

Package ‘broadSeq’

July 9, 2025

Title broadSeq : for streamlined exploration of RNA-seq data

Version 1.2.2

Description This package helps user to do easily RNA-seq data analysis with multiple methods (usually which needs many different input formats). Here the user will provide the expression data as a SummarizedExperiment object and will get results from different methods. It will help user to quickly evaluate different methods.

License MIT + file LICENSE

URL <https://github.com/dasroy/broadSeq>

BugReports <https://github.com/dasroy/broadSeq/issues>

Encoding UTF-8

Roxygen list(markdown = TRUE)

RoxygenNote 7.2.3

biocViews GeneExpression, DifferentialExpression, RNASeq,
Transcriptomics, Sequencing, Coverage, GeneSetEnrichment, GO

Suggests knitr, limma (>= 3.54.0), rmarkdown, stats (>= 4.2.2), samr

Depends dplyr, ggpibr, SummarizedExperiment

Imports BiocStyle, DELocal, EBSeq (>= 1.38.0), DESeq2 (>= 1.38.2),
NOISeq, forcats (>= 1.0.0), genefilter, ggplot2, ggplotify,
plyr, clusterProfiler (>= 4.8.2), pheatmap, sechm (>= 1.6.0),
stringr, purrr (>= 0.3.5), edgeR (>= 3.40.1)

VignetteBuilder knitr

git_url <https://git.bioconductor.org/packages/broadSeq>

git_branch RELEASE_3_21

git_last_commit c89e96c

git_last_commit_date 2025-06-10

Repository Bioconductor 3.21

Date/Publication 2025-07-09

Author Rishi Das Roy [aut, cre] (ORCID:
<https://orcid.org/0000-0002-3276-7279>)

Maintainer Rishi Das Roy <rishi.dasroy@gmail.com>

Contents

broadSeq-package	2
combinedEnrichment	3
genes_plot	4
normalizeEdgerCPM	5
plotHeatmapCluster	6
plot_MDS	7
prcompTidy	8
round_df	9
sampleAssay_plot	10
transformDESeq2	10
use_DELocal	11
use_deseq2	12
use_EBSeq	13
use_edgeR_GLM	14
use_limma_trend	15
use_multDE	16
use NOIseq	17
use_SAMseq	18
volcanoPlot	19

Index

21

broadSeq-package	<i>broadSeq : for streamlined exploration of RNA-seq data</i>
------------------	---

Description

This package helps user to do easily RNA-seq data analysis with multiple methods (usually which needs many different input formats). Here the user will provide the expression data as a SummarizedExperiment object and will get results from different methods. It will help user to quickly evaluate different methods.

Author(s)

Maintainer: Rishi Das Roy <rishi.dasroy@gmail.com> ([ORCID](#))

See Also

Useful links:

- <https://github.com/dasroy/broadSeq>
- Report bugs at <https://github.com/dasroy/broadSeq/issues>

combinedEnrichment	<i>Provides GO gene set enrichment and over-representation analysis</i>
--------------------	---

Description

This wrapper function combines clusterProfiler::gseGO and clusterProfiler::enrichGO. The input type of these two methods are different; order ranked geneList and a vector of entrez gene id. Here combinedEnrichment function internally generates these two data types from user defined DEG_table (differentially expressed genes).

Usage

```
combinedEnrichment(
  DEG_table,
  geneCol = "ID",
  logCol = "logFoldChange",
  OrgDB = "org.Hs.eg.db",
  keyType,
  universe,
  ont = "BP",
  logfoldCut = 1,
  pvalueCutoff = 0.05,
  qvalueCutoff = 0.05
)
```

Arguments

DEG_table	A data.frame atleast with two columns.
geneCol	The column name of DEG_table which provides gene ids and should be compatible with keytype parameter.
logCol	The column name of DEG_table which provides logfold(numeric) values to create a order ranked geneList for gseGO funtion.
OrgDB	OrgDb; passed to clusterProfiler functions
keyType	keytype of input gene(geneCol). One of the keytypes(OrgDB); passed to clusterProfiler functions
universe	background genes; passed to clusterProfiler::enrichGO.
ont	one of "BP", "MF", and "CC" subontologies, or "ALL" for all three.; passed to clusterProfiler functions
logfoldCut	to filter genes based on parameter logCol
pvalueCutoff	; passed to clusterProfiler functions
qvalueCutoff	; passed to clusterProfiler functions

