Package ‘RegEnrich’

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Type Package

Title Gene regulator enrichment analysis

Version 1.14.0

Description This package is a pipeline to identify the key gene regulators in a biological process, for example in cell differentiation and in cell development after stimulation. There are four major steps in this pipeline: (1) differential expression analysis; (2) regulator-target network inference; (3) enrichment analysis; and (4) regulators scoring and ranking.

Depends R (>= 4.0.0), S4Vectors, dplyr, tibble, BiocSet, SummarizedExperiment

License GPL (>= 2)

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'plots.R' 'regFET.R' 'regSEA.R' 'regenrich_diffExpr.R'
'regenrich_enrich.R' 'regenrich_network.R'
'regenrich_rankScore.R' 'results.R' 'show.R' 'topNet.R'

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DeaSet-class  DeaSet class

Description

DeaSet class
dim,TopNetwork-method

dimention of 'TopNetwork' object

Description
dimention of 'TopNetwork' object

Slots

colData  DataFrame object, sample information, the row name is corresponding to the column names of expression matrix in the assays slot.

assays  SimpleList object of one/multiple matrix/matrices, this is the slot for storing the expression data after filtering (and after Variance Stabilizing Transformation, i.e. VST, if the differential analysis method is 'Wald_DESeq2' or 'LRT_DESeq2'). And the expression matrix is used for network inference and plotting.

NAMES  row names of expression data in assays slot and elementMetadata slot.

elementMetadata  feature information, contains at least a DataFrame of three columns, i.e. 'gene', 'p' and 'logFC', which stores gene names/IDs, differential p values and log2 expression fold changes, respectively.

metadata  DataFrame object, information of feature columns.

assayRaw  a slot for saving the raw expression data.

Examples

nrows = 100
cmps = 6

counts = matrix(rnbinom(nrows * ncolm, size = 2, mu = 500),
                 nrow = nrows)
assays = SimpleList(assayData = counts)

colData = DataFrame(Condition = rep(c("treatment", "ctrl"), 3),
                     row.names=LETTERS[1:6])
geneNames = sprintf("G%03s", seq(nrows))
elementMetadata = DataFrame(gene = geneNames,
                            p = numeric(nrows),
                            logFC = numeric(nrows))

ds = new("DeaSet",
         assays = Assays(assays),
         colData = colData,
         assayRaw = counts,
         elementMetadata = elementMetadata,
         NAMES = geneNames)

ds
Usage

## S4 method for signature 'TopNetwork'
dim(x)

Arguments

x a ‘TopNetwork’ object.

Value

Dimention of regulator-target network edge table.

Examples

nw = newTopNetwork()
dim(nw)

---

Enrich-class

Description

The ‘Enrich’ object is to store enrichment analysis results by either ‘FET’ method or ‘GSEA’ method.

Slots

topResult data frame. The enrichment results that pass thresholds (default threshold is 0.05).
allResult data frame. The enrichment results by FET or GSEA methods.
gene character vector indicating the genes used for enrichment analysis.
namedScores numeric vector, a vector of ranked scores (descendent), the names of the scores are the genes to perform enrichment analysis. Here the scores are p-value of each gene.
type character indicating enrichment method, either ‘FET’ or ‘GSEA’.
getResultsNames  

Description

Inference the name of results of DESeq analysis by a formula (or model matrix) and sample information

Usage

getResultsNames(design, pData = NULL)

Arguments

design  
either a formula or a model matrix.

pData  
a data frame, showing the information of each sample. If design is a formula, the pData must be include the columns that identical to the terms of the design formula. If design is a model matrix, then pData is not used. Default is NULL.

Value

the names of contrast parameter (list of character format) that regenrich_diffExpr and results function can use, and it is the same as the value that resultsNames function returns.

