

Package ‘SISPA’

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Type Package

Title SISPA: Method for Sample Integrated Set Profile Analysis

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Author Bhakti Dwivedi and Jeanne Kowalski

Maintainer Bhakti Dwivedi <bhakti.dwivedi@emory.edu>

Description Sample Integrated Set Profile Analysis (SISPA) is a method designed to define sample groups with similar gene set enrichment profiles.

Depends R (>= 3.5), genefilter, GSVA, changepoint

Imports data.table, plyr, ggplot2

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LazyData true

Collate 'freqplot.R' 'waterfallplot.R' 'callGSVA.R' 'sortData.R'
'cptSamples.R' 'callZSCORE.R' 'filterVars.R' 'SISPA.R' 'data.R'

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VignetteBuilder knitr

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callGSVA	<i>GSVA enrichment analysis</i>
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Description

Estimates GSVA enrichment zscores.

Usage

callGSVA(x,y)

Arguments

- x A data frame or matrix of gene or probe expression values where rows correspond to genes and columns correspond to samples
- y A list of genes as data frame or vector

Details

This function uses "zscore" gene-set enrichment method in the estimation of gene-set enrichment scores per sample.

Value

A gene-set by sample matrix of GSVA enrichment zscores.

See Also

GSVA

Examples

```

g <- 10 ## number of genes
s <- 30 ## number of samples
## sample data matrix with values ranging from 1 to 10
rnames <- paste("g", 1:g, sep="")
cnames <- paste("s", 1:s, sep="")
expr <- matrix(sample.int(10, size = g*s, replace = TRUE), nrow=g, ncol=s, dimnames=list(rnames, cnames))
## genes of interest
genes <- paste("g", 1:g, sep="")
## Estimates GSVA enrichment zscores.
callGSVA(expr,genes)

```

callZSCORE	<i>Row ZSCORES</i>
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Description

Estimates the zscores for each row in the data matrix

Usage

```
callZSCORE(x)
```

Arguments

x	A data frame or matrix of gene or probe expression values where rows correspond to genes and columns correspond to samples
---	--

Details

This function compute row zscores per sample when number of genes is less than 3

Value

A gene-set by sample matrix of zscores.

Examples

```

g <- 2 ## number of genes
s <- 60 ## number of samples
## sample data matrix with values ranging from 1 to 10
rnames <- paste("g", 1:g, sep="")
cnames <- paste("s", 1:s, sep="")
expr <- matrix(sample.int(10, size = g*s, replace = TRUE), nrow=g, ncol=s, dimnames=list(rnames, cnames))
## Estimates zscores
callZSCORE(expr)

```

cptSamples *Sample profile identifier analysis*

Description

Generate sample profile identifiers from sample zscores using change point model.

Usage

```
cptSamples(x, cpt_data, cpt_method, cpt_max)
```

Arguments

x	A matrix or data frame of sample GSVA enrichment zscores within which you wish to find a changepoint.
cpt_data	Identify changepoints for data using variance (cpt.var), mean (cpt.mean) or both (cpt.meanvar). Default is cpt.var.
cpt_method	Choice of single or multiple changepoint model. Default is "BinSeg".
cpt_max	The maximum number of changepoints to search for using "BinSeg" method. Default is 60.

Details

This function assigns samples identified in the first changepoint with the active profile ("1") while the remaining samples are grouped under inactive profile ("0").

Value

The input data frame with added sample identifiers and estimated changepoints. A plot showing the changepoint locations estimated on the data

See Also

changepoint

Examples

```
g <- 10 ## number of genes
s <- 60 ## number of samples
## sample data matrix with values ranging from 1 to 10
rnames <- paste("g", 1:g, sep="")
cnames <- paste("s", 1:s, sep="")
expr <- matrix(sample.int(10, size = g*s, replace = TRUE), nrow=g, ncol=s, dimnames=list(rnames, cnames))
## genes of interest
genes <- paste("g", 1:g, sep="")
## Estimates GSVA enrichment zscores.
gsva_results <- callGSVA(expr,genes)
cptSamples(gsva_results,cpt_data="var",cpt_method="BinSeg",cpt_max=60)
```

ExpressionSet	<i>Gene Expression (ExpressionSet) data</i>
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Description

An example of RNAseq derived gene expression data The variables are as follows

Usage

```
ExpressionSet
```

Format

A data matrix with 8 probes (rows) in 125 samples (columns)

Details

- gene
- sample 1 log2 transformed normalized expression count values
- sample nth log2 transformed normalized expression count values

@source <https://research.themmr.org/>

filterVars	<i>A filter function for the data</i>
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Description

Filter rows with zero values

Usage

```
filterVars(x,y)
```

Arguments

- | | |
|---|--|
| x | : A data frame or matrix where rows represent gene and columns represent samples |
| y | : A vector of a sample column values to apply the filtering on. |

Details

This function filter out rows with zero data value for a given sample. Both input arguments (x and y) must be of the same length

Value

The returned value is a list containing an entry for each row filtered out by zero data value

Examples

```
x = matrix(runif(3*10, 0, 1), ncol=3)
y <- x[,1]
filterVars(x,y)
```

freqplot

A plotting function for SISPA sample identifiers

Description

Given a sample changepoint data frame, will plot number of samples with and without profile activity

Usage

```
freqplot(x)
```

Arguments

x A data frame containing samples as rows followed by zscores and estimated changepoints to be plotted.