Value

a named list of three data.frames which are output of gseGO("gseResult") and enrichGO ("oraUP" and "oraDOWN").

genes_plot	<i>Expression of multiple genes/features from a single assay as boxplot (or added dotplot)</i>
------------	--

Description

Expression of multiple genes/features from a single assay as boxplot (or added dotplot)

Boxplot of a single gene/feature from multiple assays

Usage

```
genes_plot(se, features, assayName = "counts", facet.by = "feature", x, ...)  
assay_plot(se, feature, assayNames = c("counts"), x, ...)
```

Arguments

se	Object of <code>SummarizedExperiment</code> class
features	a character vector of rownames or named list of character vectors where name is one of the colnames of rowData.
assayName	One of the values from <code>SummarizedExperiment::assayNames(se)</code> ; default is "counts" assay
facet.by	must be one of the column names of <code>rowData(se)</code> . default "feature" which is equivalent to rownames of <code>rowData</code>
x	a column name of <code>colData</code> which will be used in x-axis
...	other arguments to be passed to <code>ggpubr::ggbboxplot</code>
feature	a character vector of rownames or named list of character vectors where name is one of the column of <code>rowData</code> .
assayNames	names from <code>SummarizedExperiment::assayNames(se)</code> ; default value is "counts"

Value

ggplot object

return an object of class ggarrange, which is a ggplot or a list of ggplot.

Examples

```

se <- readRDS(system.file("extdata", "rat_vole_mouseSE_salmon.rds",
  package = "broadSeq"))
# The normalized values are added with the assay name "logCPM"
se <- broadSeq::normalizeEdgerCPM(se ,method = "none", cpm.log = TRUE )

broadSeq::genes_plot(se,
  features = list(mouse_gene_id = c("ENSMUSG0000022510" ,
  "ENSMUSG0000027985"))),

```

```

facet.by = "symbol", # column of rowData
x = "stage", fill="stage")

broadSeq::genes_plot(se,
                     features = list(symbol=c("Shh", "Edar") ),
                     facet.by = "symbol", # column of rowData
                     x = "stage", fill="stage")

broadSeq::assay_plot(se, feature = c("Shh"),
                     assays = c("counts", "logCPM"),
                     x = "stage", fill="stage", add="dotplot", palette = "npg")

```

normalizeEdgerCPM*Use of edgeR package to normalize count data***Description**

Use of edgeR package to normalize count data

Usage

```
normalizeEdgerCPM(se, method = "TMM", cpm.log = TRUE, ...)
```

Arguments

<code>se</code>	Object of SummarizedExperiment class
<code>method</code>	value for <code>edgeR::normLibSizes</code> function. default "TMM"
<code>cpm.log</code>	value for <code>edgeR::cpm</code> function. default TRUE
<code>...</code>	passed to <code>normLibSizes</code> function

Value

Object of [SummarizedExperiment](#) class where a new assay is added to the input object.

Examples

```

se <- readRDS(system.file("extdata",
                           "rat_vole_mouseSE_salmon.rds",
                           package = "broadSeq"))

se <- broadSeq::normalizeEdgerCPM(se , method = "TMM", cpm.log = FALSE )
# The normalized values are added with the assay name "TMM"
SummarizedExperiment::assayNames(se)

```

`plotHeatmapCluster` *Plot clustered heatmaps*

Description

Plot clustered heatmaps from `SummarizedExperiment` with `pheatmap` and return object as `ggplot`

Usage

```
plotHeatmapCluster(
  se,
  scaledAssay = "vst",
  ntop = 500L,
  features = NULL,
  show_geneAs = NULL,
  annotation_col = NA,
  annotation_row = NA,
  ...
)
```

