

# Package ‘SpaCCr’

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**Type** Package

**Title** Spatial Convex Clustering

**Version** 0.1.0

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**Description** Genomic Region Detection via Spatial Convex Clustering. See <<https://arxiv.org/abs/1611.04696>> for details.

**Depends** R (>= 2.10)

**License** GPL-3

**LazyData** TRUE

**LinkingTo** Rcpp, RcppArmadillo

**Imports** abind, dplyr, ggplot2, parallel, Rcpp, tidy

**RoxygenNote** 5.0.1

**Suggests** testthat

**NeedsCompilation** yes

**Repository** CRAN

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## R topics documented:

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|                    |  |
|--------------------|--|
| CNVPlotSeriesMeans | <i>Plot subjects' copy number data with cluster means overlaid for a single chromosome</i> |
|--------------------|--|

---

### Description

Plot subjects' copy number data with cluster means overlaid for a single chromosome

### Usage

```
CNVPlotSeriesMeans(Location, X, Cluster, NSubj = 3, lowery = -1,
  upperry = 1)
```

### Arguments

|                 |   |
|-----------------|---|
| Location        | A vector of length p with chromosomal locations                   |
| X               | A variable (p) by subject (n) data matrix                         |
| Cluster         | A vector of length p with cluster labels                          |
| NSubj           | A positive integer; number of randomly selected subjects to plot. |
| lowery, upperry | Scalars; limits for y-axis  |

### Examples

NULL

---

|             |                                      |
|-------------|--------------------------------------|
| GetClusters | <i>Compute Clusters from fusions</i> |
|-------------|--------------------------------------|

---

### Description

Compute Clusters from fusions

### Usage

```
GetClusters(V)
```

### Arguments

|   |                         |
|---|-------------------------|
| V | An n by p-1 data matrix |
|---|-------------------------|

**Examples**

NULL

---

|            |  |
|------------|--|
| GetGammaCV | <i>Get optimal cross validated gamma values by various rules</i> |
|------------|--|

---

**Description**

Get optimal cross validated gamma values by various rules

**Usage**

```
GetGammaCV(ErrMat, rule = 1, gamma.seq)
```

**Arguments**

|           |   |
|-----------|---|
| ErrMat    | Matrix of cross validated errors outputted by GetCVErrMat   |
| rule      | A number indicating how optimal gamma should be chosen. 1 for minimum cv error, 2 for 1 standard error rule |
| gamma.seq | sequence of regularization parameters used for cross validation.  |

**Value**

A scalar. Optimal gamma selected by CV rule.

**Examples**

NULL

---

|              |  |
|--------------|--|
| GetParBlocks | <i>Function to compute blocks for parallization; should not be called directly</i> |
|--------------|--|

---

**Description**

Function to compute blocks for parallization; should not be called directly

**Usage**

```
GetParBlocks(X, w)
```

**Arguments**

|   |  |
|---|--|
| X | An n by p data matrix                      |
| w | A vector of positive scalars of length p-1 |

**Examples**

```
NULL
```

---

```
hello           Hello, World!
```

---

**Description**

Prints 'Hello, world!'.

**Usage**

```
hello()
```

**Examples**

```
hello()
```

---

```
methy           A subset of methylation data for 50 subjects
```

---

**Description**

A subset of methylation data for 50 subjects

**Usage**

```
methy
```

**Format**

A data frame with 1000 rows and 53 columns. Rows are variables and columns are:

