## Package 'biomvRCNS'

July 15, 2025

Type Package

Title Copy Number study and Segmentation for multivariate biological data

**Version** 1.49.0

Date 2021-11-20

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**Description** In this package, a Hidden Semi Markov Model (HSMM) and one homogeneous segmentation model are designed and implemented for segmentation genomic data, with the aim of assisting in transcripts detection using high throughput technology like RNA-seq or tiling array, and copy number analysis using aCGH or sequencing.

License GPL (>= 2)

LazyLoad yes

Imports methods, mvtnorm

Depends IRanges, GenomicRanges, Gviz

Suggests cluster, parallel, GenomicFeatures, dynamicTreeCut, Rsamtools, TxDb.Hsapiens.UCSC.hg19.knownGene

- **biocViews** aCGH, CopyNumberVariation, Microarray, Sequencing, Visualization, Genetics
- git\_url https://git.bioconductor.org/packages/biomvRCNS
- git\_branch devel

git\_last\_commit 4524de5

- git\_last\_commit\_date 2025-04-15
- **Repository** Bioconductor 3.22

Date/Publication 2025-07-15

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biomvRCNS-class Class "biomvRCNS"

#### Description

The default object class returned by biomvRhsmm, biomvRseg and biomvRmgmr

#### **Objects from the Class**

Objects can be created by calls of the form new("biomvRCNS", ...).

#### Slots

- x: Object of class "GRanges", with range information either from real positional data or just indices, with input data matrix stored in the meta columns. Additional meta columns for the estimated states and associated probabilities for each sample will also be appended following the input data matrix when using biomvRhsmm.
- res: Object of class "GRanges", each range represent one continuous segment identified, with sample name slot 'SAMPLE' and segment mean slot 'MEAN' stored in the meta columns
- param: Object of class "list", list of all parameters used in the corresponding model.

#### Methods

plot signature(x = "biomvRCNS", y = "ANY"): ...

show signature(object = "biomvRCNS"): ...

#### Examples

showClass("biomvRCNS")

biomvRGviz

## Description

This function could be called to plot segmentation output, together with the input signal and optional annotation. By default resulting image will be printed to file. The plot method for class biomvRCNS-class also calls this method. See the vignette for a more complete example.

## Usage

biomvRGviz(exprgr, gmgr = NULL, prange = NULL, regionID = NULL, seggr = NULL, plotstrand = "+", eps =

#### Arguments

exprgr	a GRanges object with one numeric column for the segmentation input signal in its meta DataFrame
gmgr	an optional GRanges object for the annotation, which must have one column named 'TYPE' in its meta DataFrame
prange	an optional vector defining the scope of the plot, the first 3 elements must be formatted as c('seqname', 'start_position', 'end_position')
regionID	a character for the name of the plotted region or gene name or other identifier, will be used in the title of the plot and the output file name
seggr	a GRanges object for the segmentation output, which must have one column named 'STATE' in its meta DataFrame
plotstrand	select which strand to plot, possible values are '+', '-', '*'
eps	whether to output EPS file using postscript, if FALSE then PDF files for each sequence will be generated to the current working folder.
tofile	whether to output graphics file, if FALSE then will plot on the active device and have the trackList returned.
	other arguments for plot, like main, ylab, cex, or height and width for graphic device.

## Details

See the vignette for more details and examples.

#### Value

Plot graph on the active device or output to EPS/PDF file.

## Author(s)

Yang Du

## Examples

```
data(coriell)
x<-coriell[coriell[,2]==1,]
xgr<-GRanges(seqnames=paste('chr', x[,2], sep=''), IRanges(start=x[,3], width=1, names=x[,1]))
values(xgr)<-DataFrame(x[,4:5], row.names=NULL)
xgr<-xgr[order(xgr)]

J<-2; maxk<-50
# a uniform inital sojourn, not utilizing positional information
soj<-list(J=J, maxk=maxk, type='gamma', d=cbind(dunif(1:maxk, 1, maxk), dunif(1:maxk, 1, maxk)))
soj$D <- sapply(1:J, function(j) rev(cumsum(rev(soj$d[1:maxk,j])))
sample<-colnames(coriell)[5]
runout<-hsmmRun(matrix(values(xgr)[,sample]), sample, xgr, soj, emis=list(type='norm', mu=quantile(unlist(x
biomvRGviz(exprgr=xgr, seggr=runout$res, tofile=FALSE)</pre>
```

biomvRhsmm	Estimating the most likely state sequence using Hidden Semi Markov Model

## Description

The batch function of building Hidden Semi Markov Model (HSMM) to estimate the most likely state sequences for multiple input data series.

