Package 'mCSEA'

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Type Package

Title Methylated CpGs Set Enrichment Analysis

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Description Identification of differentially methylated regions (DMRs) in predefined regions (promoters, CpG islands...) from the human genome using Illumina's 450K or EPIC microarray data.

Provides methods to rank CpG probes based on linear models and includes plotting functions.

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2 mCSEA-package

R topics documented:

	mCSEA-package	2
	exprTest	3
	mCSEAIntegrate	3
	mCSEAPlot	4
	mCSEAPlotGSEA	6
	mCSEATest	7
	precomputedmCSEA	8
	rankProbes	9
Index	1	11

mCSEA-package

Methylated CpGs Set Enrichment Analysis

Description

Identification of differentially methylated regions (DMRs) in predefined regions (promoters, CpG islands...) from the human genome using Illumina's 450K or EPIC microarray data. Provides methods to rank CpG probes based on linear models and includes plotting functions.

Author(s)

Jordi Martorell Marugán

Maintainer: Jordi Martorell Marugán<jordi.martorell@genyo.es>

```
## Not run:
library(mCSEA)
data(mcseadata)
myRank <- rankProbes(betaTest, phenoTest, refGroup = "Control")
myResults <- mCSEATest(myRank, regionsTypes = "promoters", platform = "EPIC")
## End(Not run)
data(precomputedmCSEA)
head(myResults$promoters)</pre>
```

exprTest 3

exprTest	Expression data example	

Description

exprTest is a subset of 100 genes' microarray expression data for 20 bone marrow samples: 10 from Acute Lymphoblastic Leukemia patients and 10 from healthy patients. It is useful to test mCSEAIntegrate function.

Usage

```
data(exprTest)
```

Format

matrix

Source

Obtained from the leukemiasEset data package

mCSEAIntegrate	Integrate methylation and expression	
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Description

Uses mCSEA methylation analysis results and expression values to search for significant correlations between DMRs methylation and close genes expression.

Usage

```
mCSEAIntegrate(mCSEAResults, exprData, regionType = c("promoters", "genes",
    "CGI", "custom"), geneIDs = "SYMBOL", dmrName = NULL, pcutoff = 0.05,
    minCor = 0.5, minP = 0.05, makePlot = TRUE, folder = ".", nproc = 1)
```

Arguments

mCSEAResults	The object generated by mCSEATest function
exprData	A matrix or data frame with genes in rows and samples in columns. A SummarizedExperiment object can be used too
regionType	The region types to be represented. Must be one or more of "promoters", "genes", "CGI" and "custom" $$
geneIDs	Gene identifiers used in exprData. One of "SYMBOL", "ENSEMBL", "ENTREZID", "GENEID", "REFSEQ" or "UNIGENE"

4 mCSEAPlot

dmrName	The DMR of interest to correlate with expression (e.g. gene name, CGI name). If NULL (default), all DMRs with P-Value < pcutoff are selected
pcutoff	P-Value threshold to select DMRs if dmrName = NULL
minCor	Correlation threshold to output the results
minP	Correlation P-Value threshold to output the results
makePlot	If TRUE, generate corelation and save them in the folder specified by folder parameter
folder	Directory to save the correlation plots if makePlot = TRUE
nproc	Number of processors to be used

Value

A data.frame with the integration results.

Author(s)

Jordi Martorell Marugán, <jordi.martorell@genyo.es>

See Also

```
rankProbes, mCSEATest
```

Examples

mCSEAPlot

Plot mCSEA results

Description

Generate a graphic with the genomic context of the selected DMR, showing methylation status at each CpG site of different samples groups

Usage

```
mCSEAPlot(mCSEAResults, regionType, dmrName, extend = 1000,
    chromosome = TRUE, leadingEdge = TRUE, CGI = FALSE, genes = TRUE,
    transcriptAnnotation = "transcript", makePDF = TRUE,
    col = c("blue", "magenta", "green", "red", "black"))
```

mCSEAPlot 5

Arguments

mCSEAResults The object generated by mCSEATest function

regionType The region type to be represented. Must be one of "promoters", "genes", "CGI"

or "custom"

dmrName The DMR of interest to be represented (e.g. gene name, CGI name...)