**ProbeID** Varibale ID string

**Chromosome** Chromosome Number

**Genomic\_Coordinate** Probe location on chromosome in basepairs

**TCGA-A7-A0CE-11A-21D-A10Q-05** Subject ID

**TCGA-A7-A0CH-11A-32D-A10Q-05** Subject ID

**TCGA-A7-A0DB-11A-33D-A093-05** Subject ID

**TCGA-A7-A0DC-11A-41D-A10Q-05** Subject ID

**TCGA-BH-A0AY-11A-23D-A10Q-05** Subject ID

**TCGA-BH-A0BV-11A-31D-A10Q-05** Subject ID

**TCGA-BH-A0DZ-11A-22D-A10Q-05** Subject ID

TCGA-A2-A1FV-01A-11D-A13K-05 Subject ID  
TCGA-A2-A1FW-01A-11D-A13K-05 Subject ID  
TCGA-A2-A1FX-01A-11D-A13K-05 Subject ID  
TCGA-A2-A1G0-01A-11D-A13K-05 Subject ID  
TCGA-A2-A1G1-01A-21D-A13K-05 Subject ID  
TCGA-A2-A1G4-01A-11D-A13K-05 Subject ID  
TCGA-A2-A1G6-01A-11D-A13K-05 Subject ID  
TCGA-A7-A13G-01A-11D-A13K-05 Subject ID  
TCGA-A7-A13G-01B-04D-A22R-05 Subject ID  
TCGA-A7-A13G-11A-51D-A13T-05 Subject ID  
TCGA-AO-A1KO-01A-31D-A13K-05 Subject ID  
TCGA-AO-A1KP-01A-11D-A13K-05 Subject ID  
TCGA-AO-A1KQ-01A-11D-A13K-05 Subject ID  
TCGA-AO-A1KS-01A-11D-A13K-05 Subject ID  
TCGA-AO-A1KT-01A-11D-A13K-05 Subject ID  
TCGA-AQ-A1H2-01A-11D-A13K-05 Subject ID  
TCGA-AQ-A1H3-01A-31D-A13K-05 Subject ID  
TCGA-B6-A1KC-01A-11D-A13K-05 Subject ID  
TCGA-B6-A1KC-01B-11D-A161-05 Subject ID  
TCGA-B6-A1KF-01A-11D-A13K-05 Subject ID  
TCGA-B6-A1KN-01A-11D-A13K-05 Subject ID  
TCGA-BH-A1EN-01A-11D-A13K-05 Subject ID  
TCGA-BH-A1EN-11A-23D-A13T-05 Subject ID  
TCGA-BH-A1EX-01A-11D-A13K-05 Subject ID  
TCGA-BH-A1EY-01A-11D-A13K-05 Subject ID  
TCGA-BH-A1EY-11B-21D-A13T-05 Subject ID  
TCGA-BH-A1F2-01A-31D-A13K-05 Subject ID  
TCGA-BH-A1F2-11A-32D-A13T-05 Subject ID  
TCGA-BH-A1F5-01A-12D-A13K-05 Subject ID  
TCGA-BH-A1F5-11A-43D-A13T-05 Subject ID  
TCGA-BH-A1F6-01A-11D-A13K-05 Subject ID  
TCGA-BH-A1F6-11B-94D-A13T-05 Subject ID  
TCGA-BH-A1F8-01A-11D-A13K-05 Subject ID  
TCGA-BH-A1F8-11B-21D-A13T-05 Subject ID  
TCGA-BH-A1FB-01A-11D-A13K-05 Subject ID  
TCGA-BH-A1FB-11A-33D-A13T-05 Subject ID  
TCGA-BH-A1FC-01A-11D-A13K-05 Subject ID

TCGA-BH-A1FC-11A-32D-A13T-05 Subject ID  
 TCGA-BH-A1FD-01A-11D-A13K-05 Subject ID  
 TCGA-BH-A1FD-11B-21D-A13T-05 Subject ID  
 TCGA-BH-A1FE-01A-11D-A13K-05 Subject ID  
 TCGA-BH-A1FE-06A-11D-A212-05 Subject ID  
 TCGA-BH-A1FE-11B-14D-A13T-05 Subject ID

### Source

<http://cancergenome.nih.gov/>

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|                 |  |
|-----------------|--|
| MethyRegionPlot | <i>Plots methylation data by Genomic Coordinates for a given chromosomal region with cluster means overlayed for each subject.</i> |
|-----------------|--|

---

### Description

Plots methylation data by Genomic Coordinates for a given chromosomal region with cluster means overlayed for each subject.

### Usage

```
MethyRegionPlot(X, Coord, Cluster, SubjInd = 1:3, Start, End)
```

### Arguments

|         |  |
|---------|--|
| X       | A Subject by Probe data matrix for a single chromosome of CNV data       |
| Coord   | A vector of Genomic Coordinates for a single chromosome                  |
| Cluster | Cluster labels for each probe  |
| SubjInd | A vector of numeric indices corresponding to the Subjects to be plotted. |
| Start   | Genomic Coordinate minimum   |
| End     | Genomic Coordinate maximum   |

