(0)
cpt_true = c(20, 50, 170)
y = rnorm(300) + c(rep(0,20),rep(2,30),rep(0,120),rep(2,130))
gamma_set = 1:5
DP_result = CV.search.DP.univar(y, gamma_set, delta = 5)
min_idx = which.min(DP_result$test_error)
cpt_hat = unlist(DP_result$cpt_hat[min_idx])
Hausdorff.dist(cpt_hat, cpt_true)
```

CV.search.DP.VAR1 *Grid search based on cross-validation of dynamic programming for VAR change points detection via l_0 penalty.*

Description

Perform grid search based on cross-validation of dynamic programming for VAR change points detection.

Usage

```
CV.search.DP.VAR1(DATA, gamma_set, lambda_set, delta, eps = 0.001)
```

Arguments

DATA	A numeric matrix of observations with horizontal axis being time, and vertical axis being dimensions.
gamma_set	A numeric vector of candidate tuning parameters associated with the l_0 penalty.
lambda_set	A numeric vector of candidate tuning parameters for lasso penalty.
delta	A strictly integer scalar of minimum spacing.
eps	A numeric scalar of precision level for convergence of lasso.

Value

A list with the following structure:

cpt_hat	A list of vector of estimated change points
K_hat	A list of scalar of number of estimated change points
test_error	A list of vector of testing errors (each row corresponding to each gamma, and each column corresponding to each lambda)
train_error	A list of vector of training errors

Author(s)

Daren Wang & Haotian Xu

References

Wang, Yu, Rinaldo and Willett (2019) <arxiv:1909.06359>

Examples

```

set.seed(123)
p = 10
sigma = 1
n = 20
v1 = 2*(seq(1,p,1)%%2) - 1
v2 = -v1
AA = matrix(0, nrow = p, ncol = p-2)
A1 = cbind(v1,v2,AA)*0.1
A2 = cbind(v2,v1,AA)*0.1
A3 = A1
cpt_true = c(40, 80)
data = simu.VAR1(sigma, p, 2*n+1, A1)
data = cbind(data, simu.VAR1(sigma, p, 2*n, A2, vzero=c(data[,ncol(data)])))
data = cbind(data, simu.VAR1(sigma, p, 2*n, A3, vzero=c(data[,ncol(data)])))
gamma_set = c(0.1, 0.5, 1)
lambda_set = c(0.1, 1, 3.2)
temp = CV.search.DP.VAR1(data, gamma_set, lambda_set, delta = 5)
temp$test_error # test error result
# find the indices of gamma.set and lambda.set which minimizes the test error
min_idx = as.vector(arrayInd(which.min(temp$test_error), dim(temp$test_error)))
cpt_init = unlist(temp$cpt_hat[min_idx[1], min_idx[2]])
Hausdorff.dist(cpt_init, cpt_true)

```

CV.search.DPDU.regression

Grid search based on cross-validation of dynamic programming for regression change points localisation with l_0 penalisation.

Description

Perform grid search to select tuning parameters gamma (for l_0 penalty of DP) and lambda (for lasso penalty) based on cross-validation.

Usage

```
CV.search.DPDU.regression(y, X, lambda_set, zeta_set, eps = 0.001)
```

Arguments

y	A numeric vector of response variable.
X	A numeric matrix of covariates with vertical axis being time.
lambda_set	A numeric vector of candidate tuning parameters for lasso penalty.
zeta_set	An integer vector of tuning parameter associated with l_0 penalty (minimum interval size).
eps	A numeric scalar of precision level for convergence of lasso.

Value

A list with the following structure:

cpt_hat	A list of vectors of estimated change points
K_hat	A list of scalars of number of estimated change points
test_error	A matrix of testing errors (each row corresponding to each gamma, and each column corresponding to each lambda)
train_error	A matrix of training errors
beta_hat	A list of matrices of estimated regression coefficients

Author(s)

Haotian Xu

References

Xu, Wang, Zhao and Yu (2022) <arXiv:2207.12453>.

Examples

```
d0 = 5
p = 30
n = 200
cpt_true = 100
data = simu.change.regression(d0, cpt_true, p, n, sigma = 1, kappa = 9)
lambda_set = c(0.3, 0.5, 1, 2)
zeta_set = c(10, 15, 20)
temp = CV.search.DPDU.regression(y = data$y, X = data$X, lambda_set, zeta_set)
temp$test_error # test error result
# find the indices of lambda_set and zeta_set which minimizes the test error
min_idx = as.vector(arrayInd(which.min(temp$test_error), dim(temp$test_error)))
lambda_set[min_idx[2]]
zeta_set[min_idx[1]]
cpt_init = unlist(temp$cpt_hat[min_idx[1], min_idx[2]])
beta_hat = matrix(unlist(temp$beta_hat[min_idx[1], min_idx[2]]), ncol = length(cpt_init)+1)
```

DP.poly

Dynamic programming algorithm for univariate polynomials change points detection.

Description

Perform dynamic programming algorithm for univariate polynomials change points detection.

Usage

```
DP.poly(y, r, gamma, delta)
```

Arguments

y	A numeric vector of observations.
r	An integer scalar order of polynomials.
gamma	A numeric scalar of the tuning parameter associated with the l_0 penalty.
delta	A strictly integer scalar of minimum spacing.

Value

A list with the following structure:

An object of `class` "DP", which is a list with the following structure:

partition	A vector of the best partition.
yhat	A vector of mean estimation for corresponding to the best partition.
cpt	A vector of change points estimation.

Author(s)

Haotian Xu

References

Yu and Chatterjee (2020) <arXiv:2007.09910>

Examples

```
set.seed(0)
cpt_true = c(20, 50, 170)
y = rnorm(300) + c(rep(0,20),rep(2,30),rep(0,120),rep(2,130))
plot.ts(y)
temp = DP.poly(y, r = 2, gamma = 15, delta = 5)
temp$cpt
```

DP.regression	<i>Dynamic programming algorithm for regression change points localisation with l_0 penalisation.</i>
---------------	--

Description

Perform dynamic programming algorithm for regression change points localisation.

Usage

```
DP.regression(y, X, gamma, lambda, delta, eps = 0.001)
```

Arguments

y	A numeric vector of response variable.
X	A numeric matrix of covariates with vertical axis being time.
gamma	A positive numeric scalar stands for tuning parameter associated with l_0 penalty.
lambda	A positive numeric scalar stands for tuning parameter associated with the lasso penalty.
delta	A positive integer scalar stands for minimum spacing.
eps	A numeric scalar of precision level for convergence of lasso.

Value

An object of class "DP", which is a list with the following structure:

partition	A vector of the best partition.
cpt	A vector of change points estimation.

Author(s)

Daren Wang & Haotian Xu

References

Rinaldo, Wang, Wen, Willett and Yu (2020) <arxiv:2010.10410>

Examples

```
d0 = 10
p = 20
n = 100
cpt_true = c(30, 70)
data = simu.change.regression(d0, cpt_true, p, n, sigma = 1, kappa = 9)
temp = DP.regression(y = data$y, X = data$X, gamma = 2, lambda = 1, delta = 5)
cpt_hat = temp$cpt
```

DP.SEPP

Dynamic programming for SEPP change points detection through l_0 penalty.

Description

Perform dynamic programming for SEPP change points detection.

Usage

```
DP.SEPP(DATA, gamma, lambda, delta, delta2, intercept, threshold)
```

Arguments

DATA	A numeric matrix of observations with horizontal axis being time.
gamma	A numeric scalar of tuning parameter associated with the l_0 penalty.
lambda	A numeric scalar of tuning parameter for lasso penalty.
delta	An integer scalar of minimum spacing for dynamic programming.
delta2	An integer scalar representing the maximal of the change point spacing (for reducing computation cost).
intercept	A numeric scalar representing the intercept of the model, which is assumed to be known.
threshold	A numeric scalar representing the upper bound for each coordinate of X_t (for stability).

Value

An object of `class` "DP", which is a list with the following structure:

partition	A vector of the best partition.
cpt	A vector of change points estimation.

Author(s)

Daren Wang & Haotian Xu

References

Wang, D., Yu, Y., & Willett, R. (2020). Detecting Abrupt Changes in High-Dimensional Self-Exciting Poisson Processes. arXiv preprint arXiv:2006.03572.

Examples

```
p = 8 # dimension
n = 15
s = 3 # s is sparsity
factor = 0.2 # large factor gives exact recovery
threshold = 4 # thresholding makes the process stable
intercept = 1/2 # intercept of the model. Assume to be known as in the existing literature
A1 = A2 = A3 = matrix(0, p, p)
diag(A1[, -1]) = 1
diag(A1) = 1
diag(A1[-1,]) = -1
A1 = A1*factor
A1[(s+1):p, (s+1):p] = 0
diag(A2[, -1]) = 1
diag(A2) = -1
diag(A2[-1,]) = 1
A2 = A2*factor
A2[(s+1):p, (s+1):p] = 0
data1 = simu.SEPP(intercept, n, A1, threshold, vzero = NULL)
data2 = simu.SEPP(intercept, n, A2, threshold, vzero = data1[,n])
```

```

data = cbind(data1, data2)
gamma = 0.1
delta = 0.5*n
delta2 = 1.5*n
intercept = 1/2
threshold = 6
DP_result = DP.SEPP(data, gamma = gamma, lambda = 0.03, delta, delta2, intercept, threshold)
cpt_hat = DP_result$cpt

```

DP.univar	<i>Dynamic programming for univariate mean change points detection through l_0 penalty.</i>
-----------	--

Description

Perform dynamic programming for univariate mean change points detection.

Usage

```
DP.univar(y, gamma, delta)
```

Arguments

y	A numeric vector of observations.
gamma	A numeric scalar of the tuning parameter associated with l_0 penalty.
delta	A positive integer scalar of minimum spacing.

Value

An object of `class` "DP", which is a list with the following structure:

partition	A vector of the best partition.
yhat	A vector of mean estimation for corresponding to the best partition.
cpt	A vector of change points estimation.

Author(s)

Haotian Xu

References

Wang, Yu and Rinaldo (2020) <doi:10.1214/20-EJS1710>

Examples

```

set.seed(123)
cpt_true = c(20, 50, 170)
y = rnorm(300) + c(rep(0,20),rep(1,30),rep(0,120),rep(1,130))
DP_result = DP.univar(y, gamma = 5, delta = 5)
cpt_hat = DP_result$cpt
Hausdorff.dist(cpt_hat, cpt_true)

```

DP.VAR1	<i>Dynamic programming for VAR1 change points detection through l_0 penalty.</i>
---------	---

Description

Perform dynamic programming for VAR1 change points detection through l_0 penalty.

Usage

```
DP.VAR1(X_futu, X_curr, gamma, lambda, delta, eps = 0.001)
```

Arguments

X_futu	A numeric matrix of time series at one step ahead, with horizontal axis being time.
X_curr	A numeric matrix of time series at current step, with horizontal axis being time.
gamma	A numeric scalar of the tuning parameter associated with l_0 penalty.
lambda	A numeric scalar of the tuning parameter for lasso penalty.
delta	A strictly integer scalar of minimum spacing.
eps	A numeric scalar of precision level for convergence of lasso.

Value

An object of `class` "DP", which is a list with the following structure:

partition	A vector of the best partition.
cpt	A vector of change points estimation.

Author(s)

Daren Wang & Haotian Xu

References

Wang, Yu, Rinaldo and Willett (2019) <arxiv:1909.06359>

Examples

```

p = 10
sigma = 1
n = 20
v1 = 2*(seq(1,p,1)%%2) - 1
v2 = -v1
AA = matrix(0, nrow = p, ncol = p-2)
A1 = cbind(v1,v2,AA)*0.1
A2 = cbind(v2,v1,AA)*0.1
A3 = A1
data = simu.VAR1(sigma, p, 2*n+1, A1)
data = cbind(data, simu.VAR1(sigma, p, 2*n, A2, vzero=c(data[,ncol(data)])))
data = cbind(data, simu.VAR1(sigma, p, 2*n, A3, vzero=c(data[,ncol(data)])))
N = ncol(data)
X_curr = data[,1:(N-1)]
X_futu = data[,2:N]
DP_result = DP.VAR1(X_futu, X_curr, gamma = 1, lambda = 1, delta = 5)
DP_result$cpt

```

DPDU.regression

Dynamic programming with dynamic update algorithm for regression change points localisation with l_0 penalisation (earlier version).

Description

Perform DPDU algorithm for regression change points localisation.

Usage

```
DPDU.regression(y, X, lambda, zeta, eps = 0.001)
```

Arguments

y	A numeric vector of response variable.
X	A numeric matrix of covariates with vertical axis being time.
lambda	A positive numeric scalar of tuning parameter for lasso penalty.
zeta	A positive integer scalar of tuning parameter associated with l_0 penalty (minimum interval size).
eps	A numeric scalar of precision level for convergence of lasso.

Value

An object of class "DP", which is a list with the following structure:

partition	A vector of the best partition.
cpt	A vector of change points estimation.

Author(s)

Haotian Xu

References

Xu, Wang, Zhao and Yu (2022) <arXiv:2207.12453>.

Examples

