Package 'deGPS'

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Type Package
Title Differential Expression Tests Based on Generalized Poisson Statistic
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Depends R (>= 3.0.0), foreach, doParallel
Suggests LPE, limma, edgeR
Description Use methods based on Generalized Poisson Distribution to do RNA-seq differential expression tests.
License GPL-2
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Description

This package is proposed to analyze RNA-seq data in two steps: normalization and differential expression test.

New normalization methods based on generalized Poisson distribution are contained in the main analysis functions, in which GP-Theta is suggested. Other popular normalization methods, such as TMM, LOWESS, Quantile, is also availbale in the package. More than one method can be specified in one run, in which case the resulting p values are a p value matrix, with each column representing one method.

Differential expression test is designed for RNA-seq data, especially those of small sample size, based on permutation strategy. Note that deGPS can only handle DE tests between two groups, but the novel GP-based normalization methods can be applied on any RNA-seq data. More Details about the GP-based normalization method and the advantages and limitations of our DE test can be found in our article.

The permutation step may become time-consuming in large sample size context or in mRNA read count data analysis. Parallel computation is introduced in the main functions to deal with the computational burden. Note that in situations where parallel computation is not necessary, to force parallelling may result in more run time.

The package also contains function to generate GP distributed data to be an example of RNA-seq data. It has to be pointed out that the simulated data in this package is FAR AWAY from real RNA-seq data, it is just simple GP distributed samples. For more appropriately simulated RNA-seq, compcodeR package is suggested, or, alternatively, you can generate H0 and H1 data from real data. A simple example is given in the following example session. More R codes referring to the real data based simulation or compcodeR based simulation can be found in the supplementary materials of our article.

Please do not hesitate to contact the author if you have any questions or find any flaws in the package.

Details

Package: deGPS Type: Package Version: 1.0

Date: 2014-02-24 License: GPL-2

To do DE test for miRNA, call GPSmle. To do DE test for mRNA, call deGPS_mRNA. If only the normalization step is required, call GPSmle and specify type as "normalization".

Some denotations used in this manual are as follow:

• nSample

sample size of the data. If there are more than one replicate for each sample, nSample represents the sample size multiplied by the number of replicates.

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nRNA

the number of genes in the data.

• n-miRNA

the number of miRNAs in the data.

• groupSize

the number of samples in either group.

• permutationTimes

the times of permutation used to obtain the empirical distribution of T-stat.

• round(.)

the round function in R. The value of round(x) is the closest integer of x.

• ceiling(.)

the ceiling function in R. The value of ceiling(x) is the integer part of x added by 1.

• ncol(.)

the ncol function in R. The value of ncol(data) is the number of columns of data.

• mod(.)

the mod function in R. The value of mod(x) is the remainder of x.

Author(s)

Chen Chu

Maintainer: Chen Chu <chuchenblueblues@gmail.com>

References

deGPS: a Powerful and Flexible Framework for Detecting Differential Expression in RNA-Sequencing Studies

```
## Not run:
###Analyze real RNA-seq data
data(bottomlyData)
group <- c(rep(1:2, each = 10), 2)
simuData <- as.matrix(bottomlyData[ , -1])</pre>
dimnames(simuData)[[1]] <- as.character(bottomlyData$gene)</pre>
### apply deGPS on bottomly data
bottomlyRes <- deGPS_mRNA(data = simuData, group = group,</pre>
ncore = 4, nSubcore = 4, method = "GP-Theta", maxIter = 150)
pvalue <- as.vector(bottomlyRes$pvalue)</pre>
adj.p <- p.adjust(pvalue, method = "BH")</pre>
topTags(pvalue)
##Generate Random samples from GP(theta, lambda)
examData <- newExampleData(nRNA = 100, groupSize = 6, lambda = 0.9,
theta = 3, ptol = 1e-15)
str(examData)
##Differential Expression Tests
examRes <- deGPS_mRNA(data = examData$data, group = examData$group,</pre>
```