Arguments

<code>se</code>	Object of <code>SummarizedExperiment</code> class
<code>scaledAssay</code>	an scaled assay name from <code>SummarizedExperiment::assayNames(se)</code>
<code>ntop</code>	number of most-variable genes to select. Igored if "features" is specified.
<code>features</code>	character vector features/genes to be used to measure distance between the samples
<code>show_geneAs</code>	a character vector of colnames of <code>rowData(se)</code>
<code>annotation_col</code>	a character vector of colnames of <code>colData(se)</code>
<code>annotation_row</code>	a list of character vector of colnames of <code>rowData(se)</code>
<code>...</code>	other arguments like color or shape whose values should be similar to <code>colData</code> columns names passed to <code>pheatmap</code>

Value

`ggplot` object

Examples

```
se <- readRDS(system.file("extdata","rat_vole_mouseSE_salmon.rds", package = "broadSeq"))

se <- broadSeq::normalizeEdgerCPM(se ,method = "none", cpm.log = TRUE )

broadSeq::plotHeatmapCluster(
  se,
  scaledAssay = "logCPM",
```

```

annotation_col = c("species", "stage"),
annotation_row = c("Class", "gene_biotype"),
ntop = 30, show_geneAs = "symbol",
cluster_cols = TRUE, cluster_rows = FALSE,
show_rownames = TRUE, show_colnames = FALSE,
main = "Top 30 variable gene vst"
)

```

plot_MDS*Classical multidimensional scaling***Description**

Classical multidimensional scaling is based on measuring the distance between the samples.

Usage

```
plot_MDS(se, scaledAssay = "vst", ntop = 500L, features = NULL, ...)
```

Arguments

<code>se</code>	Object of SummarizedExperiment class
<code>scaledAssay</code>	an scaled assay name from <code>SummarizedExperiment::assayNames(se)</code>
<code>ntop</code>	number of most-variable genes to select. Igored if "features" is specified.
<code>features</code>	character vector features/genes to be used to measure distance between the samples
<code>...</code>	other arguments like color or shape whose values should be similar to <code>colData</code> columns names passed to ggpubr::ggsca ter

Value

ggplot object

Examples

```

se <- readRDS(system.file("extdata", "rat_vole_mouseSE_salmon.rds", package = "broadSeq"))

se <- broadSeq::transformDESeq2(se, method = "vst" )

broadSeq::plot_MDS(se, scaledAssay = "vst", ntop=500,
                    color = "species", shape = "stage",
                    ellipse=TRUE, legend = "bottom")

```

prcompTidy*Perform Principal Components Analysis*

Description

This function returns the results of stats::[prcomp](#) in a tidy list format. This is more flexible for further custom PCA , biplot and exploring gene(factor) loading of the PCA.

Usage

```
prcompTidy(se, scaledAssay = "vst", ntop = 500L, features = NULL, ...)
plotAnyPC(computedPCA, x = 1, y = 2, ...)
biplotAnyPC(computedPCA, x = 1, y = 2, genes = NULL, genesLabel = NULL, ...)
getFeatureLoadRanking(computedPCA, pcs = seq_len(5), topN = 10, keep)
```

Arguments

<code>se</code>	Object of SummarizedExperiment class
<code>scaledAssay</code>	an scaled assay name from <code>SummarizedExperiment::assayNames(se)</code>
<code>ntop</code>	number of most-variable genes to select. Igored if "features" is specified.
<code>features</code>	character vector features/genes to be used for PCA
<code>...</code>	other arguments like color or shape whose values should be similar to <code>colData</code> columns names passed to <code>ggpubr::ggscatter</code>
<code>computedPCA</code>	a list of data.frame returned by prcompTidy
<code>x</code>	PC number for x-axis default 1
<code>y</code>	PC number for y-axis default 2
<code>genes</code>	if <code>genes</code> is NULL then top max and min loaded genes of each PCs are plotted
<code>genesLabel</code>	one of <code>rowData</code> column names
<code>pcs</code>	The numbers of PCs
<code>topN</code>	Number of features per PC
<code>keep</code>	the column names of <code>rowData</code> to keep the corresponding information

Details

[Reused code](#)

Value

a list with four `data.frame` objects: `pc_scores`, `eigen_values`, `loadings` (eigen vectors) and the original data.