#### Usage

biomvRhsmm(x, maxk=NULL, maxbp=NULL, J=3, xPos=NULL, xRange=NULL, usePos='start', emis.type='norm'

#### Arguments

x	input data matrix, or a GRanges object with input stored in the meta DataFrame, assume ordered.
maxk	maximum length of stay for the sojourn distribution
maxbp	maximum length of stay in bp for the sojourn distribution, given positional in- formation specified in xPos / xRange
J	number of states
xPos	a vector of positions for each x row
xRange	a IRanges/GRanges object, same length as x rows
usePos	character value to indicate whether the 'start', 'end' or 'mid' point position should be used to estimate the sojourn distribution
emis.type	type of the emission distribution, only the following types are supported: 'norm', 'mvnorm', 'pois', 'nbinom', 'mvt', 't'
com.emis	whether to set a common emission prior across different seqnames. if TRUE, the emission will not be updated during individual runs.
xAnno	a optional TxDb / GRanges / GRangesList / list object used in sojournAnno to infer parameters for the sojourn distribution
soj.type	type of the sojourn distribution, only the following types are supported: 'non-para', 'gamma', 'pois', 'nbinom'

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q.alpha	a quantile factor controlling the estimated prior for the mean of the emission of each states, seq(from=q.alpha, to=1-q.alpha, length.out=J)
r.var	a ratio factor controlling the estimated prior for the variance / covariance struc- ture of each states. A value larger than 1 tend to allow larger variation in extreme states; a value smaller than 1 will decrease the probability of having extreme state
useMC	TRUE if mclapply should be used to speed up the calculation, use options(mc.cores=n) to set number of parallel processes
cMethod	C algorithm used for the most likely state sequence, 'F-B' or 'Viterbi'
maxit	max iteration of the EM run with Forward-Backward algorithm
maxgap	max distance between neighbouring feature to consider a split
tol	tolerance level of the likelihood change to terminate the EM run
grp	vector of group assignment for each sample, with a length the same as columns in the data matrix, samples within each group would be processed simultane- ously if a multivariate emission distribution is available
cluster.m	clustering method for prior grouping, possible values are 'ward','single','complete','average','mcquit
avg.m	method to calculate average value for each segment, 'median' or 'mean' possi- bly trimmed
prior.m	method to select emission prior for each state, 'quantile' uses different levels of quantile; the 'cluster' method uses clara function from cluster
trim	the fraction (0 to $0.5$ ) of observations to be trimmed from each end of x before the mean is computed. Values of trim outside that range are taken as the nearest endpoint.
na.rm	TRUE if NA value should be ignored

## Details

This is the batch function of building Hidden Semi Markov Model (HSMM) to estimating the most likely state sequences for multiple input data series. The function will sequentially process each region identified by the distinctive seqnames in x or in xRange if available, or assuming all data from the same region. A second layer of stratification is introduced by the argument grp, which could be used to reflect experimental design. The assumption is that profiles from the same group could be considered homogeneous, thus processed together if emis.type is compatible (currently only with 'mvnorm'). Argument for the sojourn density will be initialized as flat prior or estimated from other data before calling the work horse function hsmmRun. Then for each batch run results will be combined and eventually a biomvRCNS-class object will be returned. See the vignette for more details and examples.

## Value

A biomvRCNS-class object:

x:	Object of class "GRanges", with range information either from real positional data or just indices, with input data matrix stored in the meta columns. Additional meta columns for the estimated states and associated probabilities for each sample or group will also be appended following the input data matrix.
res:	Object of class "GRanges", each range represent one continuous segment iden- tified, with sample name slot 'SAMPLE', estimated state slot 'STATE' and seg- ment mean slot 'MEAN' stored in the meta columns
param:	Object of class "list", list of all parameters used in the model run, plus the re-estimated emission and sojourn parameters.

biomvRmgmr

#### Author(s)

Yang Du

#### References

Guedon, Y. (2003). Estimating hidden semi-Markov chains from discrete sequences. Journal of Computational and Graphical Statistics, 12(3), 604-639.