### Examples

```
library(dplyr)
library(tidyr)
data("methy")
methy <- methy[1:20,1:10]
Coordinates <- methy$Genomic_Coordinate
methy %>%
  tbl_df() %>%
  select(-Chromosome, -Genomic_Coordinate) %>%
  gather(Subject, Value, -ProbeID) %>%
  spread(ProbeID, Value) -> X
```

```

SubjectLabels <- X$Subject
X <- X[,-1] %>% as.matrix()
nsubj <- nrow(X)
nprobes <- ncol(X)
nweights <- choose(nprobes,2)
diff.vals <- diff(Coordinates)
too.far <- diff.vals > 20000
sig = 1/5e3
w.values <- exp(-sig*diff.vals)
w.values[too.far] = 0

verbose=TRUE
tol.base = 1e-4
tol.miss = 1e-4
max.iter.base=5000
max.iter.miss=500
ngam = 20
gamma.seq <- exp(seq(log(1e-1),log(1e1),length.out=ngam))
CVRes <- SpaCC_CV(X=t(scale(t(X),center=TRUE,scale=FALSE)),
                 w=w.values,
                 gamma.seq=gamma.seq,
                 nfolds=5,
                 nu=1/nsubj,
                 verbose=TRUE,
                 tol.base=tol.base,
                 tol.miss=tol.miss,
                 max.iter.base=max.iter.base,
                 max.iter.miss=max.iter.miss,
                 parallel=FALSE,frac = .1)
PlotCV(CVRes$ErrMat,gamma.seq = CVRes$gamma.seq,rule = 1)
best.gam <- GetGammaCV(CVRes$ErrMat,rule = 1,gamma.seq = CVRes$gamma.seq)
bo <-t(scale(t(X),center=TRUE,scale=FALSE))
bo[is.na(bo)] <- mean(bo,na.rm=TRUE)
Sol <- SpaCC_Missing(t(scale(t(X),center=TRUE,scale=FALSE)),
                   w.values,
                   gamma = best.gam,
                   nu=1/nsubj,
                   verbose=TRUE,
                   tol.base=tol.base,
                   tol.miss=tol.miss,
                   max.iter.base=max.iter.base,
                   max.iter.miss=max.iter.miss,
                   bo,
                   t(diff(t(bo))),
                   t(diff(t(bo))))

VThreshed <- Sol$V
clustsThreshed <- GetClusters(VThreshed)
NEstRegion <- length(unique(clustsThreshed$cluster))
NEstRegion
VThreshed <- ThreshV(Sol$V,X,mult = 1)
clustsThreshed <- GetClusters(VThreshed)
NEstRegion <- length(unique(clustsThreshed$cluster))
NEstRegion

```

```

start.coord <- 2e5
end.coord <- 4e5
MethyRegionPlot(X,Coordinates,clustsThreshed$cluster,SubjInd = 1:3,Start=start.coord,End=end.coord)

```

---

PlotCV

*A function for plotting cross validation errors*


---

### Description

A function for plotting cross validation errors

### Usage

```
PlotCV(ErrMat, rule = 2, gamma.seq)
```

### Arguments

|           |  |
|-----------|--|
| ErrMat    | A matrix of error outputted by SpaCC_CV        |
| rule      | An interger indicating which CV rule to choose |
| gamma.seq | The sequence of regularization parameters      |

### Examples

```

library(dplyr)
library(tidyr)
data("methy")
methy <- methy[1:20,1:10]
Coordinates <- methy$Genomic_Coordinate
methy %>%
  tbl_df() %>%
  select(-Chromosome,-Genomic_Coordinate) %>%
  gather(Subject,Value,-ProbeID) %>%
  spread(ProbeID,Value) -> X
SubjectLabels <- X$Subject
X <- X[,-1] %>% as.matrix()
nsubj <- nrow(X)
nprobes <- ncol(X)
nweights <- choose(nprobes,2)
diff.vals <- diff(Coordinates)
too.far <- diff.vals > 20000
sig = 1/5e3
w.values <- exp(-sig*diff.vals)
w.values[too.far] = 0

verbose=TRUE
tol.base = 1e-4
tol.miss = 1e-4
max.iter.base=5000
max.iter.miss=500

```



```

ngam = 20
gamma.seq <- exp(seq(log(1e-1),log(1e1),length.out=ngam))
CVRes <- SpaCC_CV(X=t(scale(t(X),center=TRUE,scale=FALSE)),
                 w=w.values,
                 gamma.seq=gamma.seq,
                 nolds=5,
                 nu=1/nsubj,
                 verbose=TRUE,
                 tol.base=tol.base,
                 tol.miss=tol.miss,
                 max.iter.base=max.iter.base,
                 max.iter.miss=max.iter.miss,
                 parallel=FALSE,frac = 1)
PlotCV(CVRes$ErrMat,gamma.seq = CVRes$gamma.seq,rule = 1)

```

---

|       |  |
|-------|--|
| SpaCC | <i>Base function for computing SpaCC solution for single regularization value.</i> |
|-------|--|

---

### Description

Base function for computing SpaCC solution for single regularization value.