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```
method = "GP-Theta", nSubcore = 2, ncore = 2, geneid = paste("G", 1:100, sep = ""))
str(examRes)
###Generate simulated RNA-seq data from compcodeR package
require(compcodeR)
samples.per.cond <- 5</pre>
random.outlier.high.prob <- 0.1</pre>
n.vars <- 10000
examData <- generateSyntheticData(dataset = "simuData",</pre>
n.vars = n.vars, samples.per.cond = samples.per.cond, n.diffexp = floor(n.vars * 0.1),
repl.id = 1, seqdepth = 1e+07, fraction.upregulated = 0.5,
between.group.diffdisp = FALSE, filter.threshold.total = 1,
filter.threshold.mediancpm = 0, fraction.non.overdispersed = 0,
random.outlier.high.prob = random.outlier.high.prob,
output.file = "simuData_repl1.rds")
group <- examData@sample.annotations$condition</pre>
examRes <- deGPS_mRNA(data = examData@count.matrix, group = group,
method = "GP-Theta", nSubcore = 2, ncore = 2, geneid = paste("G", 1:n.vars, sep = ""))
str(examRes)
## End(Not run)
```

bottomlyData

A real mouse RNA-seq data set

Description

A real mouse RNA-seq data set

Usage

```
data(bottomlyData)
```

Details

A mouse RNA-seq data set downloaded at Bottomly Data, study bottomly

References

Evaluating gene expression in C57BL/6J and DBA/2J mouse striatum using RNA-Seq and microarrays, Bottomly D and etc., PLoS One, 2011 Mar 24;6(3):e17820.

```
## Not run:
### load required packages
require(edgeR)
require(DESeq)
```

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```
require(DESeq2)
require(ggplot2)
require(gridExtra)
require(deGPS)
data(bottomlyData)
group <- c(rep(1:2, each = 10), 2)
simuData <- as.matrix(bottomlyData[ , -1])</pre>
### apply deGPS on bottomly data
bottomlyRes <- deGPS_mRNA(data = simuData, group = group,</pre>
ncore = 4, nSubcore = 4, method = "GP-Theta", maxIter = 300)
pvalue <- as.vector(bottomlyRes$pvalue)</pre>
### compare result with edgeR and DESeq, DESeq2
d0 <- DGEList(counts = simuData, group = group)</pre>
design <- model.matrix(~ group, data = d0$samples)</pre>
d <- try(calcNormFactors(d0, method = "TMM"))</pre>
d <- try(estimateGLMCommonDisp(d, design, verbose = TRUE))</pre>
d <- try(estimateGLMTrendedDisp(d, design))</pre>
efit <- try(glmQLFTest(d, design, coef = 2))</pre>
edgeRes <- efit$table$PValue
cds1 <- newCountDataSet(simuData, group)</pre>
cds2 <- estimateSizeFactors(cds1)</pre>
cds3 <- try(estimateDispersions(cds2))</pre>
if("try-error"
if("try-error"
fitType = "local"))
res <- nbinomTest(cds3, 1, 2)</pre>
deseqRes <- res$pval</pre>
deseqRes[is.na(deseqRes)] <- 1</pre>
dds <- DESeqDataSetFromMatrix(countData = simuData,</pre>
colData = data.frame(group = group),
design = ~ group)
dds <- DESeq(dds)
res <- results(dds)</pre>
deseq2Res <- res$pvalue</pre>
deseq2Res[is.na(deseq2Res)] <- 1</pre>
### plot the overlap of DEs
pAll <- cbind(pvalue, edgeRes, deseq2Res)</pre>
pAll <- apply(pAll, 2, p.adjust, method = "BH")
overLapTemp <- overLapTemp1 <- matrix(NA, ncol(pAll), ncol(pAll))</pre>
for(ii in 1:ncol(pAll)){
for(jj in 1:ncol(pAll)){
overLapTemp[ii, jj] \leftarrow sum(pAll[, ii] < 0.05 & pAll[, jj] < 0.05) / sum(pAll[, ii] < 0.05)
}
}
for(ii in 1:ncol(pAll)){
for(jj in 1:ncol(pAll)){
```

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```
overLapTemp1[ii, jj] <- sum(pAll[, ii] < 0.05 & pAll[, jj] < 0.05)
dimnames(overLapTemp) <- dimnames(overLapTemp1) <- list(c("deGPS", "edgeR", "DESeq", "DESeq2"), c("deGPS",</pre>
jpeg("bottomlyOverLap.jpg", width = 1600, height = 700)
a <- levelplot(overLapTemp, xlab = "", ylab = "", main = list("Overlap Proportion", cex = 2),</pre>
scale = list(cex = 1.3),
colorkey = list(labels = list(cex = 1.2)),
panel=function(...) {
arg <- list(...)
panel.levelplot(...)
panel.text(rep(1:nrow(overLapTemp), ncol(overLapTemp)),
rep(1:ncol(overLapTemp), each = nrow(overLapTemp)), round(as.vector(overLapTemp), 2), cex = 1.5)}
b <- levelplot(overLapTemp1, xlab = "", ylab = "", main = list("Overlap Number", cex = 2),
scale = list(cex = 1.3),
colorkey = list(labels = list(cex = 1.2)),
panel=function(...) {
arg <- list(...)
panel.levelplot(...)
panel.text(rep(1:nrow(overLapTemp1), ncol(overLapTemp1)),
rep(1:ncol(overLapTemp1), each = nrow(overLapTemp1)), as.vector(overLapTemp1), cex = 1.5)}
grid.arrange(a, b, ncol=2)
dev.off()
## End(Not run)
```

deGPS_mRNA

Normalization and Two-group Differential Expression Test for mRNA Read Count Data

Description

Normalization and Two-group Differential Expression Test for mRNA Read Count Data

Usage

