

Package ‘CEDA’

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Type Package

Title CRISPR Screen and Gene Expression Differential Analysis

Version 1.1.0

Description Provides analytical methods for analyzing CRISPR screen data at different levels of gene expression. Multi-component normal mixture models and EM algorithms are used for modeling.

Depends R(>= 3.5.0), limma

Imports stats, mixtools, ggplot2, dplyr, ggsci, ggribes, ggprism

Suggests knitr, rmarkdown

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alphaBeta *Calculating a significance score of a gene based on the corresponding sgRNAs' p-values of the gene.*

Description

Code was adapted from R package gscreend.

Usage

alphaBeta(pvec)

Arguments

pvec A numeric vector of p-values.

Value

A min value of the kth smallest value based on the beta distribution $B(k, n-k+1)$, where the n is the number of probabilities in the vector. This min value is the significance score of the gene.

calculateGeneLFC *Calculating gene-level log fold ratios*

Description

Log fold ratios of all sgRNAs of a gene are averaged to obtain the gene level log fold ratio.

Usage

calculateGeneLFC(lfcs, genes)

Arguments

lfcs A numeric vector containing log fold change of sgRNAs.
 genes A character string containing gene names corresponding to sgRNAs.

Value

A numeric vector containing log fold ratio of genes.

calculateGenePval	<i>Calculating gene level p-values using modified robust rank aggregation (alpha-RRA method) on sgRNAs' p-values</i>
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Description

Code was adapted from R package gscreend. The alpha-RRA method is adapted from MAGeCK.

Usage

```
calculateGenePval(pvec, genes, alpha, nperm = 20)
```

Arguments

pvec	A numeric vector containing p-values of sgRNAs.
genes	A character string containing gene names corresponding to sgRNAs.
alpha	A numeric number denoting the alpha cutoff (i.e. 0.05).
nperm	Number of permutations, default is 20

Value

A list with four elements: 1) a list of genes with their p-values; 2) a numeric matrix of rho null, each column corresponding to a different number of sgRNAs per gene; 3) a numeric vector of rho; 4) a numeric vector of number of sgRNAs per gene.

densityPlot	<i>2D density contour plot of gene log2 fold ratios against gene expression levels</i>
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Description

This function generates a scatter plot with 2D density contour of log2 fold ratios of sgRNAs against the corresponding gene expression levels.

Usage

```
densityPlot(data, ...)
```

Arguments

data	A data frame from the output of preparePlotData function
...	Other graphical parameters

Value

No return value

EMFit	<i>Fitting multi-component normal mixture models by R package mixtools</i>
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Description

The function normalmixEM in R package mixtools is employed for fitting multi-component normal mixture models.

Usage

```
EMFit(x, k0, mean_constr, sd_constr, npara, d0)
```

Arguments

x	A numeric vector
k0	Number of components in the normal mixture model
mean_constr	A constrain on means of components
sd_constr	A constrain on standard deviations of components
npara	Number of parameters
d0	Number of times for fitting mixture model using different starting values

Value

Normal mixture model fit and BIC value of the log-likelihood

makeRhoNull	<i>Generating the null distribution of the significance score of a gene.</i>
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Description

Code was adapted from R package gscreend.

Usage

```
makeRhoNull(n, p, nperm)
```

Arguments

n	An integer representing sgRNA number of a gene.
p	A numeric vector which contains the percentiles of the p-values that meet the cut-off (alpha).
nperm	Number of permutation runs.

Value

A numeric vector which contains all the significance scores (rho) of genes generated by a permutation test where the sgRNAs are randomly assigned to genes.

mda231	<i>CRISPR screen data of cell line MDA-MB-231.</i>
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Description

A dataset containing the expression data of sgRNAs in a CRISPR screen experiment of cell line MDA-MB-231.

Usage

```
mda231
```

Format

A data frame with a list of two elements:

sgRNA Raw Read counts of sgRNAs

negene A list of non-essential genes

medianNormalization	<i>Median normalization of sgRNA counts</i>
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Description

This function adjusts sgRNA counts by the median ratio method. The normalized sgRNA read counts are calculated as the raw read counts divided by a size factor. The size factor is calculated as the median of all size factors calculated from negative control sgRNAs (eg., sgRNAs corresponding to non-targeting or non-essential genes).

Usage

```
medianNormalization(data, control)
```

Arguments

data	A numeric matrix containing raw read counts of sgRNAs with rows corresponding to sgRNAs and columns corresponding to samples.
control	A numeric matrix containing raw read counts of negative control sgRNAs with rows corresponding to sgRNAs and columns corresponding to samples. Sample ordering is the same as in data.