Examples

# formula with intercept
design = ~condition
pData = data.frame(condition = factor(c('A', 'A', 'A', 'B', 'B', 'B'),
                                        c('A', 'B')))  
getResultsNames(design, pData)

# formula without intercept
design = ~0+condition
getResultsNames(design, pData)

# formula with two terms
design = ~condition+treatment  
pData = data.frame(condition = factor(rep(c('A', 'B'), each= 4),
                                        c('A', 'B')),
                       treatment = factor(rep_len(c('Ctrl', 'Treat'), 8),
                                        c('Ctrl', 'Treat')))  
getResultsNames(design, pData)

# formula with two terms and an interaction term
design = ~condition+treatment+condition:treatment  
getResultsNames(design, pData)
# design is a model matrix
pData = data.frame(condition = factor(rep(c('A', 'B'), each= 4),
   c('A', 'B')),
   treatment = factor(rep_len(c('Ctrl', 'Treat'), 8),
   c('Ctrl', 'Treat')))

design = model.matrix(~condition+treatment, pData)
getResultsNames(design)

---

**head,Score-method**  
*head or tail of Score object*

## Description
head or tail of Score object

## Usage

```r
## S4 method for signature 'Score'
head(x, ...)

## S4 method for signature 'Score'
tail(x, ...)
```

## Arguments

- `x` an Score object.
- `...` arguments to be passed to or from other methods.

## Value
Head or tail table of Score object.

## Examples

```r
s = newScore(letters, seq(26), seq(26), seq(26), seq(2, 0, len = 26))
s1 = head(s)
s1

s2 = tail(s)
s2
```
Example RNaseq dataset [Human]

Description

Data from an RNA sequencing experiment on peripheral mononuclear blood cells (PBMC) of Lyme disease patients against healthy controls. It contains a gene expression (FPKM) table (data frame) and a sample information table (data frame).

Usage

data(Lyme_GSE63085)

Format

A list of 2 elements: FPKM and sampleInfo. FPKM is the 'Fragments Per Kilobase of transcript per Million mapped reads' data, which is a 5000 (genes) * 52 (samples) data frame. sampleInfo is the information of samples, which is 52 (samples) * 9 (features) data frame. The full version of FPKM table contains 23615 rows, which can be downloaded from GEO database.

Source

URL

References

Bouquet et al. (2016) mBio 7(1): e00100-16 (PubMed)
**Arguments**

- **assayRaw**: A matrix of gene expression data. This can be the same as the matrix-like element in assays parameter.
- **rowData**: A DataFrame object describing the rows.
- **assays**: A list or SimpleList of matrix-like element, or a matrix-like object. The matrix-like element can be the same as assayRaw parameter.
- **colData**: A DataFrame describing the sample information.
- **metadata**: An optional list of arbitrary content describing the overall experiment.

**Value**

A DeaSet object.

**Examples**

```r
# Empty DeaSet object
newDeaSet()

# 100 * 6 DeaSet object
nrows = 100
ncols = 6
counts = matrix(rnbinom(nrows * ncols, size = 2, mu = 500),
                nrow = nrows)
assays = SimpleList(counts = counts)

colData = DataFrame(Condition = rep(c("treatment", "ctrl"), 3),
                    row.names = LETTERS[1:6])
geneNames = sprintf("G%03s", seq(nrows))
elementMetadata = DataFrame(gene = geneNames,
                            p = numeric(nrows),
                            logFC = numeric(nrows))

newDeaSet(assayRaw = counts,
          rowData = elementMetadata,
          assays = SimpleList(assayData = counts),
          colData = colData)
```

---

**newTopNetwork**

*TopNetwork object creator*

**Description**

This function creates a TopNetwork object using a 3-column edge table.
plotOrders

Usage

newTopNetwork(
    networkEdgeTable,
    reg = '',
    directed = TRUE,
    networkConstruction = c("new", "COEN", "GRN"),
    percent = 100
)

Arguments

networkEdgeTable
    a data frame of 3 columns, representing 'from.gene' ('regulators'), 'to.gene'
    ('targets') and 'weight', respectively.
reg
    a vector of gene regulators.
directed
    logical, whether the network is directed. Default is TRUE.
networkConstruction
    the method to construct this network. Possible can be: 'COEN', coexpression
    network; 'GRN', gene regulatory network by random forest; 'new' (default),
    meaning a network provided by user, rather than inferred based on the expression
    data.
percent
    the percentage of edges in the original whole network. Default is 100, meaning
    100% edges in whole network.