```
deGPS_mRNA(data, dataNormal = NULL, empirical.T.stats = NULL,
group = rep(1:2, each = 5), method = "GP-Theta", nSubcore = 4,
ncore = 4, paired = FALSE, maxIter = 150, geneid = NULL)
```

Arguments

data a matrix containing mRNA gene-level read count data. The column represents

samples while the row represents genes. Biologcial or technical replicates must

be made as new columns in the data.

dataNormal not used anymore.

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empirical.T.stats

A list of empirical T statistics with names as normalization methods, the format of which must be the same as the empirical.T.stats in the returned list of deGPS_mRNA or GPSmle. If null, the empirical T statistics will be calculated. Otherwise, only p values are calculated using given empirical T stats.

group The group indicator. The length of group must equal to ncol(data).

method the methods of normalization. It can be a single charactor or a charactor vector,

the values can be "Lowess", "GP-Quantile", "Quantile", "TMM" or "GP-Theta"

.

nSubcore The parallel computation strategy splits rows of the data, i.e. mRNAs, into

nSubcore parts. The empirical T-stats are calculated for each part of the data, and so are the p values. The total number of cores needed in the computation is

nSubcore * nMethod.

ncore The total cpu cores used for the calculations. The specified ncore can be less

than the total number of cores needed (i.e. nSubcore * nMethod), in which case the cores will be used repeatedly. Apparently, the maximum utilization of the ncore cores is reached when mod(nSubcore * nMethod / ncore) == 0. You may just make ncore = nSubcore to ensure the efficiency of the parallel

computation.

paired The current version of deGPS only contain unpaired test.

maxIter The default value of maxIter is 150. When sample size is large, instead of

transversing every possible permutation, randomly shuffling is applied for maxIter times to obtain the empirical distributions. Larger maxIter costs longer run time. Note that movIter is formed to be not lorger than permutation Times.

time. Note that $\max \texttt{Iter}$ is forced to be not larger than permutation Times.

geneid Gene id of the specified data. Biological or technical replicates must be new

columns in the data, i.e., duplicates in geneid are not allowed.

Details

This function is to analysis mRNA gene-level read count data in two steps: nomalization and permutation based differential expression test.

More than one normalization method can be specified in one run, method GP-Theta is suggested.

In permutation DE test, p values are calculated according to the empirical T statistics obtained by randomly shuffling the samples. You can also specify your own empirical T-stats (or the one you get from another run of the function, which is useful in real data based simulations) in argument empirical.T.stats.

To deal with the burden of computation, e.g. when SampleSize > 10, parallel computing is embeded in the function. By specify nSubcore and ncore, parallel computation is applied in the calculations. The abundant genes are divided into nSubcore subsets, for each of which the empirical T stats are therefore calculated parallelly. The calculation of p values are also parallelly applied on the subsets of mRNAs using empirical T stats obtained by binding nSubcore subsets. nSubcore * nMethod cores are needed in total, where nMethod represents the number of applied methods.

If the number of cores needed in parallel computing process is larger than ncore, cores are iteratively used by the introduced R function foreach (Windows) or mcapply (Linux). The maximum utilization of the ncore cores is reached when mod(nSubcore * nMethod / ncore) == 0.

Besides of specifying large nSubcore and ncore, specify smaller maxIter can be also useful to make the function more efficiency. Note that maxIter should not be too small, where the empirical distribution may not be reliable.

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Value

A GPSmle object is returned.

normalized.data

A list of normalized data, each element represents one speicified normalization method.

log2FoldChange The logrithm of fold change of original data.

empirical.T.stats

A list of the empirical T-stats of normalied data of different normalization meth-

ods, generated by permutation of samples. The length of the T-stats is nRNA * min(permutationTimes (nSample)

 $permutation Times = \begin{pmatrix} nSample \\ nSample/2 \end{pmatrix} / 2, \text{ if each group has equal size.}$ $permutation Times = \begin{pmatrix} nSample \\ group Size \end{pmatrix}, \text{ if each group has unequal size.}$

log2FoldChange The logarithm of fold change of original data.

pvalue The resulting pvalues. Note that the pvalues may be slightly different in dif-

ferent runs of deGPS for the same data when not all possible permutations are

transversed in the calculations of empirical T stats.

paired FALSE

method the normalization methods applied to get the result.

type "mRNA"

Author(s)

Chen Chu

See Also

GPSmle