### Usage

```
SpaCC(X, w, gamma, nu, verbose, tol, maxiter, Uinit, Vinit, Laminit)
```

### Arguments

|         |   |
|---------|---|
| X       | A subject (n) by probe (p) data matrix                              |
| w       | A vector weights for adjacent probes. Should have length nprobes -1 |
| gamma   | A scalar value for the regularization parameter                     |
| nu      | A scalar value for the step size in AMA algorithm                   |
| verbose | Logical value whether progress should be printed                    |
| tol     | A scalar value for convergence tolerance.                           |
| maxiter | Maximum number of iterations  |
| Uinit   | A matrix used for warm starts with U                                |
| Vinit   | A matrix used for warm start with V                                 |
| Laminit | A matrix used for warm starts with Lam                              |

### Value

An RcppArmadillo field object. Has three components, each holds the U,V, and Lam matrix for the current regularization

SpaCC\_CV

*Perform Cross Validation to select gamma/sparsity level***Description**

Perform Cross Validation to select gamma/sparsity level

**Usage**

```
SpaCC_CV(X, w, gamma.seq, nfolds = 5, nu = 1/nrow(X), verbose = FALSE,
  tol.base = 1e-04, tol.miss = 1e-04, max.iter.base = 5000,
  max.iter.miss = 500, parallel = FALSE, frac = 1)
```

**Arguments**

|               |   |
|---------------|---|
| X             | A subject (n) by variable (p) matrix; the data                            |
| w             | A vector of length p-1; weights for clustering                            |
| gamma.seq     | A vector of positive scalars; regularization parameter sequence           |
| nfolds        | A positive scalar; number of cross validation folds                       |
| nu            | A positive scalar; augmented Lagrangian paramter                          |
| verbose       | Logical; should messages be printed?                                      |
| tol.base      | A small positive scalar; convergence tolerance for base SpaCC problem.    |
| tol.miss      | A small positive scalar; convergence tolerance for missing data problem.  |
| max.iter.base | A positive integer; maximum number of iterations for base SpaCC problem   |
| max.iter.miss | A positive integer; maximum number of iterations for missing data problem |
| parallel      | A logical; should CV paths be done in parallel?                           |
| frac          | A positive scalar between 0 and 1; fraction of hold out set to utilize    |

**Value**

A list with elements: ErrMat - a length(gamma.seq) by nfold matrix containing error on out of fold data; SpMat - a length(gamma.seq) by nfold matrix containing sparsity levels; gamma.seq - original gamma.seq sorted largest to smallest

**Examples**

```
library(dplyr)
library(tidyr)
data("methy")
methy <- methy[1:20,1:10]
Coordinates <- methy$Genomic_Coordinate
methy %>%
  tbl_df() %>%
  select(-Chromosome, -Genomic_Coordinate) %>%
  gather(Subject, Value, -ProbeID) %>%
```

```

    spread(ProbeID,Value) -> X
    SubjectLabels <- X$Subject
    X <- X[,-1] %>% as.matrix()
    nsubj <- nrow(X)
    nprobes <- ncol(X)
    nweights <- choose(nprobes,2)
    diff.vals <- diff(Coordinates)
    too.far <- diff.vals > 20000
    sig = 1/5e3
    w.values <- exp(-sig*diff.vals)
    w.values[too.far] = 0

    verbose=TRUE
    tol.base = 1e-4
    tol.miss = 1e-4
    max.iter.base=5000
    max.iter.miss=500
    ngam = 20
    gamma.seq <- exp(seq(log(1e-1),log(1e1),length.out=ngam))
    CVRes <- SpaCC_CV(X=t(scale(t(X),center=TRUE,scale=FALSE)),
                     w=w.values,
                     gamma.seq=gamma.seq,
                     nolds=5,
                     nu=1/nsubj,
                     verbose=TRUE,
                     tol.base=tol.base,
                     tol.miss=tol.miss,
                     max.iter.base=max.iter.base,
                     max.iter.miss=max.iter.miss,
                     parallel=FALSE,frac = 1)