`ggplot` object

`ggplot` object

a `data.frame`

Examples

```
se <- readRDS(system.file("extdata","rat_vole_mouseSE_salmon.rds", package = "broadSeq"))

se <- broadSeq::normalizeEdgerCPM(se ,method = "none",cpm.log = TRUE )
computedPCA_logCPM <- broadSeq::prcompTidy(se, scaledAssay = "logCPM", ntop = 500)

plotAnyPC(computedPCA = computedPCA_logCPM, x = 1, y = 2, color = "species",
          shape = "stage",legend = "bottom")
plotAnyPC(computedPCA = computedPCA_logCPM, x = 2, y = 3, color = "species",
          shape = "stage",legend = "bottom")

computedPCA_logCPM$eigen_values %>%
  dplyr::filter(var_exp >= 0.5) %>% # Selecting PC explaining more than 1% variance
  ggbarplot(x="PC",y="var_exp", label = TRUE, label.pos = "out")
```

`round_df`

Applies round function only on numeric columns of a data.frame.

Description

Applies `round` function only on numeric columns of a `data.frame`.

Usage

```
round_df(df, digits)
```

Arguments

<code>df</code>	data.frame object
<code>digits</code>	passed to <code>round</code>

Value

`data.frame` object

Examples

```
data("iris")
iris %>% round_df(digits = 0) %>% head()
```

sampleAssay_plot	<i>Useful to visualize distribution of assay values for each sample. Plots 'boxplot' of any assay for each sample. Aesthetic can be added from colData.</i>
------------------	---

Description

Useful to visualize distribution of assay values for each sample. Plots 'boxplot' of any assay for each sample. Aesthetic can be added from colData.

Usage

```
sampleAssay_plot(se, assayName = "counts", ...)
```

Arguments

se	Object of SummarizedExperiment class
assayName	One of the values from <code>SummarizedExperiment::assayNames(se)</code>
...	other arguments to be passed to <code>ggpubr::ggbboxplot</code>

Value

ggplot object

Examples

```
se <- readRDS(system.file("extdata","rat_vole_mouseSE_salmon.rds", package = "broadSeq"))

sampleAssay_plot(se, assayName = "counts",
fill="stage", # stage is a column name of colData(se)
yscale="log2")

se <- broadSeq::normalizeEdgerCPM(se ,method = "none",cpm.log = TRUE )

sampleAssay_plot(se, assayName = "logCPM", fill="stage")
```

transformDESeq2	<i>Transform SummarizedExperiment with DESeq2 package</i>
-----------------	---

Description

To use `SummarizedExperiment` with `DESeq2`, this function makes sure that 'counts' assay should be the first in assays list and the mode is integer.

Usage

```
transformDESeq2(se, method = "vst", ...)
```

Arguments

se	Object of SummarizedExperiment class
method	"vst", "normTransform" or "rlog" to choose either of DESeq2:: varianceStabilizingTransformation DESeq2:: normTransform and DESeq2:: rlog function. default is "vst"
...	arguments passed to varianceStabilizingTransformation normTransform and rlog

Value

Object of [SummarizedExperiment](#) class where a new assay is added to the input object.

Examples

```
se <- readRDS(system.file("extdata",
  "rat_vole_mouseSE_salmon.rds",
  package = "broadSeq"))

se <- broadSeq::transformDESeq2(se, method = "vst" )
# The transformed values are added with the assay name "vst"
SummarizedExperiment::assayNames(se)
```

use_DELocal

*To use SummarizedExperiment with package DELocal***Description**

A wrapper function of DELocal where input is an object of [SummarizedExperiment](#)

Usage

```
use_DELocal(se, colData_id, control, treatment, rank = FALSE, ...)
```

Arguments

se	Object of SummarizedExperiment class
colData_id	One of the columns of colData(se). It should be factors of more than one value.
control	Base level and one of the factor values of colData(se)[[colData_id]]
treatment	one of the factor values of colData(se)[[colData_id]]
rank	Logical value default FALSE. If true the result will have an additional column named "rank" and the results are ranked on "relative.logFC"
...	other arguments to be passed to main function DELocal:: DELocal .

