

Package ‘selectKSigs’

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Type Package

Title Selecting the number of mutational signatures using a perplexity-based measure and cross-validation

Depends R(>= 3.6)

Imports HiLDA, magrittr, gtools, methods, Rcpp

Suggests knitr, rmarkdown, testthat, BiocStyle, ggplot2, dplyr, tidyr

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Description A package to suggest the number of mutational signatures in a collection of somatic mutations using calculating the cross-validated perplexity score.

URL <https://github.com/USCbiostats/selectKSigs>

BugReports <https://github.com/USCbiostats/HiLDA/selectKSigs>

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biocViews Software, SomaticMutation, Sequencing, StatisticalMethod, Clustering

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calcPMSLikelihood	<i>A function for calculating the log-likelihood from the data and parameters</i>
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Description

A function for calculating the log-likelihood from the data and parameters

Usage

```
calcPMSLikelihood(p, y)
```

Arguments

p	this variable includes the parameters for mutation signatures and membership parameters
y	this variable includes the information on the mutation features, the number of mutation signatures specified and so on

Value

a value

Calculate_Likelihood_test

Output the maximum potential scale reduction statistic of all parameters estimated

Description

Output the maximum potential scale reduction statistic of all parameters estimated

Usage

```
Calculate_Likelihood_test(train, test, paramG)
```

Arguments

train	a MutationFeatureData S4 class output of training data.
test	a MutationFeatureData S4 class output of test data.
paramG	an estimatedParameters S4 class with estimated parameters

Value

the likelihood of the test data

convertFromTurbo_F *Restore the converted parameter F for turboEM*

Description

Restore the converted parameter F for turboEM

Usage

```
convertFromTurbo_F(turboF, fdim, signatureNum, isBackground)
```

Arguments

turboF	F (converted for turboEM)
fdim	a vector specifying the number of possible values for each mutation signature
signatureNum	the number of mutation signatures
isBackground	the logical value showing whether a background mutation features is included or not

Value

a vector

convertFromTurbo_Q *Restore the converted parameter Q for turboEM*

Description

Restore the converted parameter Q for turboEM

Usage

```
convertFromTurbo_Q(turboQ, signatureNum, sampleNum)
```

Arguments

turboQ	Q (converted for turboEM)
signatureNum	the number of mutation signatures
sampleNum	the number of cancer genomes

Value

a vector

convertToTurbo_F *Convert the parameter F so that turboEM can treat*

Description

Convert the parameter F so that turboEM can treat

Usage

```
convertToTurbo_F(vF, fdim, signatureNum, isBackground)
```

Arguments

vF	F (converted to a vector)
fdim	a vector specifying the number of possible values for each mutation signature
signatureNum	the number of mutation signatures
isBackground	the logical value showing whether a background mutation features is included or not

Value

a vector

convertToTurbo_Q	<i>Convert the parameter Q so that turboEM can treat</i>
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Description

Convert the parameter Q so that turboEM can treat

Usage

```
convertToTurbo_Q(vQ, signatureNum, sampleNum)
```

Arguments

vQ	Q (converted to a vector)
signatureNum	the number of mutation signatures
sampleNum	the number of cancer genomes

Value

a vector

cv_PMSignature	<i>Output the maximum potential scale reduction statistic of all parameters estimated</i>
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Description

Output the maximum potential scale reduction statistic of all parameters estimated

Usage

```
cv_PMSignature(inputG, Kfold = 3, nRep = 3, Klimit = 8)
```

Arguments

inputG	a MutationFeatureData S4 class.
Kfold	an integer number of the number of cross-validation folds.
nRep	an integer number of replications.
Klimit	an integer of the maximum value of number of signatures.