```
## Not run:
### See the example in "bottomlyData" for real data analysis and the comparison between deGPS
### and other widely-used methods
##Generate Random samples from GP(theta, lambda)
examData <- newExampleData(nRNA = 100, groupSize = 6, lambda = 0.9,
theta = 3, ptol = 1e-15)
str(examData)
##Differential Expression Tests
examRes <- deGPS_mRNA(data = examData$data, group = examData$group,</pre>
method = "GP-Theta", nSubcore = 2, ncore = 2, geneid = paste("G", 1:100, sep = ""))
str(examRes)
topTags(examRes, n = 10, method = "BH")
###Generate simulated RNA-seq data from compcodeR package
require(compcodeR)
samples.per.cond <- 5</pre>
random.outlier.high.prob <- 0.1</pre>
n.vars <- 10000
```

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```
examData <- generateSyntheticData(dataset = "simuData",
n.vars = n.vars, samples.per.cond = samples.per.cond, n.diffexp = floor(n.vars * 0.1),
repl.id = 1, seqdepth = 1e+07, fraction.upregulated = 0.5,
between.group.diffdisp = FALSE, filter.threshold.total = 1,
filter.threshold.mediancpm = 0, fraction.non.overdispersed = 0,
random.outlier.high.prob = random.outlier.high.prob,
output.file = "simuData_repl1.rds")

group <- examData@sample.annotations$condition

examRes <- deGPS_mRNA(data = examData@count.matrix, group = group,
method = "GP-Theta", nSubcore = 2, ncore = 2, geneid = paste("G", 1:n.vars, sep = ""))
str(examRes)
topTags(examRes, n = 10, method = "BH")

## End(Not run)</pre>
```

GPSmle

GPSmle.default

Description

GPSmle.default

Usage

```
GPSmle(data, group = rep(1:2, each = 5),
type = c("normalization", "ecdf", "pvalue"),
method = c("Lowess", "GP-Quantile", "Quantile", "TMM", "GP-Theta"),
maxIter = 500, paired = FALSE, ncpu = 1, geneid = NULL, empirical.T.stats = NULL)
```

Arguments

```
data
                GPSmle.default
                GPSmle.default
group
type
                GPSmle.default
method
                GPSmle.default
maxIter
                GPSmle.default
paired
                GPSmle.default
                GPSmle.default
ncpu
geneid
                GPSmle.default
empirical.T.stats
                GPSmle.default
```

Details

GPSmle.default

See Also

GPSmle.default

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GPSmle.default	Generalized Poisson Statistical Maximum Likelihood Estimation (default)
----------------	---

Description

the default method for the function GPSmle.

Usage

```
## Default S3 method:
GPSmle(data, group = rep(1:2, each = 5),
type = c("normalization", "ecdf", "pvalue"),
method = c("Lowess", "GP-Quantile", "Quantile", "TMM", "GP-Theta"),
maxIter = 500, paired = FALSE, ncpu = 1, geneid = NULL, empirical.T.stats = NULL)
```

Arguments

data a matrix containing microRNA read count data. The column represents samples

while the row represents miRNAs. Biologcial or technical replicates must be

made as new columns in the data.

group The group indicator. The length of group must equal to ncol(data).

type can be "normalization", "ecdf" or "pvalue", to which step GPSmle will

stop. "normalization" means that only normalized data is returned and no DE test is conducted; "ecdf" means the empirical T-stats are generated after nomalization, and the output contains both the normalized data sets and empirical values; "pvalue" means p-values are calculated after the empirical T-stats are obtained, and the output contains the normalized data sets, empirical T-stats and

the p-values of DE test.

method The methods of normalization. More than one method can be specified. The

value can be "Lowess", "Quantile", "TMM", "GP-Quantile", "GP-Theta" or

"GP-MLE2L". See the reference for more details.

maxIter The default value of maxIter is 500. When sample size is large, instead of

transversing every possible permutations, randomly sampling is applied for maxIter to obtain the empirical distributions. Larger maxIter costs longer run time.

