# Package 'SUITOR'

July 6, 2020

Title Selecting the number of mutational signatures through

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Maintainer Bill Wheeler <wheelerb@imsweb.com> Depends R (&gt;= 3.5.0), doParallel, foreach, parallel</wheelerb@imsweb.com>															
•	Date 2020-07-06  Author Donghyuk Lee <donghyuk.lee@nih.gov> and Bin Zhu <bin.zhu@nih.gov>  Description Selecting the number of mutational signatures through cross-validation</bin.zhu@nih.gov></donghyuk.lee@nih.gov>														
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Version	0.0.4														

# Description

Selecting the number of mutational signatures through cross-validation

# Author(s)

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data

Data for examples

# **Description**

Example input data and results

# **Details**

Contains an example input data object of size 96 by 30 and an example matrix of results. It is generated by rpois with mean WH where W (96 by 1) is profile of signature 8 obtained from https://cancer.sanger.ac.uk/cosmic/signatures/SBS/SBS8.tt and H (1 by 30) is rounded integer generated by uniform distribution between 20,000 and 40,000.

#### See Also

suitor

# **Examples**

```
data(data, package="SUITOR")

# Display a subset of data objects
data[1:5, 1:5]
names(results)
```

getSummary

Compute summary results

# **Description**

Compute summary results and the optimal rank from the matrix containing all results

# Usage

```
getSummary(obj, NC, NR=96)
```

# Arguments

obj	Matrix containing all results in the return list from suitor.
NC	The number of columns in data when suitor was called.

NR The number of rows in data when suitor was called. The default is 96.

# **Details**

The input matrix obj must have column 1 as the rank, column 2 as the value of k in 1:k.fold, column 4 as the training errors, and column 5 as the testing errors.

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#### Value

A list containing the objects:

- rank: The optimal rank
- all.results: Matrix containing training and testing errors for all values of seeds, ranks, folds. NA values appear for runs in which the EM algorithm did not converge.
- summary: Data frame of summarized results for each possible rank created from all.results. The MSErr column is defined as sqrt({fold1 + ... +foldK}/{nrow(data)\*ncol(data)})

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#### See Also

```
plotErrors
```

# **Examples**

```
data(data, package="SUITOR")
ret <- getSummary(results$all.results, ncol(data))
ret$summary
ret$rank</pre>
```

plotErrors

Plot train and test errors

# Description

Plot train and test errors

#### Usage

```
plotErrors(x)
```

#### **Arguments**

Х

Data frame of summary results in the return list from suitor or from getSummary.

#### **Details**

The optimal rank is the minimum at which the test error is attained, and appears as a red dot on the graph.

### Value

NULL

#### Author(s)

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# **Examples**

```
data(data, package="SUITOR")
s <- getSummary(results$all.results, ncol(data))
plotErrors(s$summary)</pre>
```

# Description

Selecting the number of mutational signatures through cross-validation

# Usage

```
suitor(data, op=NULL)
```

# **Arguments**

data	Data frame or matrix containing mutational signatures. This object must contain
	non-negative values
ор	List of options (see details). The default is NULL.

# **Details**

The algorithm finds the optimal rank by applying k-fold cross validation.

# Options list op:

Name	Description	<b>Default Value</b>
em.eps	EM algorithm stopping tolerance	1e-5
get.summary	0 or 1 to create summary results	1
k.fold	Number of folds	10
max.iter	Maximum number of iterations in EM algorithm	2000
max.rank	Maximum rank	10
min.rank	Minimum rank	1
min.value	Minimum value of matrix before factorizing	1e-4
n.cores	Number of cores to use (for non-Windows only)	1
n.seeds	Number of seeds (starting points)	30
plot	0 or 1 to produce an error plot	1
print	0-3 to print info (0=no printing)	1
seeds	Vector of seeds (takes precendence over n.seeds)	NULL
kfold.vec	Vector of values in 1:k.fold when running on a cluster	NULL

# Utilizing a cluster

When running on a cluster, the option get.summary should be set to 0. For fastest running jobs, set the options min.rank = max.rank, kfold.vec to a single integer in 1:k.fold, and seeds to a single value.

# Value

A list containing the objects:

suitor 5

- rank: The optimal rank
- all.results: Matrix containing training and testing errors for all values of seeds, ranks, folds.

• summary: Data frame of summarized results for each possible rank created from all.results. The MSErr column is defined as  $sqrt(fold1 + ... + foldK)/\{nrow(data)*ncol(data)\})$ 

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#### See Also

```
getSummary, plotErrors
```

# **Examples**

```
data(data, package="SUITOR")
# Using the default options will take several minutes to run
#ret <- suitor(data)</pre>
```

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