Value

a matrix of measures

Examples

```
load(system.file("extdata/sample.rdata", package = "selectKSigs"))
results <- cv_PMSignature(G, Kfold = 3)
```

getBG *Get the status of using the background signature*

Description

Get the status of using the background signature

Usage

```
getBG(object)
```

Arguments

object the EstimatedParameters class (the result of pmgetSignature)

Value

the status of using the background signature

getCounts *Get the count data in a matrix*

Description

Get the count data in a matrix

Usage

```
getCounts(object)
```

Arguments

object the MutationFeatureData class

Value

the count data in a matrix

getExposures *Get a matrix of mutational exposures of signatures*

Description

Get a matrix of mutational exposures of signatures

Usage

getExposures(object)

Arguments

object the EstimatedParameters class (the result of pmgetSignature)

Value

a matrix of mutational exposures of signatures

getFeatures *Get a vector of possible features*

Description

Get a vector of possible features

Usage

getFeatures(object)

Arguments

object the EstimatedParameters class (the result of pmgetSignature)

Value

a vector of possible features

getFeatureVec *Get a matrix of feature vector list*

Description

Get a matrix of feature vector list

Usage

getFeatureVec(object)

Arguments

object the MutationFeatureData class

Value

a matrix of feature vector list

getK *Get the number of signatures*

Description

Get the number of signatures

Usage

getK(object)

Arguments

object the EstimatedParameters class (the result of pmgetSignature)

Value

the number of signatures in pmgetSignature in HiLDA

getLL	<i>Get the values of loglikelihood</i>
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Description

Get the values of loglikelihood

Usage

```
getLL(object)
```

Arguments

object the EstimatedParameters class (the result of pmgetSignature)

Value

likelihood values estimated by pmgetSignature in HiLDA

getLogLikelihoodC	<i>Calculate the value of the log-likelihood for given parameters</i>
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Description

Calculate the value of the log-likelihood for given parameters

Usage

```
getLogLikelihoodC(  
  vPatternList,  
  vSparseCount,  
  vF,  
  vQ,  
  fdim,  
  signatureNum,  
  sampleNum,  
  patternNum,  
  samplePatternNum,  
  isBackground,  
  vF0  
)
```

Arguments

vPatternList	The list of possible mutation features (converted to a vector)
vSparseCount	The table showing (mutation feature, sample, the number of mutation) (converted to a vector)
vF	F (converted to a vector)
vQ	Q (converted to a vector)
fdim	a vector specifying the number of possible values for each mutation signature
signatureNum	the number of mutation signatures
sampleNum	the number of cancer genomes
patternNum	the number of possible combinations of all the mutation features
samplePatternNum	the number of possible combination of samples and mutation patterns
isBackground	the logical value showing whether a background mutation features is included or not
vF0	a background mutation features

Value

a value

getSamplelist	<i>Get the sample list</i>
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Description

Get the sample list

Usage

```
getSamplelist(object)
```

Arguments

object	the EstimatedParameters class (the result of pmgetSignature)
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Value

the sample list of named elements.

`getSamplelistG` *Get the sample list*

Description

Get the sample list

Usage

`getSamplelistG(object)`

Arguments

`object` the `MutationFeatureData` class

Value

the sample list of named elements.

`getSignatures` *Get an array of signature feature distributions*

Description

Get an array of signature feature distributions

Usage

`getSignatures(object)`

Arguments

`object` the `EstimatedParameters` class (the result of `pmgetSignature`)

Value

an array of signature feature distributions

getTranscription	<i>Get the status of specifying the transcription bias</i>
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Description

Get the status of specifying the transcription bias

Usage

```
getTranscription(object)
```

Arguments

object the MutationFeatureData class

Value

the status of specifying the transcription bias

select_kth_fold	<i>Output the training data or test data</i>
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Description

Output the training data or test data

Usage

```
select_kth_fold(inputG, k, f_s, folds, include)
```

Arguments

inputG	a MutationFeatureData S4 class output by the pmsignature.
k	an integer number of the number of cross-validation folds.
f_s	a primary key of combining the feature pattern and sample ID.
folds	the assignment to each fold.
include	a boolean indicator of whether to include kth fold or not.

Value

a MutationFeatureData S4 class of either include or exclude kth fold.

splitG	<i>Output the maximum potential scale reduction statistic of all parameters estimated</i>
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Description

Output the maximum potential scale reduction statistic of all parameters estimated

Usage

```
splitG(inputG, Kfold = 3)
```

Arguments

inputG	a MutationFeatureData S4 class output by the pmsignature.
Kfold	an integer number of the number of cross-validation folds.

Value

a matrix made of perplexity from the results of cross-validation.

Examples

```
load(system.file("extdata/sample.rdata", package = "selectKSigs"))  
G_split <- splitG(G, Kfold = 3)
```

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