```
d0 = 10
p = 20
n = 100
cpt_true = c(30, 70)
data = simu.change.regression(d0, cpt_true, p, n, sigma = 1, kappa = 9)
temp = DPDU.regression(y = data$y, X = data$X, lambda = 1, zeta = 20)
cpt_hat = temp$cpt
```

DPDU2.regression	<i>Dynamic programming with dynamic update algorithm for regression change points localisation with l_0 penalisation (current version).</i>
------------------	--

Description

Perform DPDU algorithm for regression change points localisation.

Usage

```
DPDU2.regression(y, X, lambda, zeta, eps = 0.001)
```

Arguments

y	A numeric vector of response variable.
X	A numeric matrix of covariates with vertical axis being time.
lambda	A positive numeric scalar of tuning parameter for lasso penalty.
zeta	A positive integer scalar of tuning parameter associated with l_0 penalty (minimum interval size).
eps	A numeric scalar of precision level for convergence of lasso.

ValueAn object of `class` "DP", which is a list with the following structure:

partition	A vector of the best partition.
cpt	A vector of change points estimation.

Author(s)

Haotian Xu

References

Xu, Wang, Zhao and Yu (2022) <arXiv:2207.12453>.

Examples

```
d0 = 10
p = 20
n = 100
cpt_true = c(30, 70)
data = simu.change.regression(d0, cpt_true, p, n, sigma = 1, kappa = 9)
temp = DPDU2.regression(y = data$y, X = data$X, lambda = 1, zeta = 20)
cpt_hat = temp$cpt
```

gen.cov.mat

Generate population covariance matrix with dimension p.

Description

Generate population covariance matrix with dimension p.

Usage

```
gen.cov.mat(p, sigma2, type)
```

Arguments

p	A integer scalar of dimensionality.
sigma2	A positive numeric scalar representing the variance of each entry.
type	Specify the type of a covariance matrix: Diagonal structure ("diagonal"); Equal correlation structure ("equal"); Power decay structure ("power").

Value

A numeric p-by-p matrix.

Author(s)

Haotian Xu

Examples

```
gen.cov.mat(p = 5, sigma2 = 1, type = "diagonal")
```

gen.missing	<i>Function to generate a matrix with values 0 or 1, where 0 indicating the entry is missing</i>
-------------	--

Description

Function to generate a matrix with values 0 or 1, where 0 indicating the entry is missing

Usage

```
gen.missing(pi_mat, symm = TRUE)
```

Arguments

pi_mat	A numeric p x p matrix, for each entry, the value representing the probability of missing.
symm	A logic scalar indicating if the output matrix needs to be symmetric.

Value

A numeric p x p matrix.

Author(s)

Haotian Xu

Examples

```
p = 5
pi_mat = matrix(0.9, p, p)
eta_mat = gen.missing(pi_mat, symm = TRUE)
```

gen.piece.poly	<i>Generate univariate data from piecewise polynomials of degree at most r.</i>
----------------	---

Description

Generate univariate data from piecewise polynomials (currently, only the linear, quadratic functions and cubic functions are considered).

Usage

```
gen.piece.poly(init_coef_vec, cpt_vec, kappa_mat, n, sigma)
```

Arguments

<code>init_coef_vec</code>	A (r+1)-dim numeric vector of coefficients for the first segment.
<code>cpt_vec</code>	A K-dim integer vector of change points.
<code>kappa_mat</code>	A (r+1)xK numeric matrix where the i-th column represents the jump sizes for coefficients associated with the i-th change point.
<code>n</code>	An integer scalar of sample size.
<code>sigma</code>	A numeric scalar of standard deviation of error terms.

Value

A vector of data generated from piecewise polynomials.

Author(s)

Haotian Xu

References

Yu and Chatterjee (2020) <arXiv:2007.09910>.

Examples

```
r = 2
init_coef_vec = c(-2, 2, 9)
cpt_true = c(100, 200)
n = 300
sigma = 1
kappa_mat = cbind(c(3, 9, -27), c(-3, 9, -27))
plot.ts(gen.piece.poly(init_coef_vec, cpt_true, kappa_mat, n, sigma), ylab = "y")
```

gen.piece.poly.noiseless

Mean function of piecewise polynomials.

Description

Compute mean function of piecewise polynomials (currently, only the linear, quadratic functions and cubic functions are considered).

Usage

```
gen.piece.poly.noiseless(init_coef_vec, cpt_vec, kappa_mat, n)
```

Arguments

<code>init_coef_vec</code>	A numeric vector of coefficients for the first segment.
<code>cpt_vec</code>	An integer vector of change points.
<code>kappa_mat</code>	A numeric matrix where the <i>i</i> -th column represents the jump sizes for coefficients associated with the <i>i</i> -th change point.
<code>n</code>	An integer scalar of sample size.

Value

A vector of mean function of piecewise polynomials.

Author(s)

Haotian Xu

References

Yu and Chatterjee (2020) <arXiv:2007.09910>

Examples

```
r = 2
init_coef_vec = c(-2, 2, 9)
cpt_true = c(100, 200)
n = 300
kappa_mat = cbind(c(3, 9, -27), c(-3, 9, -27))
plot.ts(gen.piece.poly.noiseless(init_coef_vec, cpt_true, kappa_mat, n),
        ylab = "Values of piecewise polynomials")
```

Hausdorff.dist	<i>Bidirectional Hausdorff distance.</i>
----------------	--

Description

Compute the bidirectional Hausdorff distance between two sets.

Usage

```
Hausdorff.dist(vec1, vec2)
```

Arguments

<code>vec1</code>	A integer vector forms a subset of 1, 2, ..., n.
<code>vec2</code>	A integer vector forms a subset of 1, 2, ..., n.

Value

An integer scalar of bidirectional Hausdorff distance.

Author(s)

Daren Wang

Examples

```
vec1 = sample.int(1000, size = 50)
vec2 = sample.int(2000, size = 100)
Hausdorff.dist(vec1, vec2)
```

huber_mean

Element-wise adaptive Huber mean estimator.

Description

Computes the element-wise adaptive Huber mean estimator.

Usage

```
huber_mean(x, tau)
```

Arguments

`x` A numeric vector of observations.
`tau` A numeric scalar corresponding to the robustification parameter (larger than 0).

Value

A numeric scalar corresponding to the adaptive Huber mean estimator.

Author(s)

Haotian Xu

Examples

```
set.seed(123)
y = rnorm(100)
mean(y)
huber_mean(y, 1.345)
```

kde.biwei.eval *Multivariate kernel density estimation based on Biweight kernel.*

Description

Perform multivariate kernel density estimation and evaluated the estimated densities at specified points.

Usage

```
kde.biwei.eval(x, bw, eval.points)
```

Arguments

x	A numeric matrix of observations with horizontal axis being dimension, and vertical axis being time.
eval.points	A numeric matrix of evaluated data points with horizontal axis being dimension, and vertical axis being time..
h	A numeric scalar of bandwidth parameter.

Value

A numeric vector of evaluated densities.

Author(s)

Haotian Xu

Examples

```
n = 100
p = 10
x = matrix(rnorm(n*p), nrow = n)
h = 2*(1/n)^(1/(4+p)) # bandwidth
kde.biwei.eval(x, h, x)
```

kde.epan.eval *Multivariate kernel density estimation based on Epanechnikov kernel.*

Description

Perform multivariate kernel density estimation and evaluated the estimated densities at specified points.

Usage

```
kde.epan.eval(x, bw, eval.points)
```

Arguments

`x` A numeric matrix of observations with horizontal axis being dimension, and vertical axis being time.

`eval.points` A numeric matrix of evaluated data points with horizontal axis being dimension, and vertical axis being time..

`h` A numeric scalar of bandwidth parameter.

Value

A numeric vector of evaluated densities.

Author(s)

Haotian Xu

Examples

```
n = 100
p = 10
x = matrix(rnorm(n*p), nrow = n)
h = 2*(1/n)^{1/(4+p)} # bandwidth
kde.epan.eval(x, h, x)
```

kde.eval

Multivariate kernel density estimation based on Gaussian kernel.

Description

Perform multivariate kernel density estimation and evaluated the estimated densities at specified points.

Usage

```
kde.eval(x, H, eval.points)
```

Arguments

`x` A numeric matrix of observations with horizontal axis being dimension, and vertical axis being time.

`H` A numeric (symmetric and positive definite) matrix of bandwidth parameters.

`eval.points` A numeric matrix of evaluated data points with horizontal axis being dimension, and vertical axis being time..

Value

A numeric vector of evaluated densities.

Author(s)

Haotian Xu

Examples

```
n = 100
p = 10
x = matrix(rnorm(n*p), nrow = n)
h = 2*(1/n)^(1/(4+p)) # bandwidth
kde.eval(x, h*diag(p), x)
```

kde.triwei.eval *Multivariate kernel density estimation based on Triweight kernel.*

Description

Perform multivariate kernel density estimation and evaluated the estimated densities at specified points.

Usage

```
kde.triwei.eval(x, bw, eval.points)
```

Arguments

x	A numeric matrix of observations with horizontal axis being dimension, and vertical axis being time.
eval.points	A numeric matrix of evaluated data points with horizontal axis being dimension, and vertical axis being time..
h	A numeric scalar of bandwidth parameter.

Value

A numeric vector of evaluated densities.

Author(s)

Haotian Xu

Examples

```
n = 100
p = 10
x = matrix(rnorm(n*p), nrow = n)
h = 2*(1/n)^(1/(4+p)) # bandwidth
kde.triwei.eval(x, h, x)
```

lambda.network.missing

Function to compute the default thresholding parameter for leading singular value in the soft-impute algorithm.

Description

Function to compute the default thresholding parameter for leading singular value in the soft-impute algorithm.

Usage

```
lambda.network.missing(s, e, t, alpha, rho, pi_ub, p, C_lambda)
```

Arguments

s	An integer scalar of the starting index.
e	An integer scalar of the ending index.
t	An integer scalar of the splitting index.
alpha	A numeric scalar in (0,1) representing the desired false alarm rate.
rho	A numeric scalar of the sparsity parameter.
pi_ub	A numeric scalar of the upper bound of the missing probability.
p	An integer scalar of the dimensionality of the graphon matrix.
C_lambda	A numeric scalar of an absolute constant, which is set to be 2/3 by default.

Details

The default thresholding parameter is given in Theorem 2 of the reference.

Value

The default thresholding parameter for leading singular value in the soft-impute algorithm

References

Dubey, Xu and Yu (2021) <arxiv:2110.06450>

local.refine.CV.VAR1 *Local refinement for VAR1 change points detection.*

Description

Perform local refinement for VAR1 change points detection.

Usage

```
local.refine.CV.VAR1(cpt_init, DATA, zeta_set, delta_local)
```

Arguments

cpt_init	A integer vector of initial change points estimation (sorted in strictly increasing order).
DATA	A numeric matrix of observations with with horizontal axis being time, and vertical axis being dimensions.
zeta_set	A numeric vector of candidate tuning parameters for group lasso penalty.
delta_local	A strictly integer scalar of minimum spacing for group lasso.

Value

A list with the following structure:

cpt_hat	A vector of estimated change point locations (sorted in strictly increasing order)
zeta	A scalar of selected zeta by cross-validation

Author(s)

Daren Wang & Haotian Xu

References

Wang, Yu, Rinaldo and Willett (2019) <arxiv:1909.06359>

local.refine.DPDU.regression

Local refinement for DPDU regression change points localisation.

Description

Perform local refinement for regression change points localisation.

Usage

```
local.refine.DPDU.regression(cpt_init, beta_hat, y, X, w = 0.9)
```

Arguments

cpt_init	An integer vector of initial changepoints estimation (sorted in strictly increasing order).
beta_hat	A numeric (px(K_hat+1))matrix of estimated regression coefficients.
y	A numeric vector of response variable.
X	A numeric matrix of covariates with vertical axis being time..
w	A numeric scalar in (0,1) representing the weight for interval truncation.

Value

A vector of locally refined change points estimation.

Author(s)

Haotian Xu

References

Xu, Wang, Zhao and Yu (2022) <arXiv:2207.12453>.

Xu, Wang, Zhao and Yu (2022) <arXiv:2207.12453>.

Examples

```
d0 = 5
p = 30
n = 200
cpt_true = 100
data = simu.change.regression(d0, cpt_true, p, n, sigma = 1, kappa = 9)
lambda_set = c(0.3, 0.5, 1, 2)
zeta_set = c(10, 15, 20)
temp = CV.search.DPDU.regression(y = data$y, X = data$X, lambda_set, zeta_set)
temp$test_error # test error result
# find the indices of lambda_set and zeta_set which minimizes the test error
min_idx = as.vector(arrayInd(which.min(temp$test_error), dim(temp$test_error)))
```