extend The number of base pairs to extend the plot around the DMR of interest (default

= 1000 bp

chromosome If TRUE, represent the chromosome and genome axis

leadingEdge If TRUE, represent the bars indicating if each CpG belongs to the mCSEA lead-

ing edge (green) or not (red)

CGI If TRUE, represent the annotated CpG islands

genes If TRUE, represent the annotated genes

transcriptAnnotation

Labels showed at the genes track. Must be one of "transcript" (default), "sym-

bol", "gene", "exon" or "feature"

makePDF If TRUE, save the plot in pdf format in the working directory. Otherwise, draw

the plot in the active graphics window

col Vector with colors to plot methylation in different groups

Value

'NULL'

Author(s)

Jordi Martorell Marugán, <jordi.martorell@genyo.es>

See Also

```
rankProbes, mCSEATest, mCSEAPlotGSEA
```

```
library(mCSEAdata)
data(mcseadata)
## Not run:
myRank <- rankProbes(betaTest, phenoTest, refGroup = "Control")
set.seed(123)
myResults <- mCSEATest(myRank, betaTest, phenoTest,
regionsTypes = "promoters", platform = "EPIC")

## End(Not run)
data(precomputedmCSEA)
mCSEAPlot(myResults, "promoters", "CLIC6",
transcriptAnnotation = "symbol", makePDF = FALSE)</pre>
```

6 mCSEAPlotGSEA

mCSEAPlotGSEA

Plot mCSEA results

Description

Generate an enrichment plot

Usage

```
mCSEAPlotGSEA(rank, mCSEAResults, regionType, dmrName)
```

Arguments

rank A named numeric vector with the ranking statistic of each CpG site

mCSEAResults The object generated by mCSEATest function

regionType The region type to be represented. Must be one of "promoters", "genes", "CGI"

or "custom"

dmrName The DMR of interest to be represented (e.g. gene name, CGI name...)

Value

'NULL'

Author(s)

Jordi Martorell Marugán, <jordi.martorell@genyo.es>

References

Subramanian, A. et al (2005). Gene set enrichment analysis: A knowledge-based approach for interpreting genome-wide expression profiles . PNAS 102, 15545-15550.

See Also

```
rankProbes, mCSEATest, mCSEAPlot
```

```
## Not run:
library(mCSEAdata)
data(mcseadata)
myRank <- rankProbes(betaTest, phenoTest, refGroup = "Control")
set.seed(123)
myResults <- mCSEATest(myRank, regionsTypes = "promoters",
platform = "EPIC")
## End(Not run)
data(precomputedmCSEA)
mCSEAPlotGSEA(myRank, myResults, "promoters", "CLIC6")</pre>
```

mCSEATest 7

Description

Perform a methylated CpG sites enrichment analysis in predefined genomic regions

Usage

```
mCSEATest(rank, methData, pheno = NULL, column = 1,
  regionsTypes = c("promoters", "genes", "CGI"), customAnnotation = NULL,
  minCpGs = 5, nproc = 1, nperm = NULL, platform = "450k")
```

Arguments

rank	A named numeric vector with the ranking statistic of each CpG site	
methData	A data frame or a matrix containing Illumina's CpG probes in rows and samples in columns. A SummarizedExperiment object can be used too	
pheno	A data frame or a matrix containing samples in rows and covariates in columns. If NULL (default), pheno is extracted from the SummarizedExperiment object	
column	The column name or number from pheno used to split the samples into groups (first column is used by default)	
regionsTypes	A character or character vector indicating the predefined regions to be analyzed. NULL to skip this step and use customAnnotation.	
customAnnotation		
	An optional list with the CpGs associated to each feature (default = NULL)	
minCpGs	Minimum number of CpGs associated to a region. Regions below this threshold are not tested	
nproc	Number of processors to use in GSEA step (default = 1)	
nperm	(deprecated) Number of permutations to do in GSEA step in the previous implementation. Now, this parameter is ignored	
platform	Platform used to get the methylation data (450k or EPIC)	