Note that ${\tt maxIter}$ is forced to be not larger than ${\tt permutationTimes}$.

paired The current version of deGPS only contain unpaired test.

ncpu The number of cores for the parallel computing. When sample size is large,

the permutation step may be time-consuming. Specify ncpu > 1, parallel computation is applied in the function. ncpu cores are used to calculate maxIter times of permutations, each of which take responsibilies of part of the permutation task. The calculation of p values are also splitted into subsets for parallel

computation.

geneid Gene id of the specified data. Biological or technical replicates must be new

columns in the data, i.e., duplicates in geneid are not allowed.

empirical.T.stats

A list of empirical T statistics with names as normalization methods, the format of which must be the same as the empirical.T.stats in the returned list of deGPS_mRNA or GPSmle. If null, the empirical T statistics will be calculated. Otherwise, only p values are calculated using given empirical T stats.

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Details

This function is to analyze miRNA read count data in two steps: normalization and two-group differential expression test.

More than one normalization method can be specified in method when ncpu = 1. Method GP-Theta is suggested. There are also other choices of the normalization methods. More details about GP-Quantile, GP-Theta can be found in our article.

When sample size is large, maxIter must be specified (500 by default). Smaller maxIter may save run time but to be too small may make the empirical distribution unreliable.

Besides of specifying appropriately small value of maxIter, it is suggested to make ncpu larger than 1, where parallel computation is applied. In parallel computing process, permutation task is splitted into parts of almost equal size, each of which will be processed by a core. And the calculation of p values is also paralleled by dividing the miRs into subsets, each of which is processed by a core.

Value

A GPSmle object. See deGPS_mRNA.

References

deGPS: a Powerful and Flexible Framework for Detecting Differential Expression in RNA-Sequencing Studies

See Also

```
deGPS_mRNA
```

Examples

```
## Not run:
##Generate Random samples from GP(theta, lambda)
examData <- newExampleData(nRNA = 100, groupSize = 2, lambda = 0.9,
theta = 3, ptol = 1e-15)
str(examData)

##Differential Expression Tests for miRNA
examRes <- GPSmle(data = examData$data, group = examData$group, method = "GP-Theta",
type = "pvalue", ncpu = 1, geneid = paste("G", 1:100, sep = ""))
str(examRes)

topTags(examRes, n = 10, method = "BH")
plot(examRes)

## End(Not run)</pre>
```

GPSmleEst

GPSmle.default

Description

```
GPSmle.default
```

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Usage

```
GPSmleEst(data, group = rep(1:2, each = 5),
type = c("normalization", "ecdf", "pvalue"),
dataNormal = NULL, empirical.T.stats = NULL,
method = c("Lowess", "GP", "Quantile", "TMM", "GP2"),
maxIter = 500, paired = FALSE, ncpu = 1, geneid = NULL)
```

Arguments

```
GPSmle.default
data
                GPSmle.default
group
type
                GPSmle.default
dataNormal
                no longer validated
method
                GPSmle.default
maxIter
                GPSmle.default
paired
                GPSmle.default
                GPSmle.default
ncpu
                GPSmle.default
geneid
empirical.T.stats
                GPSmle.default
```

Details

```
GPSmle.default
```

See Also

GPSmle.default

newExampleData

Generate example data for GPSmle and deGPS_mRNA

Description

Randomly generate Generalized Poisson distributed samples with given theta and lambda.

Usage

```
newExampleData(nRNA = 100, groupSize = 5, lambda = 0, theta = 1, ptol = 1e-10)
```

Arguments

nRNA The number of genes or miRs.

groupSize A integer represents The number of samples in each group. Note that the func-

tion can only generate two equal gropus.

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1ambda The lambda parameter of GP distribution. The values must be within (0, 1). Since miRNA/mRNA read counts tend to be overdispersed, we constain the lambda of example data larger than zero to be similar to the real cases. Note that the length of lambda can be either one or two, representing the equal or

unequal lambda for each group.

theta The theta parameter of GP distribution. Must be positive values. It can be a

single value or a numeric vector with length two.

ptol The tolerance of probabilities of GP distribution. The default value is 1e-15,

since regular R can not tell the difference smaller than 1e-15. See details for

more explanations.