```

lambda_set[min_idx[2]]
zeta_set[min_idx[1]]
cpt_init = unlist(temp$cpt_hat[min_idx[1], min_idx[2]])
beta_hat = matrix(unlist(temp$beta_hat[min_idx[1], min_idx[2]]), ncol = length(cpt_init)+1)
cpt_refined = local.refine.DPDU.regression(cpt_init, beta_hat, data$y, data$X, w = 0.9)

```

```
local.refine.multi.nonpar.L2
```

Local refinement for multivariate nonparametric change points localisation based on L2 distance.

Description

Perform local refinement for multivariate nonparametric change points localisation based on L2 distance.

Usage

```

local.refine.multi.nonpar.L2(
  cpt_init,
  Y,
  kappa_hat,
  r = 2,
  w = 0.9,
  c_kappa = 10
)

```

Arguments

<code>cpt_init</code>	An integer vector of initial change points estimation (sorted in strictly increasing order).
<code>Y</code>	A numeric matrix of observations with with horizontal axis being time, and vertical axis being dimension.
<code>kappa_hat</code>	A numeric vector of jump sizes estimator.
<code>r</code>	An integer scalar of smoothness parameter of the underlying Holder space.
<code>w</code>	A numeric scalar in (0,1) representing the weight for interval truncation.
<code>c_kappa</code>	A numeric scalar to be multiplied by kappa estimator as the bandwidth in the local refinement.

Value

A vector of locally refined change points estimation.

Author(s)

Haotian Xu

Examples

```

n = 150
v = c(floor(n/3), 2*floor(n/3)) # location of change points
r = 2
p = 6
Y = matrix(0, p, n) # matrix for data
mu0 = rep(0, p) # mean of the data
mu1 = rep(0, p)
mu1[1:floor(p/2)] = 2
Sigma0 = diag(p) #Covariance matrices of the data
Sigma1 = diag(p)
# Generate data
for(t in 1:n){
  if(t <= v[1] || t > v[2]){
    Y[,t] = MASS::mvrnorm(n = 1, mu0, Sigma0)
  }
  if(t > v[1] && t <= v[2]){
    Y[,t] = MASS::mvrnorm(n = 1, mu1, Sigma1)
  }
}## close for generate data
M = 100
intervals = WBS.intervals(M = M, lower = 1, upper = ncol(Y)) #Random intervals
h = 2*(1/n)^(1/(2*r+p)) # bandwidth
temp = WBS.multi.nonpar.L2(Y, 1, ncol(Y), intervals$Alpha, intervals$Beta, h, delta = 15)
cpt_init = tuneBSmultinonpar(temp, Y)
kappa_hat = kappa.multi.nonpar.L2(cpt_init, Y, h_kappa = 0.01)
local.refine.multi.nonpar.L2(cpt_init, Y, kappa_hat, r = 2, w = 0.9, c_kappa = 2)

```

```
local.refine.multi.nonpar.L2.biwei
```

Local refinement for multivariate nonparametric change points localisation based on L2 distance.

Description

Perform local refinement for multivariate nonparametric change points localisation based on L2 distance.

Usage

```

local.refine.multi.nonpar.L2.biwei(
  cpt_init,
  Y,
  kappa_hat,
  r = 2,
  w = 0.9,
  c_kappa = 10
)

```

Arguments

cpt_init	An integer vector of initial change points estimation (sorted in strictly increasing order).
Y	A numeric matrix of observations with with horizontal axis being time, and vertical axis being dimension.
kappa_hat	A numeric vector of jump sizes estimator.
r	An integer scalar of smoothness parameter of the underlying Holder space.
w	A numeric scalar in (0,1) representing the weight for interval truncation.
c_kappa	A numeric scalar to be multiplied by kappa estimator as the bandwidth in the local refinement.

Value

A vector of locally refined change points estimation.

Author(s)

Haotian Xu

Examples

```

n = 150
v = c(floor(n/3), 2*floor(n/3)) # location of change points
r = 2
p = 6
Y = matrix(0, p, n) # matrix for data
mu0 = rep(0, p) # mean of the data
mu1 = rep(0, p)
mu1[1:floor(p/2)] = 2
Sigma0 = diag(p) #Covariance matrices of the data
Sigma1 = diag(p)
# Generate data
for(t in 1:n){
  if(t <= v[1] || t > v[2]){
    Y[,t] = MASS::mvrnorm(n = 1, mu0, Sigma0)
  }
  if(t > v[1] && t <= v[2]){
    Y[,t] = MASS::mvrnorm(n = 1, mu1, Sigma1)
  }
}## close for generate data
M = 100
intervals = WBS.intervals(M = M, lower = 1, upper = ncol(Y)) #Random intervals
h = 2*(1/n)^(1/(2*r+p)) # bandwidth
temp = WBS.multi.nonpar.L2(Y, 1, ncol(Y), intervals$Alpha, intervals$Beta, h, delta = 15)
cpt_init = tuneBSmultinonpar(temp, Y)
kappa_hat = kappa.multi.nonpar.L2(cpt_init, Y, h_kappa = 0.01)
local.refine.multi.nonpar.L2(cpt_init, Y, kappa_hat, r = 2, w = 0.9, c_kappa = 2)

```

local.refine.multi.nonpar.L2.epan

Local refinement for multivariate nonparametric change points localisation based on L2 distance.

Description

Perform local refinement for multivariate nonparametric change points localisation based on L2 distance.

Usage

```
local.refine.multi.nonpar.L2.epan(  
  cpt_init,  
  Y,  
  kappa_hat,  
  r = 2,  
  w = 0.9,  
  c_kappa = 10  
)
```

Arguments

<code>cpt_init</code>	An integer vector of initial change points estimation (sorted in strictly increasing order).
<code>Y</code>	A numeric matrix of observations with with horizontal axis being time, and vertical axis being dimension.
<code>kappa_hat</code>	A numeric vector of jump sizes estimator.
<code>r</code>	An integer scalar of smoothness parameter of the underlying Holder space.
<code>w</code>	A numeric scalar in (0,1) representing the weight for interval truncation.
<code>c_kappa</code>	A numeric scalar to be multiplied by kappa estimator as the bandwidth in the local refinement.

Value

A vector of locally refined change points estimation.

Author(s)

Haotian Xu

Examples

```

n = 150
v = c(floor(n/3), 2*floor(n/3)) # location of change points
r = 2
p = 6
Y = matrix(0, p, n) # matrix for data
mu0 = rep(0, p) # mean of the data
mu1 = rep(0, p)
mu1[1:floor(p/2)] = 2
Sigma0 = diag(p) #Covariance matrices of the data
Sigma1 = diag(p)
# Generate data
for(t in 1:n){
  if(t <= v[1] || t > v[2]){
    Y[,t] = MASS::mvrnorm(n = 1, mu0, Sigma0)
  }
  if(t > v[1] && t <= v[2]){
    Y[,t] = MASS::mvrnorm(n = 1, mu1, Sigma1)
  }
}## close for generate data
M = 100
intervals = WBS.intervals(M = M, lower = 1, upper = ncol(Y)) #Random intervals
h = 2*(1/n)^(1/(2*r+p)) # bandwidth
temp = WBS.multi.nonpar.L2(Y, 1, ncol(Y), intervals$Alpha, intervals$Beta, h, delta = 15)
cpt_init = tuneBSmultinonpar(temp, Y)
kappa_hat = kappa.multi.nonpar.L2(cpt_init, Y, h_kappa = 0.01)
local.refine.multi.nonpar.L2(cpt_init, Y, kappa_hat, r = 2, w = 0.9, c_kappa = 2)

```

```
local.refine.multi.nonpar.L2.triwei
```

Local refinement for multivariate nonparametric change points localisation based on L2 distance.

Description

Perform local refinement for multivariate nonparametric change points localisation based on L2 distance.

Usage

```

local.refine.multi.nonpar.L2.triwei(
  cpt_init,
  Y,
  kappa_hat,
  r = 2,
  w = 0.9,
  c_kappa = 10
)

```

Arguments

cpt_init	An integer vector of initial change points estimation (sorted in strictly increasing order).
Y	A numeric matrix of observations with with horizontal axis being time, and vertical axis being dimension.
kappa_hat	A numeric vector of jump sizes estimator.
r	An integer scalar of smoothness parameter of the underlying Holder space.
w	A numeric scalar in (0,1) representing the weight for interval truncation.
c_kappa	A numeric scalar to be multiplied by kappa estimator as the bandwidth in the local refinement.

Value

A vector of locally refined change points estimation.

Author(s)

Haotian Xu

Examples

```

n = 150
v = c(floor(n/3), 2*floor(n/3)) # location of change points
r = 2
p = 6
Y = matrix(0, p, n) # matrix for data
mu0 = rep(0, p) # mean of the data
mu1 = rep(0, p)
mu1[1:floor(p/2)] = 2
Sigma0 = diag(p) #Covariance matrices of the data
Sigma1 = diag(p)
# Generate data
for(t in 1:n){
  if(t <= v[1] || t > v[2]){
    Y[,t] = MASS::mvrnorm(n = 1, mu0, Sigma0)
  }
  if(t > v[1] && t <= v[2]){
    Y[,t] = MASS::mvrnorm(n = 1, mu1, Sigma1)
  }
}## close for generate data
M = 100
intervals = WBS.intervals(M = M, lower = 1, upper = ncol(Y)) #Random intervals
h = 2*(1/n)^(1/(2*r+p)) # bandwidth
temp = WBS.multi.nonpar.L2(Y, 1, ncol(Y), intervals$Alpha, intervals$Beta, h, delta = 15)
cpt_init = tuneBSmultinonpar(temp, Y)
kappa_hat = kappa.multi.nonpar.L2(cpt_init, Y, h_kappa = 0.01)
local.refine.multi.nonpar.L2(cpt_init, Y, kappa_hat, r = 2, w = 0.9, c_kappa = 2)

```

local.refine.network *Local refinement for network change points detection.*

Description

Perform local refinement for network change points detection.

Usage

```
local.refine.network(  
  cpt_init,  
  data_mat1,  
  data_mat2,  
  self = FALSE,  
  w = 0.5,  
  tau2,  
  tau3 = Inf  
)
```

Arguments

cpt_init	A integer vector of initial change points estimation (sorted in strictly increasing order).
data_mat1	A numeric matrix of observations with with horizontal axis being time, and with each column be the vectorized adjacency matrix.
data_mat2	A independent copy of data_mat1.
self	A logic scalar indicating if adjacency matrices are required to have self-loop.
w	A numeric scalar in (0,1) indicating the level of shrinkage (large shrinkage if w is small) on the interval between the (k-1)th and (k+1)th preliminary change-point estimator.
tau2	A positive numeric scalar for USVT corresponding to the threshold for singular values of input matrix.
tau3	A positive numeric scalar for USVT corresponding to the threshold for entries of output matrix.

Value

A numeric vector of locally refined change point locations.

Author(s)

Daren Wang & Haotian Xu

References

Wang, Yu and Rinaldo (2018) <arxiv:1809.09602>.

Examples

```

p = 15 # number of nodes
rho = 0.5 # sparsity parameter
block_num = 3 # number of groups for SBM
n = 100 # sample size for each segment
# connectivity matrix for the first and the third segments
conn1_mat = rho * matrix(c(0.6,1,0.6,1,0.6,0.5,0.6,0.5,0.6), nrow = 3)
# connectivity matrix for the second segment
conn2_mat = rho * matrix(c(0.6,0.5,0.6,0.5,0.6,1,0.6,1,0.6), nrow = 3)
set.seed(1)
can_vec = sample(1:p, replace = FALSE) # randomly assign nodes into groups
sbm1 = simu.SBM(conn1_mat, can_vec, n, symm = TRUE, self = TRUE)
sbm2 = simu.SBM(conn2_mat, can_vec, n, symm = TRUE, self = TRUE)
data_mat = cbind(sbm1$obs_mat, sbm2$obs_mat)
data_mat1 = data_mat[,seq(1,ncol(data_mat),2)]
data_mat2 = data_mat[,seq(2,ncol(data_mat),2)]
M = 10
intervals = WBS.intervals(M = M, lower = 1, upper = ncol(data_mat1))
temp = WBS.network(data_mat1, data_mat2, 1, ncol(data_mat1), intervals$Alpha,
                  intervals$Beta, delta = 5)
rho_hat = quantile(rowMeans(data_mat), 0.95)
tau = p*rho_hat*(log(n))^2/20 # default threshold given in the paper
cpt_init = unlist(thresholdBS(temp, tau)$cpt_hat[,1])
cpt_refined = local.refine.network(cpt_init, data_mat1, data_mat2, self = TRUE,
                                  tau2 = p*rho_hat/3, tau3 = Inf)

cpt_WBS = 2*cpt_init
cpt_refined = 2*cpt_refined

```

local.refine.poly *Local refinement for univariate polynomials change point detection.*

Description

Perform local refinement for univariate polynomials change point detection.

Usage