Value

A list with the results of each of the analyzed regions. For each region type, a data frame with the results and a list with the probes associated to each region are generated. In addition, this list also contains the input methData, pheno and platform objects

Author(s)

Jordi Martorell Marugán, <jordi.martorell@genyo.es>

8 precomputedmCSEA

References

Subramanian, A. et al (2005). Gene set enrichment analysis: A knowledge-based approach for interpreting genome-wide expression profiles . PNAS 102, 15545-15550.

See Also

```
rankProbes, mCSEAPlot, mCSEAPlotGSEA
```

Examples

```
## Not run:
library(mCSEAdata)
data(mcseadata)
myRank <- rankProbes(betaTest, phenoTest, refGroup = "Control")
set.seed(123)
myResults <- mCSEATest(myRank, betaTest, phenoTest,
regionsTypes = "promoters", platform = "EPIC")

## End(Not run)
data(precomputedmCSEA)
head(myResults[["promoters"]])
head(myResults[["promoters_association"]])</pre>
```

precomputedmCSEA

Precomputed mCSEA results

Description

myRank is a result of rankProbes function. myResults is a result of mCSEATest function.

Usage

```
data(precomputedmCSEA)
```

Format

```
vector (myRank) and list data.frame (myResults)
```

Source

Both objects were obtained with the example commands in the mCSEA vignette.

rankProbes 9

rankProbes	Rank CpG probes		
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Description

Apply a linear model to Illumina's 450k or EPIC methylation data to get the t-value of each CpG probe

Usage

```
rankProbes(methData, pheno = NULL, paired = FALSE, explanatory = 1,
  covariates = c(), pairColumn = c(), caseGroup = 1, refGroup = 2,
  continuous = NULL, typeInput = "beta", typeAnalysis = "M")
```

Arguments

methData	A data frame or a matrix containing Illumina's CpG probes in rows and samples in columns. A SummarizedExperiment object can be used too
pheno	A data frame or a matrix containing samples in rows and covariates in columns. If NULL (default), pheno is extracted from the SummarizedExperiment object
paired	Perform a paired t-test (default = FALSE)
explanatory	The column name or position from pheno used to perform the comparison between groups (default = first column)
covariates	A list or character vector with column names from pheno used as data covariates in the linear model
pairColumn	Only for paired analysis. The column name or position from pheno used to connect the paired samples (default = NULL)
caseGroup	The group name or position from explanatory variable used as cases to perform the comparison (default = first group)
refGroup	The group name or position from explanatory variable used as reference to perform the comparison (default = second group)
continuous	A list or character vector with columns names from pheno which should be treated as continuous variables (default = none)
typeInput	Type of input methylation data. "beta" for Beta-values and "M" for M-values
typeAnalysis	"M" to use M-values to rank the CpG probes (default). "beta" to use Beta-values instead

Value

A named vector containing the t-values from the linear model for each CpG probe

Author(s)

Jordi Martorell Marugán, <jordi.martorell@genyo.es>

10 rankProbes

References

Smyth, G. K. (2005). *Limma: linear models for microarray data*. Bioinformatics and Computational Biology Solutions using R and Bioconductor, 397-420.

See Also

 ${\tt mCSEATest}$

```
data(mcseadata)
myRank <- rankProbes(betaTest, phenoTest, refGroup = "Control")
head(myRank)</pre>
```

Index

```
* datasets
exprTest, 3
precomputedmCSEA, 8

exprTest, 3

mCSEA (mCSEA-package), 2
mCSEA-package, 2
mCSEAIntegrate, 3
mCSEAPlot, 4, 6, 8
mCSEAPlotGSEA, 5, 6, 8
mCSEATest, 4-6, 7, 10
myRank (precomputedmCSEA), 8
myResults (precomputedmCSEA), 8
precomputedmCSEA, 8

rankProbes, 4-6, 8, 9
```