#### See Also

biomvRseg

#### Examples

```
data(coriell)
xgr<-GRanges(seqnames=paste('chr', coriell[,2], sep=''), IRanges(start=coriell[,3], width=1, names=coriell[
values(xgr)<-DataFrame(coriell[,4:5], row.names=NULL)
xgr<-sort(xgr)
reshsmm</pre>-biomvRhsmm(x=xgr, maxbp=4E4, J=3, soj.type='gamma', emis.type='norm', grp=c(1,2))
## access model parameters
reshsmm@param$soj.par
reshsmm@param$emis.par
```

```
## states assigned and associated probabilities
mcols(reshsmm@x)[,-(1:2)]
```

biomvRmgmr	Batch process multiple sequences and samples using max-gap-min-
	run algorithm for 2 states segmentation

## Description

This is a wrapper function for batch processing multiple sequences and samples using max-gapmin-run algorithm for 2 states segmentation

#### Usage

```
biomvRmgmr(x, xPos=NULL, xRange=NULL, usePos='start', cutoff=NULL, q=0.9, high=TRUE, minrun=5, maxg
```

#### Arguments

x	input data matrix, or a GRanges object with input stored in the meta DataFrame, assume ordered.
xPos	a vector of positions for each x row
xRange	a IRanges/GRanges obejct, same length as x rows
usePos	character value to indicate whether the 'start', 'end' or 'mid' point position should be used
cutoff	threshold level above which is considered extreme
q	relative quantile threshold level instead of absolute value for the cutoff

## biomvRmgmr

high	TRUE if the cutoff or q here is the lower bound and values greater than the threshold are considered
minrun	minimum run length for the resulting segments
maxgap	maximum genomic distance below which two adjacent qualified tiles can be joined
splitLen	numeric value, maximum length of segments, split if too long
poolGrp	TRUE if samples within the same group should be pooled using median for each feature
grp	vector of group assignment for each sample, with a length the same as columns in the data matrix, samples within each group would be processed simultane- ously if a multivariate emission distribution is available
cluster.m	clustering method for prior grouping, possible values are 'ward','single','complete','average','mcquit
avg.m	method to calculate average value for each segment, 'median' or 'mean' possi- bly trimmed
trim	the fraction (0 to 0.5) of observations to be trimmed from each end of x before the mean is computed. Values of trim outside that range are taken as the nearest endpoint.
na.rm	TRUE if NA value should be ignored

## Details

This is the batch function to apply maxGapminRun multiple sequence.

## Value

A biomvRCNS-class object:

x:	Object of class "GRanges", with range information either from real positional data or just indices, with input data matrix stored in the meta columns.
res:	Object of class "GRanges", each range represent one continuous segment iden- tified, with sample name slot 'SAMPLE' and segment mean slot 'MEAN' stored in the meta columns
param:	Object of class "list", list of all parameters used in the model run.

## Author(s)

Yang Du

## See Also

biomvRhsmm maxGapminRun

```
data(coriell)
xgr<-GRanges(seqnames=paste('chr', coriell[,2], sep=''), IRanges(start=coriell[,3], width=1, names=coriell[,
values(xgr)<-DataFrame(coriell[,4:5], row.names=NULL)
xgr<-xgr[order(xgr)]
resseg<-biomvRmgmr(x=xgr, minrun=3000, maxgap=1500, q=0.9, grp=c(1,2))</pre>
```

#### biomvRseg

## Description

The function will perform a two stage segmentation on multi-sample genomic data from array experiment or high throughput sequencing data.

## Usage

biomvRseg(x, maxk=NULL, maxbp=NULL, maxseg=NULL, xPos=NULL, xRange=NULL, usePos='start', family='n