```
local.refine.poly(cpt_init, y, r, delta_lr)
```

Arguments

cpt_init	An integer vector of initial change points estimation (sorted in strictly increasing order).
y	A numeric vector of univariate time series.
r	An integer scalar order of polynomials.
delta_lr	A positive integer scalar of minimum spacing for local refinement.

Value

An integer vector of locally refined change point estimation.

Author(s)

Haotian Xu

References

Yu and Chatterjee (2020) <arXiv:2007.09910>

Examples

```
set.seed(0)
cpt_true = c(20, 50, 170)
y = rnorm(300) + c(rep(0,20),rep(2,30),rep(0,120),rep(2,130))
plot.ts(y)
gamma_set = 3:9
DP_result = CV.search.DP.poly(y, r = 2, gamma_set, delta = 5)
min_idx = which.min(DP_result$test_error)
cpt_init = unlist(DP_result$cpt_hat[min_idx])
local.refine.poly(cpt_init, y, r = 2, delta_lr = 5)
```

local.refine.regression

Local refinement for regression change points localisation.

Description

Perform local refinement for regression change points localisation.

Usage

```
local.refine.regression(cpt_init, y, X, zeta)
```

Arguments

cpt_init	An integer vector of initial change points estimation (sorted in strictly increasing order).
y	A numeric vector of response variable.
X	A numeric matrix of covariates with vertical axis being time..
zeta	A numeric scalar of tuning parameter for the group lasso.

Value

A vector of locally refined change points estimation.

Author(s)

Daren Wang & Haotian Xu

References

Rinaldo, A., Wang, D., Wen, Q., Willett, R., & Yu, Y. (2021, March). Localizing changes in high-dimensional regression models. In International Conference on Artificial Intelligence and Statistics (pp. 2089-2097). PMLR.

Rinaldo, Wang, Wen, Willett and Yu (2020) <arxiv:2010.10410>

Examples

```
d0 = 10
p = 20
n = 100
cpt_true = c(30, 70)
data = simu.change.regression(d0, cpt_true, p, n, sigma = 1, kappa = 9)
gamma_set = c(0.01, 0.1, 1)
lambda_set = c(0.01, 0.1, 1, 3)
temp = CV.search.DP.regression(y = data$y, X = data$X, gamma_set, lambda_set, delta = 2)
temp$test_error # test error result
# find the indices of gamma_set and lambda_set which minimizes the test error
min_idx = as.vector(arrayInd(which.min(temp$test_error), dim(temp$test_error)))
gamma_set[min_idx[1]]
lambda_set[min_idx[2]]
cpt_init = unlist(temp$cpt_hat[min_idx[1], min_idx[2]])
local.refine.regression(cpt_init, data$y, X = data$X, zeta = 0.5)
```

local.refine.univar *Local refinement of an initial estimator for univariate mean change points detection.*

Description

Perform local refinement for univariate mean change points detection.

Usage

```
local.refine.univar(cpt_init, y)
```

Arguments

cpt_init	An integer vector of initial change points estimation (sorted in strictly increasing order).
y	A numeric vector of univariate time series.

Value

An integer vector of locally refined change point estimation.

Author(s)

Haotian Xu

References

Wang, Yu and Rinaldo (2020) <doi:10.1214/20-EJS1710>.

Examples

```
set.seed(0)
cpt_true = c(20, 50, 170)
y = rnorm(300) + c(rep(0,20),rep(2,30),rep(0,120),rep(2,130))
gamma_set = 1:5
DP_result = CV.search.DP.univar(y, gamma_set, delta = 5)
min_idx = which.min(DP_result$test_error)
cpt_hat = unlist(DP_result$cpt_hat[min_idx])
Hausdorff.dist(cpt_hat, cpt_true)
cpt_LR = local.refine.univar(cpt_hat, y)
Hausdorff.dist(cpt_LR, cpt_true)
```

local.refine.VAR1 *Local refinement for VAR1 change points detection.*

Description

Perform local refinement for VAR1 change points detection.

Usage

```
local.refine.VAR1(cpt_init, DATA, zeta)
```

Arguments

cpt_init	A integer vector of initial change points estimation (sorted in strictly increasing order).
DATA	A numeric matrix of observations with with horizontal axis being time, and vertical axis being dimensions.
zeta	A numeric scalar of lasso penalty.

Value

An integer vector of locally refined change points estimation.

Author(s)

Daren Wang & Haotian Xu

References

Wang, Yu, Rinaldo and Willett (2019) <arxiv:1909.06359>.

See Also

[local.refine.CV.VAR1](#).

lowertri2mat	<i>Transform a vector containing lower diagonal entries into a symmetric matrix of dimension p.</i>
--------------	---

Description

Transform a vector containing lower diagonal entries into a symmetric matrix of dimension p.

Usage

```
lowertri2mat(lowertri_vec, p, diag = FALSE)
```

Arguments

`lowertri_vec` A numeric vector containing lower diagonal entries.
`p` A integer scalar of dimensionality.
`diag` A logic scalar indicating if the diagonal entries are contained in `lowertri_vec`.

Value

A numeric $p \times p$ symmetric matrix.

Author(s)

Haotian Xu

Examples

```
A = matrix(1:16, 4, 4)
B = lowertri2mat(A[lower.tri(A)], 4, diag = FALSE)
C = lowertri2mat(A[lower.tri(A, diag = TRUE)], 4, diag = TRUE)
```

LRV.regression	<i>Long-run variance estimation for regression settings with change points.</i>
----------------	---

Description

Estimating long-run variance for regression settings with change points.

Usage

```
LRV.regression(cpt_init, beta_hat, y, X, w = 0.9, block_size)
```

Arguments

cpt_init	An integer vector of initial changepoints estimation (sorted in strictly increasing order).
beta_hat	A numeric ($p \times (K_{\text{hat}}+1)$) matrix of estimated regression coefficients.
y	A numeric vector of response variable.
X	A numeric matrix of covariates with vertical axis being time.
w	A numeric scalar in (0,1) representing the weight for interval truncation.
block_size	An integer scalar corresponding to the block size S in the paper.

Value

A vector of long-run variance estimators associated with all local refined intervals.

Author(s)

Haotian Xu

References

Xu, Wang, Zhao and Yu (2022) <arXiv:2207.12453>.
 Xu, Wang, Zhao and Yu (2022) <arXiv:2207.12453>.

Examples

```
d0 = 5
p = 10
n = 200
cpt_true = c(70, 140)
data = simu.change.regression(d0, cpt_true, p, n, sigma = 1, kappa = 9)
lambda_set = c(0.3, 0.5, 1, 2)
zeta_set = c(10, 15, 20)
temp = CV.search.DPDU.regression(y = data$y, X = data$X, lambda_set, zeta_set)
temp$test_error # test error result
# find the indices of lambda_set and zeta_set which minimizes the test error
```

```

min_idx = as.vector(arrayInd(which.min(temp$test_error), dim(temp$test_error)))
lambda_set[min_idx[2]]
zeta_set[min_idx[1]]
cpt_init = unlist(temp$cpt_hat[min_idx[1], min_idx[2]])
beta_hat = matrix(unlist(temp$beta_hat[min_idx[1], min_idx[2]]), ncol = length(cpt_init)+1)
interval_refine = trim_interval(n, cpt_init)
# choose S
block_size = ceiling(sqrt(min(floor(interval_refine[,2]) - ceiling(interval_refine[,1])))/2)
LRV_est = LRV.regression(cpt_init, beta_hat, data$y, data$X, w = 0.9, block_size)

```

online.network

Online change point detection for network data.

Description

Perform online change point detection for network data by controlling the false alarm rate at level α or controlling the average run length γ . The default choice of the tuning parameters τ_1 , τ_2 and τ_3 are used (see Section 4.1 of the reference).

Usage

```

online.network(
  data_mat1,
  data_mat2,
  self = TRUE,
  b_vec = NULL,
  train_mat = NULL,
  alpha = NULL,
  gamma = NULL,
  permu_num = NULL
)

```

Arguments

data_mat1	A numeric matrix of observations with with horizontal axis being time, and with each column be the vectorized adjacency matrix.
data_mat2	A numeric matrix of observations with with horizontal axis being time, and with each column be the vectorized adjacency matrix (data_mat1 and data_mat2 are independent and have the same dimensions).
self	A logic scalar indicating if adjacency matrices are required to have self-loop.
b_vec	A numeric vector of thresholds b_t with $t \geq 2$.
train_mat	A numeric matrix of training data from a pre-change distribution(no change point), which is only needed to when b_vec is NULL in order to calibrate b_t .
alpha	A numeric scalar in (0,1) representing the level.
gamma	An integer scalar of desired average run length.
permu_num	An integer scalar of number of random permutation for calibration.

Value

A list with the following structure:

cpt	Estimated change point
score	A numeric vector of computed cumsum statistics
b_vec	A numeric vector of thresholds b_t with $t \geq 2$

Author(s)

Oscar Hernan Madrid Padilla & Haotian Xu

References

Yu, Padilla, Wang and Rinaldo (2021) <arxiv:2101.05477>

Examples

```
set.seed(123)
p = 15 # number of nodes
rho = 0.5 # sparsity parameter
block_num = 3 # number of groups for SBM
n = 100 # sample size for each segment
# connectivity matrix for the first and the third segments
conn1_mat = rho * matrix(c(0.6,1,0.6,1,0.6,0.5,0.6,0.5,0.6), nrow = 3)
# connectivity matrix for the second segment
conn2_mat = rho * matrix(c(0.6,0.5,0.6,0.5,0.6,1,0.6,1,0.6), nrow = 3)
set.seed(1)
can_vec = sample(1:p, replace = FALSE) # randomly assign nodes into groups
sbm1 = simu.SBM(conn1_mat, can_vec, n, symm = TRUE, self = TRUE)
sbm2 = simu.SBM(conn2_mat, can_vec, n, symm = TRUE, self = TRUE)
data_mat = cbind(sbm1$obs_mat, sbm2$obs_mat)
data_mat1 = data_mat[,seq(1,ncol(data_mat),2)]
data_mat2 = data_mat[,seq(2,ncol(data_mat),2)]
train_mat = simu.SBM(conn1_mat, can_vec, n = 150, symm = TRUE, self = TRUE)$obs_mat
temp = online.network(data_mat1, data_mat2, self = TRUE, b_vec = NULL, train_mat, alpha = 0.05,
                      gamma = NULL, permu_num = 20)
cpt_hat = 2 * temp$cpt
```

online.network.missing

Online change point detection for network data with missing values.

Description

Perform online change point detection for network with missing values by controlling the false alarm rate at level alpha.

Usage

```
online.network.missing(  
  data_incomplete_list,  
  eta_list,  
  alpha_grid,  
  thresholds_array,  
  rho_hat,  
  pi_ub_hat,  
  C_lambda = 2/3,  
  delta = 5  
)
```

Arguments

data_incomplete_list	A list of adjacency matrices (with entries being 0 or 1) with missing values being coercing to 0.
eta_list	A list of matrices associated with data_incomplete_list, each matrix indicates the missing entries in corresponding adjacency matrix.
alpha_grid	A numeric vector in (0,1) representing the desired false alarm rate.
thresholds_array	A numeric array of calibrated thresholds.
rho_hat	A numeric scalar of the sparsity parameter.
pi_ub_hat	A numeric scalar of the upper bound of the missing probability.
C_lambda	A numeric scalar of an absolute constant, which is set to be 2/3 by default.
delta	An integer scalar of minimum spacing.

Value

Online change point estimator.

Author(s)

Haotian Xu

References

Dubey, Xu and Yu (2021) <arxiv:2110.06450>

See Also

[calibrate.online.network.missing](#) for calibrating thresholds.

online.univar	<i>Online change point detection with controlled false alarm rate or average run length.</i>
---------------	--

Description

Perform online change point detection with controlled false alarm rate or average run length.

Usage

```
online.univar(
  y_vec,
  b_vec = NULL,
  train_vec = NULL,
  alpha = NULL,
  gamma = NULL,
  permu_num = NULL
)
```

Arguments

y_vec	A numeric vector of observations.
b_vec	A numeric vector of thresholds b_t with $t \geq 2$.
train_vec	A numeric vector of training data from a pre-change distribution (no change point), which is only needed to when <code>b_vec</code> is <code>NULL</code> in order to calibrate b_t .
alpha	A numeric scalar of desired false alarm rate.
gamma	An integer scalar of desired average run length.
permu_num	An integer scalar of number of random permutation for calibration.

Value

A list with the following structure:

cpt_hat	An integer scalar of estimated change point location
b_vec	A numeric vector of thresholds b_t with $t \geq 2$

Author(s)

Haotian Xu

References

Yu, Padilla, Wang and Rinaldo (2020) <arxiv:2006.03283>

Examples