```

---

SpaCC\_Methy

*Performs Spatial Convex Clustering for methylation data*


---

## Description

Performs Spatial Convex Clustering for methylation data

## Usage

```

SpaCC_Methy(X, Coordinates, gamma.seq, dist.cutoff = 20000, sig = 1/5000,
            weights = NULL, center = TRUE, scale = FALSE, nolds = 5, nu = NULL,
            tol.base = 1e-04, tol.miss = 1e-04, max.iter.base = 5000,
            max.iter.miss = 500, frac = 0.1, parallel = FALSE, gam.rule = 2,
            thresh.mult = 1, thresh.value = NULL)

```

## Arguments

X                    A subject (n) by variable (p) matrix; the data

|               |  |
|---------------|--|
| Coordinates   | a vector listing genomic coordinates                   |
| gamma.seq     | a vector of regularization parameters                  |
| dist.cutoff   | maximum distance at which probes should be regularized |
| sig           | positive scalar controlling spatial weight decay       |
| weights       | a vector of spatial weights                            |
| center        | should data be centered                                |
| scale         | should data be scaled                                  |
| nfolds        | number of folds for cross validation                   |
| nu            | parameter for augmented lagrangian                     |
| tol.base      | tolerance level for base function                      |
| tol.miss      | tolerance for missing function                         |
| max.iter.base | maximum number of iterations for base function         |
| max.iter.miss | maximum number of iterations for missing function      |
| frac          | fraction of fold to use for cross validation           |
| parallel      | should algorithm be run in parallel                    |
| gam.rule      | cross validation rule                                  |
| thresh.mult   | multiplier for threshold value                         |
| thresh.value  | value of threshold                                     |

### Value

Labels a vector of cluster labels

### Examples

```

data("methy")
methy <- methy[1:20,1:10]
library(dplyr)
library(tidyr)
Coordinates <- methy$Genomic_Coordinate
methy %>%
  tbl_df() %>%
  select(-Chromosome, -Genomic_Coordinate) %>%
  gather(Subject, Value, -ProbeID) %>%
  spread(ProbeID, Value) -> X
SubjectLabels <- X$Subject
X <- X[,-1] %>% as.matrix()
verbose=TRUE
tol.base = 1e-4
tol.miss = 1e-4
max.iter.base=5000
max.iter.miss=500
ngam = 20
gamma.seq <- exp(seq(log(1e-1), log(1e1), length.out=ngam))
ClusterLabels <- SpaCC_Methy(X = X, Coordinates = Coordinates, gamma.seq = gamma.seq)

```

SpaCC\_Missing

*Solve Spatial Convex Clustering problem for missing data***Description**

Solve Spatial Convex Clustering problem for missing data

**Usage**

```
SpaCC_Missing(X, w, gamma, nu = 1/nrow(X), verbose = FALSE,
  tol.base = 1e-04, tol.miss = 1e-04, max.iter.base = 5000,
  max.iter.miss = 500, Uinit, Vinit, Laminit)
```

**Arguments**

|               |   |
|---------------|---|
| X             | A subject (n) by variable (p) matrix; the data                            |
| w             | A vector of length p-1; weights for clustering                            |
| gamma         | A positive scalar; regularization parameter                               |
| nu            | A positive scalar; augmented Lagrangian parameter                         |
| verbose       | Logical; should messages be printed?                                      |
| tol.base      | A small positive scalar; convergence tolerance for base SpaCC problem.    |
| tol.miss      | A small positive scalar; convergence tolerance for missing data problem.  |
| max.iter.base | A positive integer; maximum number of iterations for base SpaCC problem   |
| max.iter.miss | A positive integer; maximum number of iterations for missing data problem |
| Uinit         | An n by p matrix; initial value for U                                     |
| Vinit         | An n by p-1 matrix; initial value for V                                   |
| Laminit       | An n by p-1 matrix; initial value for Lam                                 |