## Arguments

х		input data matrix, or a GRanges object with input stored in the meta DataFrame
ma	axk	maximum length of a segment
ma	axbp	maximum length of a segment in bp, given positional information specified in xPos / xRange / or $x$
ma	axseg	maximum number of segment the function will try
xF	Pos	a vector of positions for each x row
xF	Range	a IRanges/GRanges object, same length as x rows
us	sePos	character value to indicate whether the 'start', 'end' or 'mid' point position should be used
fa	amily	family of x distribution, only the following types are supported: 'norm', 'nbi- nom', 'pois'
pe	enalty	penalty method used for determining the optimal number of segment using like- lihood, possible values are 'none', AIC', 'AICc', 'BIC', 'SIC', 'HQIC', 'mBIC'
t١	woStep	TRUE if a second stage merging will be performed after the initial group seg- mentation
se	egDisp	TRUE if a segment-wise estimation of dispersion parameter rather than using a overall estimation
us	seMC	TRUE if mclapply should be used to speed up the calculation for nbinom dis- persion estimation
us	seSum	TRUE if using grand sum across sample / x columns, like in the tilingArray solution
С	omVar	TRUE if assuming common variance across samples (x columns)
ma	axgap	max distance between neighbouring feature to consider a split
to	ol	tolerance level of the likelihood change to determining the termination of the EM run
gı	rp	vector of group assignment for each sample, with a length the same as columns in the data matrix, samples within each group would be processed simultane- ously if a multivariate emission distribution is available
c	luster.m	clustering method for prior grouping, possible values are 'ward','single','complete','average','mcquit
a	vg.m	method to calculate average value for each segment, 'median' or 'mean' possi- bly trimmed

#### biomvRseg

trim	the fraction $(0 \text{ to } 0.5)$ of observations to be trimmed from each end of x before the mean is computed. Values of trim outside that range are taken as the nearest endpoint.
na.rm	TRUE if NA value should be ignored

## Details

A homogeneous segmentation algorithm, using dynamic programming like in tilingArray; however capable of handling count data from sequencing.

### Value

A biomvRCNS-class object:

x:	Object of class "GRanges", with range information either from real positional data or just indices, with input data matrix stored in the meta columns.
res:	Object of class "GRanges", each range represent one continuous segment iden- tified, with sample name slot 'SAMPLE' and segment mean slot 'MEAN' stored in the meta columns
param:	Object of class "list", list of all parameters used in the model run.

#### References

Piegorsch, W. W. (1990). Maximum likelihood estimation for the negative binomial dispersion parameter. Biometrics, 863-867.

Picard,F. et al. (2005) A statistical approach for array CGH data analysis. BMC Bioinformatics, 6, 27.

Huber, W. et al. (2006) Transcript mapping with high density oligonucleotide tiling arrays. Bioinformatics, 22, 1963-1970.

Zhang, N. R. and Siegmund, D. O. (2007). A Modified Bayes Information Criterion with Applications to the Analysis of Comparative Genomic Hybridization Data. Biometrics 63 22-32.

Robinson MD and Smyth GK (2008). Small-sample estimation of negative binomial dispersion, with applications to SAGE data. Biostatistics, 9, 321-332

#### See Also

biomvRhsmm

```
data(coriell)
xgr<-GRanges(seqnames=paste('chr', coriell[,2], sep=''), IRanges(start=coriell[,3], width=1, names=coriell[
values(xgr)<-DataFrame(coriell[,4:5], row.names=NULL)
xgr<-xgr[order(xgr)]
resseg<-biomvRseg(x=xgr, maxbp=4E4, maxseg=10, family='norm', grp=c(1,2))</pre>
```

coriell

#### Description

These are two data array CGH studies sets of Corriel cell lines taken from the reference below.

#### Format

A data frame containing five variables: first is clone name, second is clone chromosome, third is clone position, fourth and fifth are log2ratio for two cell lines.

#### References

http://www.nature.com/ng/journal/v29/n3/suppinfo/ng754\\_S1.html

Snijders et al., Assembly of microarrays for genome-wide measurement of DNA copy number, Nature Genetics, 2001

encodeTP53

mapped RNA-seq data from ENCODE

#### Description

The data contains gene expression and transcript annotations in the region of the human TP53 gene (region (chr17:7,560,001-7,610,000 from the Human February 2009 (GRCh37/hg19) genome assembly), which is part of the long RNA-seq data generated by ENCODE/Cold Spring Harbor Lab, containing 2 cell types (GM12878 and K562) with 2 replicates each.

The alignment files were pulled from UCSC (http://hgdownload.cse.ucsc.edu/goldenPath/hg19/encodeDCC/wgEncodeC And subsequently reads were counted in each non-overlapping 25bp window for the region (chr17:7,560,001-7,610,000). The example code to generate this count GRanges is available in the vignette.

The regional annotation of TP53 RNAs isoforms were derived from the ENCODE Gene Annotations (GENCODE), sub-setted to only isoforms of TP53 gene. http://hgdownload.cse.ucsc.edu/goldenPath/hg19/encodeDCC/wgEncodeGencodeV4/wgEncodeGencodeManualV4.gtf.gz).