Value

a data.frame from DELocal

Examples

```
se <- readRDS(system.file("extdata",
  "rat_vole_mouseSE_salmon.rds",
  package = "broadSeq"))

# To reduce runtime
se <- se[rowData(se)$chromosome_name == 2,colData(se)$species == "Mouse"]

result <-
  use_DELocal(se = se,
    colData_id = "stage", control = "Bud", treatment = "Cap",
    rank = TRUE)
```

use_deseq2

To use SummarizedExperiment with package DESeq2

Description

A wrapper function of DESeq2 where input is an object of [SummarizedExperiment](#)

Usage

```
use_deseq2(se, colData_id, control, treatment, rank = FALSE, ...)
```

Arguments

se	Object of SummarizedExperiment class
colData_id	One of the columns of colData(se). It should be factors of more than one value.
control	Base level and one of the factor values of colData(se)[[colData_id]]
treatment	one of the factor values of colData(se)[[colData_id]]
rank	Logical value default FALSE. If true the result will have an additional column named "rank" and the results are ranked on "padj"
...	other arguments to be passed to main function DESeq2:: results .

Value

a data.frame converted from DESeq2::[DESeqResults](#)

Examples

```
se <- readRDS(system.file("extdata",
  "rat_vole_mouseSE_salmon.rds",
  package = "broadSeq"))

# To reduce runtime
se <- se[rowData(se)$chromosome_name == 2,colData(se)$species == "Mouse"]
```

```
result <-
  use_deseq2(se = se,
             colData_id = "stage", control = "Bud", treatment = "Cap",
             rank = TRUE)
```

use_EBSeq*To use SummarizedExperiment with package EBSeq***Description**

A wrapper function of EBSeq where input is an object of [SummarizedExperiment](#)

Usage

```
use_EBSeq(se, colData_id, control, treatment, rank = FALSE, ...)
```

Arguments

<code>se</code>	Object of SummarizedExperiment class
<code>colData_id</code>	One of the columns of <code>colData(se)</code> . It should be factors of more than one value.
<code>control</code>	Base level and one of the factor values of <code>colData(se)[[colData_id]]</code>
<code>treatment</code>	one of the factor values of <code>colData(se)[[colData_id]]</code>
<code>rank</code>	Logical value default FALSE. If true the result will have an additional column named "rank" and the results are ranked on "PPDE"
<code>...</code>	other arguments to be passed to main function <code>EBSeq::GetDEResults</code> .

Value

a data.frame object converted from the output of `EBSeq::GetDEResults`.

Examples

```
se <- readRDS(system.file("extdata",
                           "rat_vole_mouseSE_salmon.rds",
                           package = "broadSeq"))

# To reduce runtime
se <- se[rowData(se)$chromosome_name == 2,colData(se)$species == "Mouse"]

result <-
  use_EBSeq(se = se,
            colData_id = "stage", control = "Bud", treatment = "Cap",
            rank = TRUE)
```

`use_edgeR_GLM`

To use SummarizedExperiment with package edgeR

Description

A wrapper function of DESeq2 where input is an object of [SummarizedExperiment](#)

Usage

```
use_edgeR_GLM(se, colData_id, control, treatment, rank = FALSE, ...)

use_edgeR_exact(se, colData_id, control, treatment, rank = FALSE, ...)

use_edgeR(
  se,
  colData_id,
  control,
  treatment,
  rank = FALSE,
  edgeR.n = Inf,
  edgeR.adjust.method = "BH",
  edgeR.sort.by = "PValue",
  option = "GLM",
  ...
)
```

Arguments

<code>se</code>	Object of SummarizedExperiment class
<code>colData_id</code>	One of the columns of <code>colData(se)</code> . It should be factors of more than one value.
<code>control</code>	Base level and one of the factor values of <code>colData(se)[[colData_id]]</code>
<code>treatment</code>	one of the factor values of <code>colData(se)[[colData_id]]</code>
<code>rank</code>	Logical value default FALSE. If true the result will have an additional column named "rank"
<code>...</code>	other arguments to be passed to <code>edgeR::glmLRT</code> or <code>edgeR::exactTest</code>
<code>edgeR.n</code>	argument for <code>edgeR::topTags</code>
<code>edgeR.adjust.method</code>	argument for <code>edgeR::topTags</code>
<code>edgeR.sort.by</code>	argument for <code>edgeR::topTags</code>
<code>option</code>	"GLM" or "exact" to indicate to use either <code>edgeR::glmLRT</code> or <code>edgeR::exactTest</code>