Value

an object of topNetwork class.

Examples

data(TFs)
edge = data.frame(from = rep(TFs$TF_name[seq(3)], seq(3)),
    to = TFs$TF_name[11:16], weight = 0.1*(6:1))
object = newTopNetwork(edge, networkConstruction = 'new', percent = 100)
object
str(object)

plotOrders

Compare the orders of two vectors

Description

compare the orders of two vectors

Usage

plotOrders(name1, name2)
plotRegTarExpr

Arguments

name1 a vector with first order.
name2 a vector with another second order.

Value

A plot of comparing two orders of vectors.

Examples

```r
a = c('a1', 'a2', 'a5', 'a4')
b = c('a2', 'a5', 'a7', 'a4', 'a6')
plotOrders(a, b)
```

Description

Plot regulator and its targets expression

Usage

```r
plotRegTarExpr(
  object,
  reg,
  n = 1000,
  scale = TRUE,
  tarCol = "black",
  tarColAlpha = 0.1,
  regCol = "#ffaa00",
  xlab = "Samples",
  ylab = "Z-scores",
  ...
)
```

Arguments

object a RegenrichSet object, to which at least regenrich_diffExpr and regenrich_network functions have been applied.
reg a regulator to plot.
n the maximum number of targets to plot.
scale logical, whether gene expression is z-score normalized.
tarCol the color of the lines for the targets of the regulator.
tarColAlpha numeric, ranging from 0 to 1, indicating transparency of target lines.
plotRegTarExpr

regCol the color of the line for the 'reg'.
xlab x label of plot.
ylab y label of plot.
... other parameters in `ggplot` function.

Value

a ggplot object.

Examples

# constructing a RegenrichSet object
colData = data.frame(patientID = paste0('Sample_', seq(50)),
week = rep(c('0', '1'), each = 25),
row.names = paste0('Sample_', seq(50)),
stringsAsFactors = TRUE)
design = ~week
reduced = ~1
set.seed(123)
cnts = matrix(as.integer(rnbinom(n=1000*50, mu=100, size=1/0.1)), ncol=50,
dimnames = list(paste0('gene', seq(1000)), rownames(colData)))
cnts[5,26:50] = cnts[5,26:50] + 50L # add reads to gene5 in some samples.
id = sample(31:1000, 20) # randomly select 20 rows, and assign reads.
cnts[id,] = vapply(cnts[5,], function(x){
as.integer(rnbinom(n = 20, size = 1/0.02, mu = x))},
FUN.VALUE = rep(1L, 20))
object = RegenrichSet(expr = cnts,
colData = colData,
method = 'LRT_DESeq2', minMeanExpr = 0,
design = design, reduced = reduced, fitType = 'local',
networkConstruction = 'COEN',
enrichTest = 'FET',
reg = paste0('gene', seq(30)))

## RegEnrich analysis
object = regenrich_diffExpr(object)

# Set a random softPower, otherwise it is difficult to achieve a
# scale-free network because of a randomly generated count data.
object = regenrich_network(object, softPower = 3)
object = regenrich_enrich(object)
object = regenrich_rankScore(object)

## plot expression of a regulator and its targets.
plotRegTarExpr(object, reg = 'gene5')
plotRegTarExpr(object, reg = 'gene27')
plotSoftPower  

Plot soft power for WGCNA analysis

**Description**

Plot soft power and corresponding scale free topology fitting index to find a proper soft power for WGCNA analysis.