Details

The resulting data set contains two GP distributed groups with the specified lambda and theta as the parameters, with each group containing groupSize samples. Note that the length of lambda and theta can be either one or two, representing the same or different GP for two groups. Moreover, the data is neither H0 (non-DE) nor H1 (with DE) data. Every element is a random sample of the given GP and one can not tell whether one single row in the data is DE. However, with large amount of RNAs in the data, it tends to be H0 data, since the variability in one particular row is caused by random assignment of two GP distributions.

The random samples of the specified GP distribution are generated from a multinomial distribution, the domain of which is from zero to a maximum value - gpMax + 1. The larger gpMax is, the closer two distributions are.

The maximum integer gpMax is determined as the minimum integer satisfying $P(x = gpMax) \ge ptol$. The probability for each value from zero to gpMax + 1 is then calculated as the probability of that in specified GP distribution. Note that the probability of $P(x = gpMax + 1) = 1 - P(x = 0) - \ldots - P(x = gpMax)$. Hence, the smaller ptol is, the closer the approximated multinomial distribution is to the specified GP distribution. And since regular R, i.e. without particular package which enables more precise calculations, can not tell differences smaller than 1e-15, ptol is set as 1e-15.

Another way to generate simulated RNA-seq data is the compcodeR package. Details can be found in the package manual and user guide document. See a simple example in the following example session.

Value

group The group indicator of the resulting data.

data The resulting data with GP distributon.

See Also

```
deGPS_mRNA, GPSmle
```

```
## Not run:
####Different Lambda and Theta for Two Groups
examData <- newExampleData(nRNA = 100, groupSize = 2, lambda = c(0.5, 0.9),
theta = c(3, 10), ptol = 1e-15)

####Same Lambda and Theta for Two Groups
examData <- newExampleData(nRNA = 100, groupSize = 2, lambda = 0.9, theta = 3,</pre>
```

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```
###Generate simulated RNA-seq data from compcodeR package
require(compcodeR)

samples.per.cond <- 5
random.outlier.high.prob <- 0.1
n.vars <- 10000

examData <- generateSyntheticData(dataset = "simuData",
n.vars = n.vars, samples.per.cond = samples.per.cond, n.diffexp = floor(n.vars * 0.1),
repl.id = 1, seqdepth = 1e+07, fraction.upregulated = 0.5,
between.group.diffdisp = FALSE, filter.threshold.total = 1,
filter.threshold.mediancpm = 0, fraction.non.overdispersed = 0,
random.outlier.high.prob = random.outlier.high.prob,
output.file = "simuData_repl1.rds")

## End(Not run)</pre>
```

plot.GPSmle

Plot GPSmle

Description

Plot the histograms of GPSmle results.

Usage

```
## S3 method for class GPSmle plot(x, ...)
```

Arguments

x the object returned by GPSmle.

... the parameters of plot

Details

See GPSmle. default for more details of the output of GPSmle.

Value

The output depends on the specification of type in GPSmle. If type = "normalization", the histograms of normalized data sets are returned. So are the "ecdf" and "pvalue" or "mRNA" in deGPS_mRNA.

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summary.GPSmle

Summary of GPSmle

Description

Summary of GPSmle

Usage

```
## S3 method for class GPSmle
summary(object, ...)
```

Arguments

object the GPSmle object returned by GPSmle or deGPS_mRNA.

... see the summary.default

Details

summary of the GPSmle

Value

summary of the GPSmle

topTags

The top significant genes or miRs

Description

The top significant genes or miRs

Usage

```
topTags(x, n = 10, method = "BH", significance = 0.05)
```

Arguments

x A GPSmle object returned by GPSmle or deGPS_mRNA or a p value vector. If it is

a GPSmle object, only one column of p values is allowed in this function.

n the number of required top significant genes or miRs.

method the adjust method of multiple testing p values, same as R function p.adjust.

significance the significant level of the DE.

Details

This function is to find the significant genes or miRs at the given significant level. If you want to call this function, make sure that library(deGPS) is called after other packages, such as edgeR, since those packages also contain function named topTags.

16 topTags

Value

A list containing:

pvalue the ordered p values of significant genes or miRs.

adj.pvalue the ordered adjusted p values by specified method.

 $\mbox{method} \qquad \mbox{p value adjusted method, same as the method in R function p.adjust}$

geneName names of the genes or miRs.

geneid the row indice of the genes or miRs in given p values.

significance the specified significant level.

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