```

y_vec = rnorm(150) + c(rep(0, 100), rep(1, 50))
train_vec = rnorm(100)
# control the false alarm rate
temp1 = online.univar(y_vec = y_vec, train_vec = train_vec, alpha = 0.05, permu_num = 20)
temp1$cpt_hat
temp1$b_vec # calibrated threshold

```

online.univar.multi *Online change point detection with potentially multiple change points.*

Description

Perform Online change point detection with potentially multiple change points.

Usage

```

online.univar.multi(
  y_vec,
  b_vec = NULL,
  train_vec = NULL,
  alpha = NULL,
  gamma = NULL,
  permu_num = NULL
)

```

Arguments

y_vec	A numeric vector of observations.
b_vec	A numeric vector of thresholds b_t with $t \geq 2$.
train_vec	A numeric vector of training data from a pre-change distribution (no change point), which is only needed to when b_vec is NULL in order to calibrate b_t .
alpha	A numeric scalar of desired false alarm rate.
gamma	An integer scalar of desired average run length.
permu_num	An integer scalar of number of random permutation for calibration.

Details

```

#' @title Online change point detection with controlled average run length. #' @description Per-
form online change point detection via CUSUM (single change point, type 2). #' @param y_vec
A numeric vector of observations. #' @param gamma A integer scalar of interval length ( $\geq$ 
2). #' @param tau_gamma A numeric scalar of threshold. #' @param ... Additional argu-
ments. #' @return An integer scalar of estimated change points location. #' @export #' @author
Haotian Xu #' @examples #' TO DO online.univar.one2 = function(y_vec, gamma, tau_gamma,
...) t = 1 FLAG = 0 while(FLAG == 0 & t <= length(y_vec)) t = t + 1 e = max(t-gamma,

```

```

0) cusum_vec = sapply((e+1):(t-1), function(s) sqrt((t-s)*(s-e)/(t-e)) * abs(mean(y_vec[(e+1):s])
- mean(y_vec[(s+1):t]))) FLAG = 1 - prod(cusum_vec <= tau_gamma)
return(t)

#' @title Online change point detection via CUSUM (single change point, type 3). #' @description
Perform online change point detection via CUSUM (single change point, type 3). #' @param y_vec
A numeric vector of observations. #' @param tau_vec A numeric vector of thresholds at time
t>= 1. #' @param ... Additional arguments. #' @return An integer scalar of estimated change
point location. #' @export #' @author Haotian Xu #' @examples #' TO DO online.univar.one3 =
function(y_vec, tau_vec, ...) if(length(y_vec) != length(tau_vec)) stop("y_vec and tau_vec should
have the same length.")

t = 1 FLAG = 0 while(FLAG == 0 & t <= length(y_vec)) t = t + 1 J = floor(log2(t)) j = 0 while(j
< J & FLAG == 0) j = j + 1 s_j = t - 2^(j-1) cusum = sqrt((t-s_j)*s_j/t) * abs(mean(y_vec[1:s_j]) -
mean(y_vec[(s_j+1):t])) FLAG = (cusum > tau_vec[t])

return(t)

```

Value

An integer vector of estimated change points.

Author(s)

Haotian Xu

References

Yu, Padilla, Wang and Rinaldo (2020) <arxiv:2006.03283>

Examples

```

y_vec = rnorm(200) + c(rep(0, 50), rep(1, 100), rep(0, 50))
train_vec = rnorm(150)
# control the false alarm rate
temp1 = online.univar.multi(y_vec = y_vec, train_vec = train_vec, alpha = 0.05, permu_num = 20)
temp1

```

simu.change.regression

Simulate a sparse regression model with change points in coefficients.

Description

Simulate a sparse regression model with change points in coefficients under temporal dependence.

Usage

```
simu.change.regression(
  d0,
  cpt_true,
  p,
  n,
  sigma,
  kappa,
  cov_type = "I",
  mod_X = "IID",
  mod_e = "IID"
)
```

Arguments

<code>d0</code>	A numeric scalar stands for the number of nonzero coefficients.
<code>cpt_true</code>	An integer vector contains true change points (sorted in strictly increasing order).
<code>p</code>	An integer scalar stands for the dimensionality.
<code>n</code>	An integer scalar stands for the sample size.
<code>sigma</code>	A numeric scalar stands for error standard deviation.
<code>kappa</code>	A numeric scalar stands for the minimum jump size of coefficient vector in l_2 norm.
<code>cov_type</code>	A character string stands for the type of covariance matrix of covariates. 'I': Identity; 'T': Toeplitz; 'E': Equal-correlation.
<code>mod_X</code>	A character string stands for the time series model followed by the covariates. 'IID': IID multivariate Gaussian; 'AR': Multivariate AR1 with $\rho = 0.3$; Multivariate MA1 $\theta = 0.3$.
<code>mod_e</code>	A character string stands for the time series model followed by the errors 'IID': IID univariate Gaussian; 'AR': Univariate AR1 with $\rho = 0.3$; Univariate MA1 $\theta = 0.3$.

Value

A list with the following structure:

<code>cpt_true</code>	A vector of true changepoints (sorted in strictly increasing order).
<code>X</code>	An n-by-p design matrix.
<code>y</code>	An n-dim vector of response variable.
<code>betafullmat</code>	A p-by-n matrix of coefficients.

Author(s)

Daren Wang, Zifeng Zhao & Haotian Xu

References

Rinaldo, Wang, Wen, Willett and Yu (2020) <arxiv:2010.10410>; Xu, Wang, Zhao and Yu (2022) <arXiv:2207.12453>.

Examples

```
d0 = 10
p = 30
n = 100
cpt_true = c(10, 30, 40, 70, 90)
data = simu.change.regression(d0, cpt_true, p, n, sigma = 1, kappa = 9)
```

simu.RDPG

Simulate a dot product graph (without change point).

Description

Simulate a dot product graph (without change point). The generated data is a matrix with each column corresponding to the vectorized adjacency (sub)matrix at a time point. For example, if the network matrix is required to be symmetric and without self-loop, only the strictly lower diagonal entries are considered.

Usage

```
simu.RDPG(x_mat, n, symm = TRUE, self = FALSE)
```

Arguments

<code>x_mat</code>	A numeric matrix representing the latent positions with horizontal axis being latent dimensions and vertical axis being nodes (each entry takes value in $[0, 1]$).
<code>n</code>	A integer scalar representing the number of observations.
<code>symm</code>	A logic scalar indicating if adjacency matrices are required to be symmetric.
<code>self</code>	A logic scalar indicating if adjacency matrices are required to have self-loop.

Value

A list with the following structure:

<code>obs_mat</code>	A matrix, with each column be the vectorized adjacency (sub)matrix. For example, if "symm = TRUE" and "self = FALSE", only the strictly lower triangular matrix is considered.
<code>graphon_mat</code>	Underlying graphon matrix.

Author(s)

Haotian Xu

Examples

```

p = 20 # number of nodes
n = 50 # sample size for each segment
lat_dim_num = 5 # number of latent dimensions
set.seed(1)
x_mat = matrix(runif(p*lat_dim_num), nrow = p, ncol = lat_dim_num)
x_tilde_mat = matrix(runif(p*lat_dim_num), nrow = p, ncol = lat_dim_num)
y_mat = rbind(x_tilde_mat[1:floor(p/4),], x_mat[(floor(p/4)+1):p,])
rdpg1 = simu.RDPG(x_mat, n, symm = TRUE, self = FALSE)
rdpg2 = simu.RDPG(y_mat, n, symm = TRUE, self = FALSE)
data1_mat = rdpg1$obs_mat
data2_mat = rdpg2$obs_mat
data_mat = cbind(data1_mat, data2_mat)

```

simu.SBM

*Simulate a Stochastic Block Model (without change point).***Description**

Simulate a Stochastic Block Model (without change point). The generated data is a matrix with each column corresponding to the vectorized adjacency (sub)matrix at a time point. For example, if the network matrix is required to be symmetric and without self-loop, only the strictly lower diagonal entries are considered.

Usage

```
simu.SBM(convec_mat, can_vec, n, symm = FALSE, self = TRUE)
```

Arguments

convec_mat	A numeric symmetric matrix representing the connectivity matrix (each entry takes value in $[0, 1]$).
can_vec	A integer p-dim vector of node indices. can_vec is then divided into subvectors corresponding to blocks.
n	A integer scalar representing the number of observations.
symm	A logic scalar indicating if adjacency matrices are required to be symmetric.
self	A logic scalar indicating if adjacency matrices are required to have self-loop.

Value

A list with the following structure:

obs_mat	A matrix, with each column be the vectorized adjacency (sub)matrix. For example, if "symm = TRUE" and "self = FALSE", only the strictly lower triangular matrix is considered.
graphon_mat	Underlying graphon matrix.

Author(s)

Daren Wang & Haotian Xu

References

Wang, Yu and Rinaldo (2018) <arxiv:1809.09602>.

Examples

```

p = 15 # number of nodes
rho = 0.5 # sparsity parameter
block_num = 3 # number of groups for SBM
n = 100 # sample size for each segment
# connectivity matrix for the first and the third segments
conn1_mat = rho * matrix(c(0.6,1,0.6,1,0.6,0.5,0.6,0.5,0.6), nrow = 3)
# connectivity matrix for the second segment
conn2_mat = rho * matrix(c(0.6,0.5,0.6,0.5,0.6,1,0.6,1,0.6), nrow = 3)
set.seed(1)
can_vec = sample(1:p, replace = FALSE) # randomly assign nodes into groups
sbm1 = simu.SBM(conn1_mat, can_vec, n, symm = TRUE, self = TRUE)
sbm2 = simu.SBM(conn2_mat, can_vec, n, symm = TRUE, self = TRUE)
data_mat = cbind(sbm1$obs_mat, sbm2$obs_mat)

```

simu.SEPP

*Simulate a (stable) SEPP model (without change point).***Description**

Simulate a (stable) SEPP model (without change point).

Usage

simu.SEPP(intercept, n, A, threshold, vzero = NULL)

Arguments

intercept	A numeric scalar representing the intercept of the model.
n	An integer scalar representing sample size.
A	A numeric $p \times p$ matrix representing the coefficient matrix.
threshold	A numeric scalar representing the upper bound for each coordinate of X_t (for stability).
vzero	A numeric vector representing the observation at time 0. If vzero = NULL, each coordinate is generated from a Poisson distribution with mean lambda.

Value

A p-by-n matrix.

Author(s)

Daren Wang & Haotian Xu

References

Wang, Yu, & Willett (2020). Detecting Abrupt Changes in High-Dimensional Self-Exciting Poisson Processes. <arXiv:2006.03572>.

Examples

```

p = 10 # dimension
n = 50
s = 5 # sparsity
factor = 0.12 # large factor gives exact recovery
threshold = 4 # thresholding makes the process stable
intercept = 1/2 # intercept of the model. Assume to be known as in the existing literature
A1 = A2 = A3 = matrix(0, p, p)
diag(A1[, -1]) = 1
diag(A1) = 1
diag(A1[-1,]) = -1
A1 = A1*factor
A1[(s+1):p, (s+1):p] = 0
diag(A2[, -1]) = 1
diag(A2) = -1
diag(A2[-1,]) = 1
A2 = A2*factor
A2[(s+1):p, (s+1):p] = 0
diag(A3[, -1]) = 1
diag(A3) = 1
diag(A3[-1,]) = -1
A3 = A3*factor
A3[(s+1):p, (s+1):p] = 0
data1 = simu.SEPP(intercept, n, A1, threshold, vzero = NULL)
data2 = simu.SEPP(intercept, n, A2, threshold, vzero = data1[,n])
data3 = simu.SEPP(intercept, n, A3, threshold, vzero = data2[,n])
data = cbind(data1, data2, data3)
dim(data)

```

simu.VAR1

Simulate from a VAR1 model (without change point).

Description

Simulate data of size n and dimension p from a VAR1 model (without change point) with Gaussian i.i.d. error terms.

Usage

```
simu.VAR1(sigma, p, n, A, vzero = NULL)
```

Arguments

sigma	A numeric scalar representing the standard deviation of error terms.
p	An integer scalar representing dimension.
n	An integer scalar representing sample size.
A	A numeric p-by-p matrix representing the transition matrix of the VAR1 model.
vzero	A numeric vector representing the observation at time 0. If vzero = NULL, it is generated following the distribution of the error terms.

Value

A p-by-n matrix.

Author(s)

Daren Wang

References

Wang, Yu, Rinaldo and Willett (2019) <arxiv:1909.06359>

Examples