Value

a `data.frame` of output from `edgeR::topTags`

Examples

```
se <- readRDS(system.file("extdata",
  "rat_vole_mouseSE_salmon.rds",
  package = "broadSeq"))

# To reduce runtime
se <- se[rowData(se)$chromosome_name == 2,colData(se)$species == "Mouse"]

result <-
  use_edgeR(se = se,
            colData_id = "stage", control = "Bud", treatment = "Cap",
            rank = TRUE)
```

use_limma_trend

To use SummarizedExperiment with package limma

Description

A wrapper function of limma where input is an object of [SummarizedExperiment](#)

Usage

```
use_limma_trend(se, colData_id, control, treatment, rank = FALSE, ...)

use_limma_voom(se, colData_id, control, treatment, rank = FALSE, ...)

use_limma(
  se,
  colData_id,
  control,
  treatment,
  rank = FALSE,
  useVoom = TRUE,
  showPlot = FALSE,
  limma.adjust = "BH",
  limma.sort.by = "p",
  limma.number = Inf,
  ...
)
```

Arguments

se	Object of SummarizedExperiment class
colData_id	One of the columns of colData(se). It should be factors of more than one value.
control	Base level and one of the factor values of colData(se)[[colData_id]]
treatment	one of the factor values of colData(se)[[colData_id]]

rank	Logical value default FALSE. If true the result will have an additional column named "rank"
...	other arguments to be passed to main function edgeR::calcNormFactors .
useVoom	whether to use limma::voom or edgeR::cpm
showPlot	whether to use limma::plotSA ; default FALSE
limma.adjust	argument for limma::topTable
limma.sort.by	argument for limma::topTable
limma.number	argument for limma::topTable

Value

a data.frame of output from limma::topTable

Examples

```
se <- readRDS(system.file("extdata",
  "rat_vole_mouseSE_salmon.rds",
  package = "broadSeq"))

# To reduce runtime
se <- se[rowData(se)$chromosome_name == 2,colData(se)$species == "Mouse"]
result <-
  use_limma(se = se,
            colData_id = "stage", control = "Bud", treatment = "Cap",
            rank = TRUE)
```

use_multDE

To identify differentially expressed genes by multiple methods

Description

To identify differentially expressed genes by multiple methods

Usage

```
use_multDE(
  deFun_list,
  return.df = FALSE,
  se,
  colData_id,
  control,
  treatment,
  ...
)
```

Arguments

deFun_list	a list of function which can perform differential expression analysis
return.df	whether to return all results aggregated form of data.frame or a list of results. Default is FALSE
se	Object of SummarizedExperiment class
colData_id	One of the columns of colData(se). It should be factors of more than one value.
control	Base level and one of the factor values of colData(se)[[colData_id]]
treatment	one of the factor values of colData(se)[[colData_id]]
...	other arguments to be passed to functions listed in deFun_list

Value

a list or data.frame

Examples

```
se <- readRDS(system.file("extdata",
  "rat_vole_mouseSE_salmon.rds",
  package = "broadSeq"))

# To reduce runtime
se <- se[rowData(se)$chromosome_name == 2,colData(se)$species == "Mouse"]

# First define a named list of functions
fun <- list(limma_trend = use_limma_trend, limma_voom = use_limma_voom,
  edgeR_exact = use_edgeR_exact, edgeR_glm = use_edgeR_GLM,
  deseq2 = use_deseq2,
  DELocal = use_DELocal, noiseq = use_NOIseq,
  EBSeq = use_EBSeq)

multi_result <- broadSeq::use_multDE(
  se <- se[rowData(se)$chromosome_name == 2,colData(se)$species == "Mouse"],
  deFun_list = fun, return.df = TRUE,
  colData_id = "stage", control = "Bud", treatment = "Cap",
  rank = TRUE)
```

Description

This is a wrapper function of NOISeq:::noiseqbio whose input class is 'eSet' and output class is Output which are not widely used. We can use as(se, "ExpressionSet") to get an eSet easily but then it will be hard to refer the treatment and control. The order of factors influence the log fold change sign. To keep it comparable to other methods the NOISeq:::readData() is used internally.