**Usage**

```r
plotSoftPower(
  expr,
  rowSample = FALSE,
  weights = NULL,
  powerVector = c(seq(10), seq(12, 20, by = 2)),
  RsquaredCut = 0.85,
  networkType = "unsigned",
  removeFirst = FALSE,
  nBreaks = 10,
  corFnc = WGCNA::cor,
  corOptions = list(use = "p")
)
```

**Arguments**

- `expr`: Gene expression data, either a matrix or a data frame. By default, each row represents a gene, each column represents a sample.
- `rowSample`: logic. If TRUE, each row represents a sample. The default is FALSE.
- `weights`: optional observation weights for expr to be used in correlation calculation.
- `powerVector`: a vector of soft thresholding powers for which the scale free topology fit indices are to be calculated.
- `RsquaredCut`: desired minimum scale free topology fitting index R^2. The default is 0.85.
- `networkType`: character, network type. Allowed values are (unique abbreviations of) "unsigned" (default), "signed", "signed hybrid". See adjacency.
- `removeFirst`: should the first bin be removed from the connectivity histogram? The default is FALSE.
- `nBreaks`: number of bins in connectivity histograms. The default is 10.
- `corFnc`: correlation function to be used in adjacency calculation. The default is the cor function in WGCNA.
- `corOptions`: a named list of options to the correlation function specified in corFnc. The default is list(use = "p").
Value

a list of three elements: `powerEstimate`, `fitIndices`, and `plot`. `powerEstimate` is an estimate of an appropriate soft-thresholding power. `fitIndices` is a data frame containing the fit indices for scale free topology. The plot is a ggplot object.

Examples

data(Lyme_GSE63085)
log2FPKM = log2(Lyme_GSE63085$FPKM + 1)
log2FPKMhi = log2FPKM[rowMeans(log2FPKM) >= 10^(-3), , drop = FALSE]
log2FPKMhi = head(log2FPKMhi, 3000) # First 3000 genes for example

softP = plotSoftPower(log2FPKMhi, RsquaredCut = 0.85)
print(softP)

plot_Enrich

Plot results of FET/GSEA enrichment analysis

Description

Plot FET/GSEA enrichment results. If the FET method is applied, the top `showCategory` regulator will be plotted. If the GSEA method is applied, the GSEA graph of regulator `reg` will be plotted.

Usage

plot_Enrich(object, ...)

## S4 method for signature 'RegenrichSet'
plot_Enrich(
  object,
  reg = NULL,
  showCategory = 20,
  regDescription = NULL,
  font.size = 12
)

Arguments

- **object** 
a RegenrichSet object.
- **...** 
other parameters.
- **reg** 
The regulator to plot. This only works when the GSEA enrichment method has used.
- **showCategory** 
the number of regulator to plot.
regDescription  NULL or a two-column data frame, in which first column is the regulator IDs (for example ENSEMBL IDs), and the second column is the description of regulators (for example gene name). Default is NULL, meaning both columns are the same regulator names/IDs in the network.

font.size  font size of axis labels and axis tick mark labels, default is 12.

Value

a ggplot object of plotting FET or GSEA enrichment result.

Examples

```r
# library(RegEnrich)
data("Lyme_GSE63085")data("TFs")
data = log2(Lyme_GSE63085$FPKM + 1)
colData = Lyme_GSE63085$sampleInfo

# Take first 2000 rows for example
data1 = data[seq(2000),]

design = model.matrix(~0 + patientID + week, data = colData)
object = RegenrichSet(expr = data1,
           colData = colData,
           method = "limma", minMeanExpr = 0,
           design = design,
           contrast = c(rep(0, ncol(design) - 1), 1),
           networkConstruction = "COEN",
           enrichTest = "FET")

# Differential expression analysis
object = regenrich_diffExpr(object)
# Network inference using "COEN" method
object = regenrich_network(object)
# Enrichment analysis by Fisher's exact test (FET)
object = regenrich_enrich(object)
# plot
plot_Enrich(object)

# Enrichment analysis by Fisher's exact test (FET)
object = regenrich_enrich(object, enrichTest = "GSEA")
# plot
plot_Enrich(object)
```

print.Score  

Print Score object

Description  

Print Score object
Usage

```r
## S3 method for class 'Score'
print(x, ...)
```

Arguments

- `x`: a Score object.
- `...`: optional arguments to print.

Value

`print.Score` returns the a Score object

Examples

```r
x = newScore(letters[1:5], 1:5, 1:5, -2:2, seq(2, 1, len = 5))
print(x)
```

---

reexports

Objects exported from other packages

Description

These objects are imported from other packages. Follow the links below to see their documentation.