```
p = 10
sigma = 1
n = 100
A = matrix(rnorm(p*p), nrow = p)*0.1 # transition matrix
simu.VAR1(sigma, p, n, A)
```

softImpute.network.missing

Estimate graphon matrix by soft-impute for independent adjacency matrices with missing values.

Description

Estimate graphon matrix by soft-impute for independent adjacency matrices with missing values.

Usage

```
softImpute.network.missing(
  data_incomplete_list,
  eta_list,
  lambda,
  a,
  it_max = 10000
)
```

Arguments

data_incomplete_list	A list of adjacency matrices (with entries being 0 or 1) with missing values being coercing to 0.
eta_list	A list of matrices associated with data_incomplete_list, each matrix indicates the missing entries in corresponding adjacency matrix.
lambda	A numeric scalar of thresholding parameter for leading singular value in the soft-impute algorithm.
a	A numeric scalar of truncation parameter in the soft-impute algorithm.
it_max	An integer scalar of maximum iteration for the soft-impute algorithm.

Value

Estimated graphon matrix

Author(s)

Haotian Xu

References

Dubey, Xu and Yu (2021) <arxiv:2110.06450>

thresholdBS	<i>Thresholding a BS object with threshold value tau.</i>
-------------	---

Description

Given a BS object, perform thresholding to find the change point locations.

Usage

```
thresholdBS(BS_object, tau)
```

Arguments

BS_object	A BS object.
tau	A positive numeric scalar of thresholding value.

Value

A list with the following structure:

BS_tree_trimmed	BS_tree with change points which do not satisfy the thresholding criteria removed
cpt_hat	A matrix contains change point locations, values of corresponding statistic, and levels at which each change point is detected

Author(s)

Haotian Xu

See Also[BS.univar](#), [BS.uni.nonpar](#), [BS.cov](#), [WBS.univar](#), [WBS.uni.nonpar](#), [WBS.multi.nonpar](#), [WBS.network](#), [WBSIP.cov](#)**Examples**

```
y = c(rnorm(100, 0, 1), rnorm(100, 10, 10), rnorm(100, 40, 10))
temp = BS.univar(y, 1, 300, 5)
plot.ts(y)
points(x = tail(temp$S[order(temp$Dval)],4), y = y[tail(temp$S[order(temp$Dval)],4)], col = "red")
thresholdBS(temp, 20)
```

`trim_interval`*Interval trimming based on initial change point localisation.*

Description

Performing the interval trimming for local refinement.

Usage

```
trim_interval(n, cpt_init, w = 0.9)
```

Arguments

<code>n</code>	An integer scalar corresponding to the sample size.
<code>cpt_init</code>	An integer vector of initial changepoints estimation (sorted in strictly increasing order).
<code>w</code>	A numeric scalar in (0,1) representing the weight for interval truncation.

Value

A matrix with each row be a trimmed interval.

Author(s)

Haotian Xu

References

Xu, Wang, Zhao and Yu (2022) <arXiv:2207.12453>.

tuneBSmultinonpar	<i>A function to compute change points based on the multivariate non-parametric method with tuning parameter selected by FDR control.</i>
-------------------	---

Description

A function to compute change points based on the multivariate nonparametric method with tuning parameter selected by FDR control.

Usage

```
tuneBSmultinonpar(BS_object, Y)
```

Arguments

BS_object	A "BS" object produced by <code>WBS.multi.nonpar</code> .
Y	A numeric matrix of observations with with horizontal axis being time, and vertical axis being dimension.

Value

A vector of estimated change points.

Author(s)

Oscar Hernan Madrid Padilla & Haotian Xu

References

Padilla, Yu, Wang and Rinaldo (2019) <arxiv:1910.13289>.

See Also

[WBS.multi.nonpar](#).

Examples

```
n = 70
v = c(floor(n/3), 2*floor(n/3)) # location of change points
p = 4
Y = matrix(0, p, n) # matrix for data
mu0 = rep(0, p) # mean of the data
mu1 = rep(0, p)
mu1[1:floor(p/2)] = 2
Sigma0 = diag(p) #Covariance matrices of the data
Sigma1 = diag(p)*2
# Generate data
for(t in 1:n){
  if(t < v[1] || t > v[2]){
```

```

      Y[,t] = MASS::mvrnorm(n = 1, mu0, Sigma0)
    }
    if(t >= v[1] && t < v[2]){
      Y[,t] = MASS::mvrnorm(n = 1, mu1, Sigma1)
    }
  }## close for generate data
  M = 8
  intervals = WBS.intervals(M = M, lower = 1, upper = ncol(Y)) #Random intervals
  K_max = 30
  h = 5*(K_max*log(n)/n)^{1/p} # bandwidth
  temp = WBS.multi.nonpar(Y, Y, 1, ncol(Y), intervals$Alpha, intervals$Beta, h, delta = 10)
  S = tuneBSmultinonpar(temp, Y)

```

tuneBSnonparRDPG	<i>Change points detection for dependent dynamic random dot product graph models.</i>
------------------	---

Description

Perform Change points detection for dependent dynamic random dot product graph models. The tuning parameter τ for WBS is automatically selected based on the BIC-type scores defined in Equation (2.4) in Zou et al. (2014).

Usage

```
tuneBSnonparRDPG(BS_object, data_mat, lowerdiag = FALSE, d)
```

Arguments

BS_object	A "BS" object produced by <code>WBS.nonpar.RDPG</code> .
data_mat	A numeric matrix of observations with horizontal axis being time, and vertical axis being vectorized adjacency matrix.
lowerdiag	A logic scalar. TRUE, if each row of data_mat is the vectorization of the strictly lower diagonal elements in an adjacency matrix. FALSE, if each row of data_mat is the vectorization of all elements in an adjacency matrix.
d	A numeric scalar of the number of leading singular values of an adjacency matrix considered in the scaled PCA algorithm.

Value

A numeric vector of estimated change points.

Author(s)

Oscar Hernan Madrid Padilla & Haotian Xu

References

Padilla, Yu and Priebe (2019) <arxiv:1911.07494>.

See Also

[WBS.nonpar.RDPG](#).

Examples

```
### generate data
p = 20 # number of nodes
n = 50 # sample size for each segment
lat_dim_num = 5 # number of latent dimensions
set.seed(1)
x_mat = matrix(runif(p*lat_dim_num), nrow = p, ncol = lat_dim_num)
x_tilde_mat = matrix(runif(p*lat_dim_num), nrow = p, ncol = lat_dim_num)
y_mat = rbind(x_tilde_mat[1:floor(p/4),], x_mat[(floor(p/4)+1):p,])
rdpg1 = simu.RDPG(x_mat, n, symm = TRUE, self = FALSE)
rdpg2 = simu.RDPG(y_mat, n, symm = TRUE, self = FALSE)
data1_mat = rdpg1$obs_mat
data2_mat = rdpg2$obs_mat
data_mat = cbind(data1_mat, data2_mat)
### detect change points
M = 20 # number of random intervals for WBS
d = 10 # parameter for scaled PCA algorithm
delta = 5
intervals = WBS.intervals(M = M, lower = 1, upper = ncol(data_mat))
WBS_result = WBS.nonpar.RDPG(data_mat, lowerdiag = TRUE, d,
                             Alpha = intervals$Alpha, Beta = intervals$Beta, delta)
cpt_hat = tuneBSnonparRDPG(WBS_result, data_mat, lowerdiag = TRUE, d)
```

tuneBSuninonpar

Wild binary segmentation for univariate nonparametric change points detection with tuning parameter selection.

Description

Perform wild binary segmentation with tuning parameter selection based on sample splitting.

Usage

```
tuneBSuninonpar(BS_object, Y, N)
```

Arguments

BS_object	A "BS" object produced by BS.uni.nonpar or WBS.uni.nonpar.
Y	A numeric matrix of observations with horizontal axis being time, and vertical axis being multiple observations on each time point.
N	A integer vector representing number of multiple observations on each time point.

Value

A vector of estimated change points (sorted in strictly increasing order).

Author(s)

Oscar Hernan Madrid Padilla & Haotian Xu

References

Padilla, Yu, Wang and Rinaldo (2021) <doi:10.1214/21-EJS1809>.

See Also

[BS.uni.nonpar](#) and [WBS.uni.nonpar](#).

Examples

```
Y = t(as.matrix(c(rnorm(100, 0, 1), rnorm(100, 0, 10), rnorm(50, 0, 40))))
W = Y # W is a copy of the matrix Y, it can be Y itself.
N = rep(1, 250)
M = 5
set.seed(123)
intervals = WBS.intervals(M = M, lower = 1, upper = ncol(Y))
BS_object = WBS.uni.nonpar(W, 1, ncol(Y), intervals$Alpha, intervals$Beta, N, delta = 5)
cpt_hat = tuneBSuninonpar(BS_object, Y, N)
```

tuneBSunivar

Univariate mean change points detection based on standard or wild binary segmentation with tuning parameter selected by sSIC.

Description

Perform univariate mean change points detection based on standard or wild binary segmentation. The threshold parameter τ for WBS is automatically selected based on the sSIC score defined in Equation (4) in Fryzlewicz (2014).

Usage

```
tuneBSunivar(BS_object, y)
```

Arguments

BS_object A "BS" object produced by `BS.univar` or `WBS.univar`.
y A numeric vector of observations.

Value

A list with the following structure:

cpt A vector of estimated change point locations (sorted in strictly increasing order).
 tau A scalar of selected threshold tau based on sSIC.

Author(s)

Daren Wang & Haotian Xu

References

Wang, Yu and Rinaldo (2020) <doi:10.1214/20-EJS1710>; Fryzlewicz (2014), Wild binary segmentation for multiple change-point detection, <DOI: 10.1214/14-AOS1245>.

See Also

[BS.univar](#) and [WBS.univar](#).

Examples

```
set.seed(0)
cpt_true = c(20, 50, 170)
y = rnorm(300) + c(rep(0,20),rep(2,30),rep(0,120),rep(2,130))
## change points detection by WBS
intervals = WBS.intervals(M = 100, lower = 1, upper = length(y))
temp2 = WBS.univar(y, 1, length(y), intervals$Alpha, intervals$Beta, delta = 5)
WBS_result = tuneBSunivar(temp2, y)
cpt_WBS = WBS_result$cpt
Hausdorff.dist(cpt_WBS, cpt_true)
```

WBS.intervals

Generate random intervals for WBS.

Description

Generate random intervals for WBS.

Usage

```
WBS.intervals(M, lower = 1, upper)
```

Arguments

M A positive integer scalar of number of random intervals.
 lower A positive integer scalar of lower bound of random intervals.
 upper A positive integer scalar of upper bound of random intervals.

Value

A list with the following structure:

Alpha	A M-dim vector representing the starting indices of random intervals
Beta	A M-dim vector representing the ending indices of random intervals

Author(s)

Oscar Hernan Madrid Padilla

Examples

```
WBS.intervals(120, lower = 1, upper = 300)
```

WBS.multi.nonpar	<i>Wild binary segmentation for multivariate nonparametric change points detection.</i>
------------------	---

Description

Perform wild binary segmentation for multivariate nonparametric change points detection.

Usage

```
WBS.multi.nonpar(Y, W, s, e, Alpha, Beta, h, delta, level = 0)
```

Arguments

Y	A numeric matrix of observations with with horizontal axis being time, and vertical axis being dimension.
W	A copy of the matrix Y, it can be Y itself.
s	A integer scalar of starting index.
e	A integer scalar of ending index.
Alpha	A integer vector of starting indices of random intervals.
Beta	A integer vector of ending indices of random intervals.
h	A numeric scalar of bandwidth parameter.
delta	A integer scalar of minimum spacing.
level	Should be fixed as 0.

Value

An object of `class` "BS", which is a list with the following structure:

S	A vector of estimated change points (sorted in strictly increasing order).
Dval	A vector of values of CUSUM statistic based on KS distance.
Level	A vector representing the levels at which each change point is detected.
Parent	A matrix with the starting indices on the first row and the ending indices on the second row.

Author(s)

Oscar Hernan Madrid Padilla & Haotian Xu

References

Padilla, Yu, Wang and Rinaldo (2019) <arxiv:1910.13289>.

See Also

[thresholdBS](#) for obtain change points estimation, [tuneBSmultinonpar](#) for a tuning version.

Examples

```
n = 70
v = c(floor(n/3), 2*floor(n/3)) # location of change points
p = 4
Y = matrix(0, p, n) # matrix for data
mu0 = rep(0, p) # mean of the data
mu1 = rep(0, p)
mu1[1:floor(p/2)] = 2
Sigma0 = diag(p) #Covariance matrices of the data
Sigma1 = diag(p)*2
# Generate data
for(t in 1:n){
  if(t < v[1] || t > v[2]){
    Y[,t] = MASS::mvrnorm(n = 1, mu0, Sigma0)
  }
  if(t >= v[1] && t < v[2]){
    Y[,t] = MASS::mvrnorm(n = 1, mu1, Sigma1)
  }
}## close for generate data
M = 10
intervals = WBS.intervals(M = M, lower = 1, upper = ncol(Y)) #Random intervals
K_max = 30
h = 5*(K_max*log(n)/n)^(1/p) # bandwidth
temp = WBS.multi.nonpar(Y, Y, 1, ncol(Y), intervals$Alpha, intervals$Beta, h, delta = 10)
result = thresholdBS(temp, median(temp$Dval))
```

WBS.multi.nonpar.L2 *Wild binary segmentation for multivariate nonparametric change points detection based on L2 distance.*

Description

Perform wild binary segmentation for multivariate nonparametric change points detection based on L2 distance.

Usage