This dataset is used in the package vignette to illustrate a use case of transcript detection.

#### Format

Containing two GRanges objects, one for the sample count and one for the regional annotation of gene TP53

#### References

http://dx.doi.org/10.1371%2Fjournal.pbio.1001046 The ENCODE Project Consortium (2011) A User's Guide to the Encyclopedia of DNA Elements (ENCODE). PLoS Biol 9(4): e1001046. doi:10.1371/journal.pbio.1001046

hsmmRun

## Description

This is the working horse of the biomvRhsmm

#### Usage

hsmmRun(x, xid="sampleid", xRange, soj, emis, cMethod='F-B', maxit=1, maxgap=Inf, tol=1e-06, avg.m=

#### Arguments

x	input data matrix or vector, ordered wrt. position
xid	name of the sample
xRange	a IRanges/GRanges object, same length as x rows
soj	a list object containing the relevant sojourn distribution parameters
emis	a list object containing the relevant emission distribution parameters
cMethod	C algorithm used for the most likely state sequence, 'F-B' or 'Viterbi'
maxit	max iteration of the EM run with Forward-Backward algorithm
maxgap	max distance between neighbouring feature to consider a split
tol	tolerance level of the likelihood change to terminate the EM run
avg.m	method to calculate average value for each segment, 'median' or 'mean' possi- bly trimmed
trim	the fraction $(0 \text{ to } 0.5)$ of observations to be trimmed from each end of x before the mean is computed. Values of trim outside that range are taken as the nearest endpoint.
na.rm	TRUE if NA value should be ignored
com.emis	whether to set a common emission prior across different seqnames. if TRUE, the emission will not be updated during individual runs.

## Details

The function fits a Hidden-semi Markov model for the input data matrix / vector, which should contains ordered data from a continuous region on one chromosome. The model will start with flat prior for the initial state probability and transition probability, while emission parameter for each state will be estimated using different quantiles of the input controlled by argument q. alpha and r.var. Argument for the sojourn density should be provided via the list object soj, which is either initialized as flat prior or estimated from other data in a previous call. The positional information in the xRange is used for the optional spiting of physically distant features and construction of returning GRanges object res.

## Value

a list object	,
---------------	---

yhat	a "Rle" object, the most likely state sequence, same length as x rows number
ур	"Rle", the associated state probability, same length as x rows number
res	Object of class "GRanges", each range represent one continuous segment iden- tified, with sample name slot 'SAMPLE', estimated state slot 'STATE' and seg- ment mean slot 'MEAN' stored in the meta columns

#### Author(s)

Yang Du

#### References

Guedon, Y. (2003). Estimating hidden semi-Markov chains from discrete sequences. Journal of Computational and Graphical Statistics, 12(3), 604-639.

#### See Also

#### biomvRhsmm

## Examples

```
data(coriell)
# select only chr1
x<-coriell[coriell[,2]==1,]
xgr<-GRanges(seqnames=paste('chr', x[,2], sep=''), IRanges(start=x[,3], width=1, names=x[,1]))
values(xgr)<-DataFrame(x[,4:5], row.names=NULL)
xgr<-xgr[order(xgr)]

J<-2 ; maxk<-50
# a uniform initial sojourn, not utilizing positional information, just the index
soj<-list(J=J, maxk=maxk, type='gamma', d=cbind(dunif(1:maxk, 1, maxk), dunif(1:maxk, 1, maxk)))
soj$D <- sapply(1:J, function(j) rev(cumsum(rev(soj$d[1:maxk,j])))
# run 1 sample only, Coriell.13330
sample<-colnames(coriell)[5]
runout<-hsmmRun(matrix(values(xgr)[,sample]), sample, xgr, soj, emis=list(type='norm', mu=range(x[,4:5]), values(xgr)[, sample])</pre>
```

maxGapminRun M	Max-gap-min-run	algorithm for 2	2 states segmentation
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#### Description

A custom Max-gap-min-run implementation using physical position for gap and run length calculation.