Details

This function expects the output from cptSamples function of SISPA package, and shows the number of samples with (orange filled bars) and without profile activity (grey filled bars).

Value

Bar plot pdf illustrating distribution of samples

Examples

```
samples <- c("s1", "s2", "s3", "s4", "s5", "s6", "s7", "s8", "s9", "s10")
zscores <- c(3.83, 2.70, 2.67, 2.31, 1.70, 1.25, -0.42, -1.01, -2.43, -3.37)
changepoints <- c(1, 1, 1, 2, 2, 3, 3, NA, NA, NA)
sample_groups <- c(1, 1, 1, 0, 0, 0, 0, 0, 0, 0)
my.data = data.frame(samples, zscores, changepoints, sample_groups)
freqplot(my.data)
```

SISPA	<i>SISPA</i>
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Description

SISPA: Method for Sample Integrated Gene Set Analysis

Usage

```
SISPA(feature=1,f1.df,f1.profile,f2.df,f2.profile,cpt_data="var",cpt_method="BinSeg",cpt_max=60)
```

Arguments

feature	Number of input feature or data types
f1.df	A data matrix of first feature (e.g., gene or probe expression values) where rows correspond to genes and columns correspond to samples
f1.profile	A flag to specify gene profile. If gene.profile="up" then samples with increased zscores are identified. If gene.profile="down" then samples with decreased zscores are identified. Default is "up".
f2.df	A data matrix of second feature (e.g., gene variant change) where rows correspond to genes and columns correspond to samples
f2.profile	A flag to specify gene profile. If gene.profile="up" then samples with increased zscores are identified. If gene.profile="down" then samples with decreased zscores are identified. Default is "up".
cpt_data	Identify changepoints for data using variance (cpt.var), mean (cpt.mean), or both (cpt.meanvar). Default is cpt.var.
cpt_method	Choice of single or multiple changepoint model. Default is "BinSeg". See changepoint R package for details
cpt_max	The maximum number of changepoints to search for using "BinSeg" method. Default is 60.

Details

Sample Integrated Gene Set Analysis (SISPA) is a method designed to define sample groups with similar gene set enrichment profiles. The user specifies a gene list of interest and sample by gene molecular data (expression, methylation, variant, or copy change data) to obtain gene set enrichment scores by each sample. The score statistics is rank ordered by the desired profile (e.g., upregulated or downregulated) for samples. A change point model is then applied to the sample scores to identify groups of samples that show similar gene set profile patterns. Samples are ranked by desired profile activity score and grouped by samples with and without profile activity. Figure 1 shows the schematic representation of the SISPA method overview.

Value

The input molecular data frame with added sample identifiers and estimated changepoints. A plot showing the changepoint locations estimated on the data. Bar plots pdf illustrating distinct distribution of samples with and without profile activity

Examples

```
g <- 10 ## number of genes
s <- 60 ## number of samples
## sample data matrix with values ranging from 1 to 10
rnames <- paste("g", 1:g, sep="")
cnames <- paste("s", 1:s, sep="")
expr <- matrix(sample.int(10, size = g*s, replace = TRUE), nrow=g, ncol=s, dimnames=list(rnames, cnames))
SISPA(feature=1,f1.df=expr,f1.profile="up")
```

sortData

Sorts the data by a column

Description

Sorts the data frame by a column index in the given order

Usage

```
sortData(x,i,b)
```

Arguments

x	A data frame
i	A numeric column index of the data frame to sort it by
b	User specified sorting order, ascending (FALSE) or descending (TRUE)

Details

defaults are used: i = 1, b = FALSE, if not specified

Value

sorted data by the input column index

Author(s)

Bhakti Dwivedi & Jeanne Kowalski

Examples

```
samples <- c("s1","s2","s3","s4","s5","s6","s7","s8","s9","s10")
zscores <- c(3.83,2.70,2.67,2.31,1.70,1.25,-0.42,-1.01,-2.43,-3.37)
my.data = data.frame(samples,zscores)
sortData(my.data,2,TRUE)
```

VariantSet	<i>Variant (VariantSet) data</i>
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Description

An example of RNAseq derived gene variant change data. The variables are as follows

Usage

```
VariantSet
```

Format

A data matrix with 380 variants (rows) in 125 samples (column)

Details

- gene
- sample 1 transformed variant proportion data
- sample nth transformed variant proportion data

@source <https://research.themmr.org/>

waterfallplot	<i>A plotting function for SISPA sample identifiers</i>
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Description

Given a sample changepoint data frame, will plot all samples zscores from that data.

Usage

```
waterfallplot(x)
```

Arguments

x A data frame containing samples as rows followed by zscores and estimated sample_groups to be plotted.

Details

This function expects the output from cptSamples function of SISPA package, and highlights the sample profile of interest in the changepoint 1 with orange filled bars.

Value

Bar plot pdf illustrating distinct SISPA sample profiles.

Examples

```
samples <- c("s1","s2","s3","s4","s5","s6","s7","s8","s9","s10")
zscores <- c(3.83,2.70,2.67,2.31,1.70,1.25,-0.42,-1.01,-2.43,-3.37)
changepoints <- c(1,1,1,2,2,3,3,NA,NA,NA)
sample_groups <- c(1,1,1,0,0,0,0,0,0,0)
my.data = data.frame(samples,zscores,changepoints,sample_groups)
waterfallplot(my.data)
```

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