```
WBS.multi.nonpar.L2(Y, s, e, Alpha, Beta, h, delta, level = 0)
```

Arguments

Y	A numeric matrix of observations with with horizontal axis being time, and vertical axis being dimension.
s	A integer scalar of starting index.
e	A integer scalar of ending index.
Alpha	A integer vector of starting indices of random intervals.
Beta	A integer vector of ending indices of random intervals.
h	A numeric scalar of bandwidth parameter.
delta	A integer scalar of minimum spacing.
level	Should be fixed as 0.

Value

An object of `class` "BS", which is a list with the following structure:

S	A vector of estimated change points (sorted in strictly increasing order).
Dval	A vector of values of CUSUM statistic based on KS distance.
Level	A vector representing the levels at which each change point is detected.
Parent	A matrix with the starting indices on the first row and the ending indices on the second row.

Author(s)

Haotian Xu

See Also

[thresholdBS](#) for obtain change points estimation, [tuneBSmultinonpar](#) for a tuning version.

Examples

```
n = 150
v = c(floor(n/3), 2*floor(n/3)) # location of change points
r = 2
p = 5
Y = matrix(0, p, n) # matrix for data
mu0 = rep(0, p) # mean of the data
mu1 = rep(0, p)
mu1[1:floor(p/2)] = 2
Sigma0 = diag(p) #Covariance matrices of the data
Sigma1 = diag(p)*2
# Generate data
for(t in 1:n){
  if(t < v[1] || t > v[2]){
    Y[,t] = MASS::mvrnorm(n = 1, mu0, Sigma0)
  }
  if(t >= v[1] && t < v[2]){
```

```

      Y[,t] = MASS::mvrnorm(n = 1, mu1, Sigma1)
    }
}## close for generate data
h = 2*(1/n)^[1/(2*r+p)] # bandwidth
M = 50
intervals = WBS.intervals(M = M, lower = 1, upper = ncol(Y)) #Random intervals
temp = WBS.multi.nonpar.L2(Y, 1, ncol(Y), intervals$Alpha, intervals$Beta, h, delta = 15)
cpt_init = tuneBSmultinonpar(temp, Y)

```

WBS.multi.nonpar.L2.biwei

Wild binary segmentation for multivariate nonparametric change points detection based on L2 distance.

Description

Perform wild binary segmentation for multivariate nonparametric change points detection based on L2 distance.

Usage

```
WBS.multi.nonpar.L2.biwei(Y, s, e, Alpha, Beta, h, delta, level = 0)
```

Arguments

Y	A numeric matrix of observations with with horizontal axis being time, and vertical axis being dimension.
s	A integer scalar of starting index.
e	A integer scalar of ending index.
Alpha	A integer vector of starting indices of random intervals.
Beta	A integer vector of ending indices of random intervals.
h	A numeric scalar of bandwidth parameter.
delta	A integer scalar of minimum spacing.
level	Should be fixed as 0.

Value

An object of `class` "BS", which is a list with the following structure:

S	A vector of estimated change points (sorted in strictly increasing order).
Dval	A vector of values of CUSUM statistic based on KS distance.
Level	A vector representing the levels at which each change point is detected.
Parent	A matrix with the starting indices on the first row and the ending indices on the second row.

Author(s)

Haotian Xu

See Also[thresholdBS](#) for obtain change points estimation, [tuneBSmultinonpar](#) for a tuning version.**Examples**

```

n = 150
v = c(floor(n/3), 2*floor(n/3)) # location of change points
r = 2
p = 5
Y = matrix(0, p, n) # matrix for data
mu0 = rep(0, p) # mean of the data
mu1 = rep(0, p)
mu1[1:floor(p/2)] = 2
Sigma0 = diag(p) #Covariance matrices of the data
Sigma1 = diag(p)*2
# Generate data
for(t in 1:n){
  if(t < v[1] || t > v[2]){
    Y[,t] = MASS::mvrnorm(n = 1, mu0, Sigma0)
  }
  if(t >= v[1] && t < v[2]){
    Y[,t] = MASS::mvrnorm(n = 1, mu1, Sigma1)
  }
}## close for generate data
h = 2*(1/n)^(1/(2*r+p)) # bandwidth
M = 50
intervals = WBS.intervals(M = M, lower = 1, upper = ncol(Y)) #Random intervals
temp = WBS.multi.nonpar.L2.biwei(Y, 1, ncol(Y), intervals$Alpha, intervals$Beta, h, delta = 15)
cpt_init = tuneBSmultinonpar(temp, Y)

```

WBS.multi.nonpar.L2.epan

Wild binary segmentation for multivariate nonparametric change points detection based on L2 distance.

Description

Perform wild binary segmentation for multivariate nonparametric change points detection based on L2 distance.

Usage

```
WBS.multi.nonpar.L2.epan(Y, s, e, Alpha, Beta, h, delta, level = 0)
```

Arguments

Y	A numeric matrix of observations with with horizontal axis being time, and vertical axis being dimension.
s	A integer scalar of starting index.
e	A integer scalar of ending index.
Alpha	A integer vector of starting indices of random intervals.
Beta	A integer vector of ending indices of random intervals.
h	A numeric scalar of bandwidth parameter.
delta	A integer scalar of minimum spacing.
level	Should be fixed as 0.

Value

An object of `class` "BS", which is a list with the following structure:

S	A vector of estimated change points (sorted in strictly increasing order).
Dval	A vector of values of CUSUM statistic based on KS distance.
Level	A vector representing the levels at which each change point is detected.
Parent	A matrix with the starting indices on the first row and the ending indices on the second row.

Author(s)

Haotian Xu

See Also

[thresholdBS](#) for obtain change points estimation, [tuneBSmultinonpar](#) for a tuning version.

Examples

```
n = 150
v = c(floor(n/3), 2*floor(n/3)) # location of change points
r = 2
p = 5
Y = matrix(0, p, n) # matrix for data
mu0 = rep(0, p) # mean of the data
mu1 = rep(0, p)
mu1[1:floor(p/2)] = 2
Sigma0 = diag(p) #Covariance matrices of the data
Sigma1 = diag(p)*2
# Generate data
for(t in 1:n){
  if(t < v[1] || t > v[2]){
    Y[,t] = MASS::mvrnorm(n = 1, mu0, Sigma0)
  }
  if(t >= v[1] && t < v[2]){
```

```

      Y[,t] = MASS::mvrnorm(n = 1, mu1, Sigma1)
    }
}## close for generate data
h = 2*(1/n)^[1/(2*r+p)] # bandwidth
M = 50
intervals = WBS.intervals(M = M, lower = 1, upper = ncol(Y)) #Random intervals
temp = WBS.multi.nonpar.L2.epan(Y, 1, ncol(Y), intervals$Alpha, intervals$Beta, h, delta = 15)
cpt_init = tuneBSmultinonpar(temp, Y)

```

WBS.multi.nonpar.L2.triwei

Wild binary segmentation for multivariate nonparametric change points detection based on L2 distance.

Description

Perform wild binary segmentation for multivariate nonparametric change points detection based on L2 distance.

Usage

```
WBS.multi.nonpar.L2.triwei(Y, s, e, Alpha, Beta, h, delta, level = 0)
```

Arguments

Y	A numeric matrix of observations with with horizontal axis being time, and vertical axis being dimension.
s	A integer scalar of starting index.
e	A integer scalar of ending index.
Alpha	A integer vector of starting indices of random intervals.
Beta	A integer vector of ending indices of random intervals.
h	A numeric scalar of bandwidth parameter.
delta	A integer scalar of minimum spacing.
level	Should be fixed as 0.

Value

An object of `class` "BS", which is a list with the following structure:

S	A vector of estimated change points (sorted in strictly increasing order).
Dval	A vector of values of CUSUM statistic based on KS distance.
Level	A vector representing the levels at which each change point is detected.
Parent	A matrix with the starting indices on the first row and the ending indices on the second row.

Author(s)

Haotian Xu

See Also

[thresholdBS](#) for obtain change points estimation, [tuneBSmultinonpar](#) for a tuning version.

Examples

```

n = 150
v = c(floor(n/3), 2*floor(n/3)) # location of change points
r = 2
p = 5
Y = matrix(0, p, n) # matrix for data
mu0 = rep(0, p) # mean of the data
mu1 = rep(0, p)
mu1[1:floor(p/2)] = 2
Sigma0 = diag(p) #Covariance matrices of the data
Sigma1 = diag(p)*2
# Generate data
for(t in 1:n){
  if(t < v[1] || t > v[2]){
    Y[,t] = MASS::mvrnorm(n = 1, mu0, Sigma0)
  }
  if(t >= v[1] && t < v[2]){
    Y[,t] = MASS::mvrnorm(n = 1, mu1, Sigma1)
  }
}## close for generate data
h = 2*(1/n)^(1/(2*r+p)) # bandwidth
M = 50
intervals = WBS.intervals(M = M, lower = 1, upper = ncol(Y)) #Random intervals
temp = WBS.multi.nonpar.L2.triwei(Y, 1, ncol(Y), intervals$Alpha, intervals$Beta, h, delta = 15)
cpt_init = tuneBSmultinonpar(temp, Y)

```

WBS.network

Wild binary segmentation for network change points detection.

Description

Perform wild binary segmentation for network change points detection.

Usage

```
WBS.network(data_mat1, data_mat2, s, e, Alpha, Beta, delta, level = 0)
```

Arguments

data_mat1	A numeric matrix of observations with with horizontal axis being time, and with each column be the vectorized adjacency matrix.
data_mat2	An independent copy of data_mat1.
s	A integer scalar of starting index.
e	A integer scalar of ending index.
Alpha	A integer vector of starting indices of random intervals.
Beta	A integer vector of ending indices of random intervals.
delta	A positive integer scalar of minimum spacing.
level	Should be fixed as 0.

Value

An object of `class` "BS", which is a list with the following structure:

S	A vector of estimated change points (sorted in strictly increasing order).
Dval	A vector of values of CUSUM statistic based on KS distance.
Level	A vector representing the levels at which each change point is detected.
Parent	A matrix with the starting indices on the first row and the ending indices on the second row.

Author(s)

Daren Wang & Haotian Xu

References

Wang, Yu and Rinaldo (2018) <arxiv:1809.09602>.

See Also

[thresholdBS](#) for obtaining change points estimation.

Examples

```
p = 15 # number of nodes
rho = 0.5 # sparsity parameter
block_num = 3 # number of groups for SBM
n = 100 # sample size for each segment
# connectivity matrix for the first and the third segments
conn1_mat = rho * matrix(c(0.6,1,0.6,1,0.6,0.5,0.6,0.5,0.6), nrow = 3)
# connectivity matrix for the second segment
conn2_mat = rho * matrix(c(0.6,0.5,0.6,0.5,0.6,1,0.6,1,0.6), nrow = 3)
set.seed(1)
can_vec = sample(1:p, replace = FALSE) # randomly assign nodes into groups
sbm1 = simu.SBM(conn1_mat, can_vec, n, symm = TRUE, self = TRUE)
sbm2 = simu.SBM(conn2_mat, can_vec, n, symm = TRUE, self = TRUE)
```

```

data_mat = cbind(sbm1$obs_mat, sbm2$obs_mat)
data_mat1 = data_mat[,seq(1,ncol(data_mat),2)]
data_mat2 = data_mat[,seq(2,ncol(data_mat),2)]
M = 10
intervals = WBS.intervals(M = M, lower = 1, upper = ncol(data_mat1))
temp = WBS.network(data_mat1, data_mat2, 1, ncol(data_mat1),
                   intervals$Alpha, intervals$Beta, delta = 5)
rho_hat = quantile(rowMeans(data_mat), 0.95)
tau = p*rho_hat*(log(n))^2/20 # default threshold given in the paper
cpt_init = unlist(thresholdBS(temp, tau)$cpt_hat[,1])
cpt_refined = local.refine.network(cpt_init, data_mat1, data_mat2,
                                  self = TRUE, tau2 = p*rho_hat/3, tau3 = Inf)
cpt_WBS = 2*cpt_init
cpt_refined = 2*cpt_refined

```

WBS.nonpar.RDPG	<i>Wild binary segmentation for dependent dynamic random dot product graph models.</i>
-----------------	--

Description

Perform wild binary segmentation for dependent dynamic random dot product graph models.

Usage