## Usage

```
maxGapminRun(x, xPos = NULL, xRange = NULL, cutoff = NULL, q = 0.9, high=TRUE, minrun = 5, maxgap = 2,
```

## Arguments

х	a numeric vector for the input signal
xPos	a numeric vector, same length as x, carrying positional information for each element of x
xRange	an IRanges object, same length as x, carrying range information for each element of x
cutoff	numeric value used as cut-off, optional if q is specified
q	numeric value used to derive cut-off of x, as the q quantile of x , optional if cutoff is specified
high	TRUE if the cutoff or q here is the lower bound and values greater than the threshold are considered
minrun	minimum run length for the resulting segments
maxgap	maximum genomic distance below which two adjacent qualified tiles can be joined
splitLen	numeric value, maximum length of segments, split if too long
na.rm	TRUE if NA value should be ignored

## Details

A custom Max-gap-min-run implementation using physical position for gap and run length calculation.

## Value

a list of segment starts and ends indices

IS	the start index for each segment
IE	the end index for each segment
CUTOFF	the cutoff value used in the run
MG	the parameter value for maxgap
MR	the parameter value for minrun
SPL	the parameter value for ${\tt splitLen}$

## Author(s)

Yang Du

## See Also

biomvRhsmm biomvRseg biomvRmgmr

```
x<-rpois(50, 10)
xpos<-rnorm(50, 300, 100)
xpos<-xpos[order(xpos)]
maxGapminRun(x, xpos, cutoff=9.5, maxgap=30, minrun=100)</pre>
```

regionSegAlphaNB

#### Description

Estimate matrix of dispersion parameter alpha (size) used in regionSegCost for negative binomial distributed x.

#### Usage

```
regionSegAlphaNB(x, maxk = NULL, segs = NULL, useMC = FALSE, tol=1e-06)
```

#### Arguments

х	The input data matrix or vector
maxk	Maximum number of index to search forward
segs	Starting indices (excluding 1) for the candidate segments, for the second stage model, maxk will be overridden with length(segs)+1.
useMC	TRUE if mclapply should be used to speed up the calculation
tol	tolerance level for the convergence criteria in the maximum likelihood estima- tion of negative binomial distribution dispersion parameter.

#### Details

Estimate matrix of dispersion parameter alpha (size) used in regionSegCost for negative binomial distributed x.

#### Value

Matrix with maxk rows and nrow(x) columns, or a length(segs)+1 square matrix for the second stage model.

#### References

Piegorsch, W. W. (1990). Maximum likelihood estimation for the negative binomial dispersion parameter. Biometrics, 863-867.

Robinson MD and Smyth GK (2008). Small-sample estimation of negative binomial dispersion, with applications to SAGE data. Biostatistics, 9, 321-332

#### See Also

## regionSegCost

```
x<-matrix(rnbinom(120, size=0.05, mu=20), ncol=3)
Aa<-regionSegAlphaNB(x, maxk=20)
dim(Aa) # [1] 20 40
Ab<-regionSegAlphaNB(x, segs=as.integer(c(3, 6, 12, 30)))
dim(Ab) # [1] 5 5</pre>
```

regionSegCost

#### Description

To calculate regional cost matrix for the initial stage and second merging stage of the segmentation model.

## Usage

regionSegCost(x, maxk = NULL, segs = NULL, family = NULL, alpha = NULL, useSum = TRUE, useMC = FALSE,

#### Arguments

The input data matrix or vector
Maximum number of index to search forward
Starting indices (excluding 1) for the candidate segments, for the second stage model, maxk will be overridden with length(segs)+1.
which exponential family the data belongs to, possible values are 'norm', 'pois' and 'nbinom'
alpha matrix for negative binomial cost calculation, estimated from regionSegAlphaNB
TRUE if using grand sum across sample / x columns, like in the tilingArray solution
TRUE if mclapply should be used to speed up
TRUE if assuming common variance across samples (x columns)

#### Details

Preparing the cost matrix for the follow-up segmentation. Using residual sum of squares for 'norm' data, and negative log-likelihood for 'pois' and 'nbinom' data. Extension of the costMatrix function in tilingArray.

#### Value

Matrix with maxk rows and nrow(x) columns, or a length(segs)+1 square matrix for the second stage model.

#### References

Piegorsch, W. W. (1990). Maximum likelihood estimation for the negative binomial dispersion parameter. Biometrics, 863-867.

Picard,F. et al. (2005) A statistical approach for array CGH data analysis. BMC Bioinformatics, 6, 27.

Huber, W. et al. (2006) Transcript mapping with high density oligonucleotide tiling arrays. Bioinformatics, 22, 1963-1970.