Usage

```
use NOIseq(se, colData_id, control, treatment, rank = FALSE, ...)
```

Arguments

se	Object of SummarizedExperiment class
colData_id	One of the columns of colData(se). It should be factors of more than one value.
control	Base level and one of the factor values of colData(se)[[colData_id]]
treatment	one of the factor values of colData(se)[[colData_id]]
rank	Logical value default FALSE. If true the result will have an additional column named "rank" which is ordered by 'prob' values returned by function NOISeq::noiseqbio.
...	other arguments to be passed to main function NOISeq::noiseqbio. The 'input' and 'factor' argument should not be used.

Value

A data.frame object from the results of NOISeq::noiseqbio(). For details check the documentation of 'NOISeq'

Examples

```
se <- readRDS(system.file("extdata",
  "rat_vole_mouseSE_salmon.rds",
  package = "broadSeq"))

# To reduce runtime
se <- se[rowData(se)$chromosome_name == 2,colData(se)$species == "Mouse"]

result_Noiseq <-
  use NOIseq(se = se,
             colData_id = "stage", control = "Bud", treatment = "Cap",
             rank = TRUE,
             r = 10) # r is an argument of NOISeq::noiseqbio
```

use_SAMseq

To use SummarizedExperiment with package samr

Description

To use SummarizedExperiment with package samr

Usage

```
use_SAMseq(se, colData_id, control, treatment, rank = FALSE, ...)
```

Arguments

se	Object of SummarizedExperiment class
colData_id	One of the columns of colData(se). It should be factors of more than one value.
control	Base level and one of the factor values of colData(se)[[colData_id]]
treatment	one of the factor values of colData(se)[[colData_id]]
rank	Logical value default FALSE. If true the result will have an other arguments to be passed to samr::SAMseq
...	

Value

a data.frame object as a result

volcanoPlot

Volcano plot with formatted x and y axis label.

Description

Volcano plot with formatted x and y axis label.

Usage

```
volcanoPlot(
  df,
  pValName,
  lFCName,
  sigThreshold = 0.05,
  logFCThreshold = 1,
  labelName = NULL,
  selectedLabel = NULL,
  palette = "nejm"
)
```

Arguments

df	a data.frame object
pValName	column name of df which provides p-values
lFCName	column name of df which provides log fold change values
sigThreshold	Threshold for p-values
logFCThreshold	Threshold for log fold change values
labelName	column name of df to label the dots
selectedLabel	which dots to highlight
palette	one of "npg" , "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "nejm" or similar to viridis::cividis(3)

20

volcanoPlot

Value

ggplot object

Index

assay_plot (genes_plot), 4
biplotAnyPC (prcompTidy), 8
broadSeq (broadSeq-package), 2
broadSeq-package, 2
combinedEnrichment, 3
cpm, 5
DELocal, 11
DESeqResults, 12
exactTest, 14
genes_plot, 4
GetDEResults, 13
getFeatureLoadRanking (prcompTidy), 8
ggboxplot, 4, 10
ggscatter, 7, 8
glmLRT, 14
noiseqbio, 17, 18
normalizeEdgerCPM, 5
normLibSizes, 5
normTransform, 11
pheatmap, 6
plot_MDS, 7
plotAnyPC (prcompTidy), 8
plotHeatmapCluster, 6
prcomp, 8
prcompTidy, 8, 8
results, 12
rlog, 11
round, 9
round_df, 9
sampleAssay_plot, 10
SummarizedExperiment, 4–8, 10–15, 17–19
topTags, 14
transformDESeq2, 10
use_DELocal, 11
use_deseq2, 12
use_EBSeq, 13
use_edgeR (use_edgeR_GLM), 14
use_edgeR_exact (use_edgeR_GLM), 14
use_edgeR_GLM, 14
use_limma (use_limma_trend), 15
use_limma_trend, 15
use_limma_voom (use_limma_trend), 15
use_multDE, 16
use_NOIseq, 17
use_SAMseq, 18
varianceStabilizingTransformation, 11
volcanoPlot, 19