```
WBS.nonpar.RDPG(data_mat, lowerdiag = FALSE, d, Alpha, Beta, delta)
```

Arguments

data_mat	A numeric matrix of observations with horizontal axis being time, and vertical axis being vectorized adjacency matrix.
lowerdiag	A logic scalar. TRUE, if each row of data_mat is the vectorization of the strictly lower diagonal elements in an adjacency matrix. FALSE, if each row of data_mat is the vectorization of all elements in an adjacency matrix.
d	A numeric scalar of the number of leading singular values of an adjacency matrix considered in the scaled PCA algorithm.
Alpha	A integer vector of starting indices of random intervals.
Beta	A integer vector of ending indices of random intervals.
delta	A positive integer scalar of minimum spacing.

Value

An object of class "BS", which is a list with the following structure:

S	A vector of estimated change point locations (sorted in strictly increasing order).
Dval	A vector of values of CUSUM statistic.
Level	A vector representing the levels at which each change point is detected.
Parent	A matrix with the starting indices on the first row and the ending indices on the second row.

Author(s)

Oscar Hernan Madrid Padilla, Haotian Xu

References

Padilla, Yu and Priebe (2019) <arxiv:1911.07494>.

See Also

[thresholdBS](#) for obtaining change points estimation, [tuneBSnonparRDPG](#) for a tuning version.

Examples

```
### generate data
p = 20 # number of nodes
n = 50 # sample size for each segment
lat_dim_num = 5 # number of latent dimensions
set.seed(1)
x_mat = matrix(runif(p*lat_dim_num), nrow = p, ncol = lat_dim_num)
x_tilde_mat = matrix(runif(p*lat_dim_num), nrow = p, ncol = lat_dim_num)
y_mat = rbind(x_tilde_mat[1:floor(p/4),], x_mat[(floor(p/4)+1):p,])
rdpg1 = simu.RDPG(x_mat, n, symm = TRUE, self = FALSE)
rdpg2 = simu.RDPG(y_mat, n, symm = TRUE, self = FALSE)
data1_mat = rdpg1$obs_mat
data2_mat = rdpg2$obs_mat
data_mat = cbind(data1_mat, data2_mat)
### detect change points
M = 30 # number of random intervals for WBS
d = 10 # parameter for scaled PCA algorithm
delta = 5
intervals = WBS.intervals(M = M, lower = 1, upper = ncol(data_mat))
WBS_result = WBS.nonpar.RDPG(data_mat, lowerdiag = TRUE, d,
                             Alpha = intervals$Alpha, Beta = intervals$Beta, delta)
```

WBS.uni.nonpar	<i>Wild binary segmentation for univariate nonparametric change points detection.</i>
----------------	---

Description

Perform wild binary segmentation for univariate nonparametric change points detection.

Usage

```
WBS.uni.nonpar(Y, s, e, Alpha, Beta, N, delta, level = 0)
```

Arguments

Y	A numeric matrix of observations with horizontal axis being time, and vertical axis being multiple observations on each time point.
s	A integer scalar of starting index.
e	A integer scalar of ending index.
Alpha	A integer vector of starting indices of random intervals.
Beta	A integer vector of ending indices of random intervals.
N	A integer vector representing number of multiple observations on each time point.
delta	A positive integer scalar of minimum spacing.
level	Should be fixed as 0.

Value

A list with the following structure:

S	A vector of estimated change points (sorted in strictly increasing order)
Dval	A vector of values of CUSUM statistic based on KS distance
Level	A vector representing the levels at which each change point is detected
Parent	A matrix with the starting indices on the first row and the ending indices on the second row

Author(s)

Oscar Hernan Madrid Padilla, Haotian Xu

References

Padilla, Yu, Wang and Rinaldo (2021) <doi:10.1214/21-EJS1809>.

See Also

[thresholdBS](#) for obtaining change points estimation, [tuneBSuninonpar](#) for a tuning version.

Examples

```
Y = t(as.matrix(c(rnorm(100, 0, 1), rnorm(100, 0, 10), rnorm(100, 0, 40))))
N = rep(1, 300)
M = 20
intervals = WBS.intervals(M = M, lower = 1, upper = ncol(Y))
temp = WBS.uni.nonpar(Y, 1, 300, intervals$Alpha, intervals$Beta, N, 5)
plot.ts(t(Y))
points(x = tail(temp$$[order(temp$Dval)], 4), y = Y[,tail(temp$$[order(temp$Dval)],4)], col = "red")
thresholdBS(temp, 2)
```

WBS.uni.rob	<i>Robust wild binary segmentation for univariate mean change points detection.</i>
-------------	---

Description

Perform a robust version of the wild binary segmentation method using Huber loss.

Usage

```
WBS.uni.rob(y, s, e, Alpha, Beta, K = 1.345, delta, level = 0)
```

Arguments

y	A numeric vector of observations.
s	A numeric scalar representing the starting index of an interval.
e	A numeric scalar representing the ending index of an interval.
Alpha	An integer vector of starting indices of random intervals.
Beta	An integer vector of ending indices of random intervals.
K	A numeric scalar representing the robustification parameter in the Huber loss.
delta	A positive integer scalar of minimum spacing.
level	Should be fixed as 0.

Value

An object of `class` "BS", which is a list with the following structure:

S	A vector of estimated change points (sorted in strictly increasing order).
Dval	A vector of values of CUSUM statistic based on KS distance.
Level	A vector representing the levels at which each change point is detected.
Parent	A matrix with the starting indices on the first row and the ending indices on the second row.

Author(s)

Mengchu Li & Haotian Xu

References

Fearnhead & Rigail (2019) <doi:10.1080/01621459.2017.1385466>.

See Also

[thresholdBS](#) for obtaining change points estimation.

WBS.univar	<i>Wild binary segmentation for univariate mean change points detection.</i>
------------	--

Description

Perform wild binary segmentation for univariate mean change points detection.

Usage

```
WBS.univar(y, s, e, Alpha, Beta, delta = 2, level = 0)
```

Arguments

y	A numeric vector of observations.
s	A integer scalar of starting index.
e	A integer scalar of ending index.
Alpha	A integer vector of starting indices of random intervals.
Beta	A integer vector of ending indices of random intervals.
delta	A positive integer scalar of minimum spacing.
level	Should be fixed as 0.

Value

An object of `class` "BS", which is a list with the following structure:

S	A vector of estimated change point locations (sorted in strictly increasing order).
Dval	A vector of values of CUSUM statistic.
Level	A vector representing the levels at which each change point is detected.
Parent	A matrix with the starting indices on the first row and the ending indices on the second row.

Author(s)

Haotian Xu

References

Wang, Yu and Rinaldo (2020) <doi:10.1214/20-EJS1710>.

See Also

[thresholdBS](#) for obtaining change points estimation, [tuneBSunivar](#) for a tuning version.

Examples

```

set.seed(0)
cpt_true = c(20, 50, 170)
y = rnorm(300) + c(rep(0,20),rep(2,30),rep(0,120),rep(2,130))
intervals = WBS.intervals(M = 300, lower = 1, upper = length(y))
temp = WBS.univar(y, 1, length(y), intervals$Alpha, intervals$Beta, delta = 5)
plot.ts(y)
points(x = tail(temp$S[order(temp$Dval)],4),
       y = y[tail(temp$S[order(temp$Dval)],4)], col = "red")
WBS_result = thresholdBS(temp, tau = 4)
print(WBS_result$BS_tree, "value")
plot(WBS_result$BS_tree)
print(WBS_result$BS_tree_trimmed, "value")
plot(WBS_result$BS_tree_trimmed)
cpt_hat = sort(WBS_result$cpt_hat[,1]) # the threshold tau is specified to be 4
Hausdorff.dist(cpt_hat, cpt_true)
cpt_LR = local.refine.univar(cpt_hat, y)
Hausdorff.dist(cpt_LR, cpt_true)

```

WBSIP.cov

Wild binary segmentation for covariance change points detection through Independent Projection.

Description

Perform wild binary segmentation for covariance change points detection through Independent Projection

Usage

```
WBSIP.cov(X, X_prime, s, e, Alpha, Beta, delta, level = 0)
```

Arguments

X	A numeric vector of observations.
X_prime	A numeric vector of observations which are independent copy of X.
s	A integer scalar of starting index.
e	A integer scalar of ending index.
Alpha	A integer vector of starting indices of random intervals.
Beta	A integer vector of ending indices of random intervals.
delta	A positive integer scalar of minimum spacing.
level	A parameter for tracking the level at which a change point is detected. Should be fixed as 0.

Value

An object of class "BS", which is a list with the following structure:

S	A vector of estimated change points (sorted in strictly increasing order)
Dval	A vector of values of CUSUM statistic based on KS distance
Level	A vector representing the levels at which each change point is detected
Parent	A matrix with the starting indices on the first row and the ending indices on the second row

Author(s)

Haotian Xu

References

Wang, Yu and Rinaldo (2021) <doi:10.3150/20-BEJ1249>.

See Also

[thresholdBS](#) for obtain change points estimation.

Examples

```
p = 10
A1 = gen.cov.mat(p, 1, "equal")
A2 = gen.cov.mat(p, 3, "power")
A3 = A1
set.seed(1234)
X = cbind(t(MASS::mvrnorm(50, mu = rep(0, p), A1)),
          t(MASS::mvrnorm(50, mu = rep(0, p), A2)),
          t(MASS::mvrnorm(50, mu = rep(0, p), A3)))
X_prime = cbind(t(MASS::mvrnorm(50, mu = rep(0, p), A1)),
                t(MASS::mvrnorm(50, mu = rep(0, p), A2)),
                t(MASS::mvrnorm(50, mu = rep(0, p), A3)))
intervals = WBS.intervals(M = 120, lower = 1, upper = dim(X)[2])
temp = WBSIP.cov(X, X_prime, 1, dim(X)[2], intervals$Alpha, intervals$Beta, delta = 5)
tau = sqrt(p*log(ncol(X)))*1.5
sort(thresholdBS(temp, tau)$cpt_hat[,1])
```

Index

aARC, 3
ARC, 5

BD_U, 6
BS.cov, 7, 67
BS.uni.nonpar, 8, 67, 71
BS.univar, 9, 67, 72

calibrate.online.network.missing, 10, 56
changepoints, 12
changepoints-package (changepoints), 12
CI.regression, 13
class, 7, 9, 10, 23–29, 73, 75, 76, 78, 79, 81, 82, 85, 86, 88
CV.search.DP.LR.regression, 15
CV.search.DP.poly, 16
CV.search.DP.regression, 17
CV.search.DP.univar, 19
CV.search.DP.VAR1, 20
CV.search.DPDU.regression, 21

DP.poly, 22
DP.regression, 23
DP.SEPP, 24
DP.univar, 26
DP.VAR1, 27
DPDU.regression, 28
DPDU2.regression, 29

gen.cov.mat, 30
gen.missing, 31
gen.piece.poly, 31
gen.piece.poly.noiseless, 32

Hausdorff.dist, 33
huber_mean, 34

kde.biwei.eval, 35
kde.epan.eval, 35
kde.eval, 36

kde.triwei.eval, 37

lambda.network.missing, 38
local.refine.CV.VAR1, 39, 52
local.refine.DPDU.regression, 40
local.refine.multi.nonpar.L2, 41
local.refine.multi.nonpar.L2.biwei, 42
local.refine.multi.nonpar.L2.epan, 44
local.refine.multi.nonpar.L2.triwei, 45
local.refine.network, 47
local.refine.poly, 48
local.refine.regression, 49
local.refine.univar, 50
local.refine.VAR1, 51
lowertri2mat, 52
LRV.regression, 53

online.network, 54
online.network.missing, 12, 55
online.univar, 57
online.univar.multi, 58

simu.change.regression, 59
simu.RDPG, 61
simu.SBM, 62
simu.SEPP, 63
simu.VAR1, 64
softImpute.network.missing, 65

thresholdBS, 8–10, 66, 74, 75, 77, 78, 80, 81, 83–86, 88
trim_interval, 67
tuneBSmultinonpar, 68, 74, 75, 77, 78, 80
tuneBSnonparRDPG, 69, 83
tuneBSuninonpar, 9, 70, 84
tuneBSunivar, 10, 71, 86

WBS.intervals, 72
WBS.multi.nonpar, 67, 68, 73
WBS.multi.nonpar.L2, 74

WBS.multi.nonpar.L2.biwei, [76](#)
WBS.multi.nonpar.L2.epan, [77](#)
WBS.multi.nonpar.L2.triwei, [79](#)
WBS.network, [67](#), [80](#)
WBS.nonpar.RDPG, [70](#), [82](#)
WBS.uni.nonpar, [67](#), [71](#), [83](#)
WBS.uni.rob, [85](#)
WBS.univar, [67](#), [72](#), [86](#)
WBSIP.cov, [67](#), [87](#)