Value

A list with two elements: 1) size factors of all samples; 2) normalized counts of sgRNAs.

Examples

```
count <- matrix(rnbinom(5000 * 6, mu=500, size=3), ncol = 6)
colnames(count) = paste0("sample", 1:6)
rownames(count) = paste0("sgRNA", 1:5000)
control <- count[1:100,]
normalizedcount <- medianNormalization(count, control)
```

normalMM

Performing empirical Bayes modeling on limma results

Description

This function perform an empirical Bayes modeling on log fold ratios and return the posterior log fold ratios.

Usage

```
normalMM(data, theta0, n.b = 5, d = 10)
```

Arguments

data	A numeric matrix containing limma results and log ₂ gene expression levels that has a column named 'lfc' and a column named 'exp.level.log2'
theta0	Standard deviation of log ₂ fold changes under permutations
n.b	Number of bins, default is 5 bins
d	Number of times for fitting mixture model using different starting values, default is 10

Value

A numeric matrix containing limma results, RNA expression levels, posterior log₂ fold ratio, log p-values, and estimates of mixture model

permutelimma	<i>Modeling CRISPR data with a permutation test between conditions by R package limma</i>
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Description

The lmFit function in R package limma is employed for group comparisons under permutations.

Usage

```
permutelimma(data, design, contrast.matrix, nperm)
```

Arguments

data	A numeric matrix containing log2 expression level of sgRNAs with rows corresponding to sgRNAs and columns to samples.
design	A design matrix with rows corresponding to samples and columns to coefficients to be estimated.
contrast.matrix	A matrix with columns corresponding to contrasts.
nperm	Number of permutations

Value

A numeric matrix containing log2 fold changes with permutations

Examples

```
y <- matrix(rnorm(1000*6),1000,6)
condition <- gl(2,3,labels=c("Control","Baseline"))
design <- model.matrix(~ 0 + condition)
contrast.matrix <- makeContrasts("conditionControl-conditionBaseline",levels=design)
fit <- permutelimma(y,design,contrast.matrix,20)
```

preparePlotData	<i>Prepare data for density plot and ridge plot</i>
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Description

Input a data frame with each gene one row, and geneID, geneLFC, geneFDR as columns. This function will stratify genes into five groups based on their FDR levels: ≤ 0.001 , (0.001,0.01], (0.01,0.05], (0.05,0.5], (0.5,1]

Usage

```
preparePlotData(data, gene.fdr)
```

Arguments

data	A data frame containing each gene in one row, and at least three columns with geneID, geneLFC, and geneFDR.
gene.fdr	A numeric variable (column) in the data frame, corresponding to the gene level FDR

Value

A data frame based on the original data frame, with an additional column "group" indicating which FDR group this gene belongs to.

ridgePlot	<i>Density ridgeline plot of gene expression levels for different FDR groups.</i>
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Description

This function generates a density ridgeline plot of gene expression levels for different FDR groups.

Usage

```
ridgePlot(data, ...)
```

Arguments

data	A data frame from the output of preparePlotData function
...	Other graphical parameters

Value

No return value

`runLimma`*Modeling CRISPR screen data by R package limma*

Description

The `lmFit` function in R package `limma` is employed for group comparisons.

Usage

```
runLimma(data, design, contrast.matrix)
```

Arguments

`data` A numeric matrix containing log₂ expression levels of sgRNAs with rows corresponding to sgRNAs and columns corresponding to samples.

`design` A design matrix with rows corresponding to samples and columns corresponding to coefficients to be estimated.

`contrast.matrix` A matrix with columns corresponding to contrasts.

Value

A data frame with rows corresponding to sgRNAs and columns corresponding to `limma` results

Examples

```
y <- matrix(rnorm(1000*6),1000,6)
condition <- gl(2,3,labels=c("Treatment","Baseline"))
design <- model.matrix(~ 0 + condition)
contrast.matrix <- makeContrasts("conditionTreatment-conditionBaseline",levels=design)
limma.fit <- runLimma(y,design,contrast.matrix)
```

`scatterPlot`*Scatter plot of log₂ fold ratios against gene expression levels*

Description

This function generates a scatter plot of log₂ fold ratios of sgRNAs against the corresponding gene expression levels.

Usage

```
scatterPlot(data, fdr, ...)
```

Arguments

<code>data</code>	A numeric matrix from the output of <code>normalMM</code> function
<code>fdr</code>	A level of false discovery rate
<code>...</code>	Other graphical parameters

Value

No return value

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