**Value**

A list with elements U, V, and Lam

**Examples**

```
library(dplyr)
library(tidyr)
data("methy")
methy <- methy[1:20,1:10]
Coordinates <- methy$Genomic_Coordinate
methy %>%
  tbl_df() %>%
  select(-Chromosome, -Genomic_Coordinate) %>%
  gather(Subject, Value, -ProbeID) %>%
  spread(ProbeID, Value) -> X
SubjectLabels <- X$Subject
```

```

X <- X[,-1] %>% as.matrix()
X[1:5,1:5]
nsubj <- nrow(X)
nprobes <- ncol(X)
nweights <- choose(nprobes,2)
diff.vals <- diff(Coordinates)
too.far <- diff.vals > 20000
sig = 1/5e3
w.values <- exp(-sig*diff.vals)
w.values[too.far] = 0

verbose=TRUE
tol.base = 1e-4
tol.miss = 1e-4
max.iter.base=5000
max.iter.miss=500
bo <-t(scale(t(X),center=TRUE,scale=FALSE))
bo[is.na(bo)] <- mean(bo,na.rm=TRUE)
best.gam = 1
Sol <- SpaCC_Missing(t(scale(t(X),center=TRUE,scale=FALSE)),
                    w.values,
                    gamma = best.gam,
                    nu=1/nsubj,
                    verbose=TRUE,
                    tol.base=tol.base,
                    tol.miss=tol.miss,
                    max.iter.base=max.iter.base,
                    max.iter.miss=max.iter.miss,
                    bo,
                    t(diff(t(bo))),
                    t(diff(t(bo))))

```

---

SpaCC\_Path

*Solve Spatial Convex Clustering problem for path of regularization parameters*


---

### Description

Solve Spatial Convex Clustering problem for path of regularization parameters

### Usage

```

SpaCC_Path(X, w, gamma.seq, nu = 1/nrow(X), verbose = FALSE,
           tol.base = 1e-04, tol.miss = 1e-04, max.iter.base = 5000,
           max.iter.miss = 500)

```

### Arguments

X                    A subject (n) by variable (p) matrix; the data

|               |   |
|---------------|---|
| w             | A vector of length p-1; weights for clustering                            |
| gamma.seq     | A vector of positive scalars; regularization parameter sequence           |
| nu            | A positive scalar; augmented Lagrangian paramter                          |
| verbose       | Logical; should messages be printed?                                      |
| tol.base      | A small positive scalar; convergence tolerance for base SpaCC problem.    |
| tol.miss      | A small positive scalar; convergence tolerance for missing data problem.  |
| max.iter.base | A positive integer; maximum number of iterations for base SpaCC problem   |
| max.iter.miss | A positive integer; maximum number of iterations for missing data problem |

**Value**

A list with elements UPath, VPath, LamPath, and gamma.seq

**Examples**

NULL

---

SpaCC\_Path\_Parallel *Solve Spatial Convex Clustering problem for path of regularization parameters in parallel*

---

**Description**

Solve Spatial Convex Clustering problem for path of regularization parameters in parallel

**Usage**

```
SpaCC_Path_Parallel(X, w, gamma.seq, nu = 1/nrow(X), verbose = FALSE,
  tol.base = 1e-04, tol.miss = 1e-04, max.iter.base = 5000,
  max.iter.miss = 500, ncores = 2)
```

**Arguments**

|               |   |
|---------------|---|
| X             | A subject (n) by variable (p) matrix; the data                            |
| w             | A vector of length p-1; weights for clustering                            |
| gamma.seq     | A vector of positive scalars; regularization parameter sequence           |
| nu            | A positive scalar; augmented Lagrangian paramter                          |
| verbose       | Logical; should messages be printed?                                      |
| tol.base      | A small positive scalar; convergence tolerance for base SpaCC problem.    |
| tol.miss      | A small positive scalar; convergence tolerance for missing data problem.  |
| max.iter.base | A positive integer; maximum number of iterations for base SpaCC problem   |
| max.iter.miss | A positive integer; maximum number of iterations for missing data problem |
| ncores        | A positive integer; number of cores to use                                |

**Value**

A list with elements UPath, VPath, LamPath, and gamma.seq

**Examples**

NULL

---

|         |                              |
|---------|------------------------------|
| ThreshV | <i>Threshold differences</i> |
|---------|------------------------------|

---

**Description**

Threshold differences

**Usage**

ThreshV(V, X, mult = 1, thresh.value = NULL)

**Arguments**

|              |  |
|--------------|--|
| V            | an n x p-1 matrix of differences         |
| X            | an n x p matrix                          |
| mult         | scalar to multiply standard deviation    |
| thresh.value | optional user specified threshold value. |

**Value**

VThreshed an n x p-1 matrix of thresholded differences

**Examples**

NULL



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