Robinson MD and Smyth GK (2008). Small-sample estimation of negative binomial dispersion, with applications to SAGE data. Biostatistics, 9, 321-332

#### See Also

regionSegAlphaNB

#### Examples

```
x<-matrix(rnorm(120), ncol=3)
Ca<-regionSegCost(x, maxk=20, family='norm')
dim(Ca) # [1] 20 40
Cb<-regionSegCost(x, segs=as.integer(c(3, 6, 12, 30)), family='norm')
dim(Cb) # [1] 5 5</pre>
```

simSegData	Simulate exemplary segmentation data.
------------	---------------------------------------

## Description

Simulate exemplary segmentation data.

## Usage

simSegData(nseg=10, J=3, soj, emis, seed=1234, toPlot=FALSE)

## Arguments

nseg	size of initial segments pool
J	states number
soj	a list object containing sojourn settings
emis	a list object containing emission settings
seed	seed for simulation
toPlot	whether to output a pdf image of the simulated series

## Value

a list object containing the simulated data and the segment info

E	a numeric vector of the simulated data serie
L	a vector of the length for each continuous segment
S	a vector of state assignment for each segment
pdf	the name of the output pdf file if any

## Examples

```
soj<-list(type='pois', lambda=c(200, 100, 10))
emis<-list(type='pois', lambda=1:3)
simSegData(soj=soj, emis=emis)</pre>
```

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sojournAnno	Estimate sojourn distribution parameters from posterior information
	like annotation data

## Description

Using prior information from previous studies or annotation data to determine sojourn distribution parameters

## Usage

sojournAnno(xAnno, soj.type = "gamma", pbdist = NULL)

#### Arguments

xAnno	a GRanges / GRangesList / TxDb object, with its first meta column to represent the possible type of the range; Or a list object with named initial value vectors matching required parameters for a specific soj.type
soj.type	type of the sojourn distribution, following types are supported: 'gamma', 'pois', 'nbinom'
pbdist	average distance between neighbouring features, in this case in the link{biomvRhsmm} call one should only use the rank rather than the position.

#### Details

Be default, the hidden-semi Markov model implemented in this package uses a uniform prior for the initial sojourn distribution. However user can provide custom data from related studies to learn the prior of the sojourn distribution. The number of possible state will also be estimated from the unique level of feature type in the first meta column of xAnno if it is not a TxDb object.

## Value

a list object containing the sojourn distribution parameter

type	type of the sojourn distribution
fttypes	unique levels of the types stored in the first meta column of xAnno, alphabeti- cally sorted
J	number of possible states
	distribution parameters, 'lambda' and 'shift' for 'pois'; 'size', 'mu' and 'shift' for 'nbinom'; 'scale' and 'shape' for 'gamma'

#### Author(s)

Yang Du

```
data(encodeTP53)
encodeTP53$gmgr # a GRanges object
soj<-sojournAnno(encodeTP53$gmgr, soj.type='gamma')</pre>
```

splitFarNeighbour

#### Description

Split segments if long gaps exist between feature positions, due to low coverage or resolution.

## Usage

```
splitFarNeighbour(intStart = NULL, intEnd = NULL, xPos = NULL, xRange = NULL, maxgap = Inf, minrun = ?
```

## Arguments

intStart	indices of start for each segment
intEnd	indices of end for each segment
xPos	position vector, the distance of neighbouring features will be counted as point to point
xRange	IRanges / GRanges object for the positions, the the distance of neighbouring features will be counted as end to start.
maxgap	maximum distance between neighbouring features
minrun	when splitting, the minimum length of the features spanning, which half will be ignored if shorter.

## Value

a list object containing the start and end indices for new segments

IS	the start indices for new segments
IE	the end indices for new segments

## Author(s)

Yang Du

```
set.seed(123)
pos<-cumsum(rnbinom(20, size=10, prob=0.01))
splitFarNeighbour(intStart=c(1, 10), intEnd=c(6, 18), xPos=pos, maxgap=1000)</pre>
```

variosm

## Description

Extracted from package BiSeq, which is a small subset of a published study using targeted bisulfite sequencing data to detect differentially methylated regions (DMRs).

### Format

Containing one GRanges object

## References

http://dx.doi.org/10.1182 Schoofs et al. DNA methylation changes are a late event in acute upromyelocytic leukemia and coincide with loss of transcription factor binding. Blood, Nov 2012.

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