- magrittr `%>%`

Examples

```r
# library(RegEnrich)
data("Lyme_GSE63085")
data("TFs")

data = log2(Lyme_GSE63085$FPKM + 1)
colData = Lyme_GSE63085$sampleInfo
data1 = data[seq(2000), ]
design = model.matrix(~0 + patientID + week, data = colData)

# Initializing a 'RegenrichSet' object
object = RegenrichSet(expr = data1,
                       colData = colData,
                       method = 'limma', minMeanExpr = 0,
                       design = design,
                       contrast = c(rep(0, ncol(design) - 1), 1),
                       networkConstruction = 'COEN',
                       enrichTest = 'FET')

# Using %>%
```
### Description

This is 'RegenrichSet' object creator function. There are four types of parameters in this function. First, parameters to provide raw data and sample information; `expr` and `colData`.

Second, parameters to perform differential expression analysis; 'method', 'minMeanExpr', 'design', 'reduced', 'contrast', 'coef', 'name', 'fitType', 'sfType', 'betaPrior', 'minReplicatesForReplace', 'useT', 'minmu', 'parallel', 'BPPARAM' (also for network inference), 'altHypothesis', 'listValues', 'cooksCutoff', 'independentFiltering', 'alpha', 'filter', 'theta', 'filterFun', 'addMLE', 'blind', 'ndups', 'spacing', 'block', 'correlation', 'weights', 'proportion', 'stdev.coef.lim', 'trend', 'robust', and 'winsor.tail.p'.

Third, parameters to perform regulator-target network inference; 'reg', 'networkConstruction', 'topNetPercent', 'directed', 'rowSample', 'softPower', 'networkType', 'TOMDenom', 'RsquaredCut', 'edgeThreshold', 'K', 'nbTrees', 'importanceMeasure', 'trace', 'BPPARAM' (also for differential expression analysis), and 'minR'.

Fourth, parameters to perform enrichment analysis: 'enrichTest', 'namedScoresCutoffs', 'minSize', 'maxSize', 'pvalueCutoff', 'qvalueCutoff', 'regAltName', 'universe', and 'nperm'.

### Usage

```r
RegenrichSet(
  expr,
  colData,
  rowData = NULL,
  method = c("Wald_DESeq2", "LRT_DESeq2", "limma", "LRT_LM"),
  minMeanExpr = NULL,
  design,
  reduced,
  contrast,
  coef = NULL,
  name,
  fitType = c("parametric", "local", "mean"),
  sfType = c("ratio", "poscounts", "iterate"),
  betaPrior,
)```
minReplicatesForReplace = 7,
useT = FALSE,
minmu = 0.5,
parallel = FALSE,
BPPARAM = bpparam(),
altHypothesis = c("greaterAbs", "lessAbs", "greater", "less"),
listValues = c(1, -1),
cooksCutoff,
independentFiltering = TRUE,
alpha = 0.1,
filter,
theta,
filterFun,
addMLE = FALSE,
blind = FALSE,
ndups = 1,
spacing = 1,
block = NULL,
correlation,
weights = NULL,
proportion = 0.01,
stddev.coef.lim = c(0.1, 4),
trend = FALSE,
robust = FALSE,
winsor.tail.p = c(0.05, 0.1),
reg = TFs$TF_name,
networkConstruction = c("COEN", "GRN", "new"),
topNetPercent = 5,
directed = FALSE,
rowSample = FALSE,
softPower = NULL,
networkType = "unsigned",
TOMDenom = "min",
RsquaredCut = 0.85,
edgeThreshold = NULL,
K = "sqrt",
nbTrees = 1000,
importanceMeasure = "IncNodePurity",
trace = FALSE,
minR = 0.3,
enrichTest = c("FET", "GSEA"),
namedScoresCutoffs = 0.05,
minSize = 5,
maxSize = 5000,
pvalueCutoff = 0.05,
qvalueCutoff = 0.2,
regAltName = NULL,
universe = NULL,
Arguments

**expr**
matrix or data.frame, expression profile of a set of genes or a set of proteins. If the method = 'Wald_DESeq2' or 'LRT_DESeq2' only non-negative integer matrix (read counts by RNA sequencing) is accepted.

**colData**
data frame, sample phenotype data. The rows of colData must correspond to the columns of expr.

**rowData**
NULL or data frame, information of each row/gene. Default is NULL, which will generate a DataFrame of three columns, i.e., "gene", "p", and "logFC".

**method**
either 'Wald_DESeq2', 'LRT_DESeq2', 'limma', or 'LRT_LM' for the differential expression analysis.

- When method = 'Wald_DESeq2', the Wald test in DESeq2 package is used;
- When method = 'LRT_DESeq2', the likelihood ratio test (LRT) in DESeq2 package is used;
- When method = 'limma', the 'ls' method and empirical Bayes method in limma package are used to calculate moderated t-statistics and differential p-values;
- When method = 'LRT_LM', a likelihood ratio test is performed for each row of 'expr' to compare two linear model specified by 'design' and 'reduced' arguments. In this case, the fold changes are not calculated but set to 0.

**minMeanExpr**
numeric, the cutoff of gene average expression for pre-filtering. The rows of 'expr' with average expression < minMeanExpr is removed. The higher 'minMeanExpr' is, the more genes are not included for testing.

**design**
either model formula or model matrix. For method = 'LRT_DESeq2' or 'LRT_LM', the design is the full model formula/matrix. For method = 'limma', and if design is a formula, the model matrix is constructed using model.matrix(design, colData), so the name of each term in the design formula must be included in the column names of 'colData'.

**reduced**
The argument is used only when method = 'LRT_DESeq2' or 'LRT_LM', it is a reduced formula/matrix to compare against. If the design is a model matrix, 'reduced' must also be a model matrix.

**contrast**
The argument is used only when method = 'LRT_DESeq2', 'Wald_DESeq2', or 'limma'. When method = 'LRT_DESeq2', or 'Wald_DESeq2', it specifies what comparison to extract from the 'DESeqDataSet' object to build a results table (when method = 'LRT_DESeq2', this does not affect the value of 'stat', 'pvalue', or 'padj'). It can be one of following three formats:

- a character vector with exactly three elements: the name of a factor in the design formula, the name of the numerator level for the fold change, and the name of the denominator level for the fold change;
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• a list of 1 or 2 character vector(s): the first element specifies the names of the fold changes for the numerator, and the second element (optional) specifies the names of the fold changes for the denominator. These names should be elements of \texttt{getResultsNames(design, colData)};

• a numeric contrast vector with one element for each element in \texttt{getResultsNames(design, colData)}.

When method = 'limma', It can be one of following two formats:

• a numeric matrix with rows corresponding to coefficients in design matrix and columns containing contrasts;

• a numeric vector if there is only one contrast. Each element of the vector corresponds to coefficients in design matrix. This is similar to the third format of contrast when method = 'LRT\_DESeq2', or 'Wald\_DESeq2'.

\texttt{coef} The argument is used only when method = 'limma'. (Vector of) column number or column name specifying which coefficient or contrast of the linear model is of interest. Default is NULL.

\texttt{name} The argument is used only when method = 'LRT\_DESeq2' or 'Wald\_DESeq2'. the name of the individual effect (coefficient) for building a results table. Use this argument rather than contrast for continuous variables, individual effects or for individual interaction terms. The value provided to name must be an element of \texttt{getResultsNames(design, colData)}.

\texttt{fitType} either 'parametric', 'local', or 'mean' for the type of fitting of dispersions to the mean intensity. This argument is used only when method = 'Wald\_DESeq2' or 'LRT\_DESeq2'. See \texttt{DESeq} from DESeq2 package for more details. Default is 'parametric'.

\texttt{sfType} either 'ratio', 'poscounts', or 'iterate' for the type of size factor estimation. This argument is used only when method = either 'Wald\_DESeq2' or 'LRT\_DESeq2'. See \texttt{DESeq} from DESeq2 package for more details. Default is 'ratio'.

\texttt{betaPrior} This argument is used only when method = either 'Wald\_DESeq2' or 'LRT\_DESeq2'. See \texttt{DESeq} from DESeq2 package for more details.

\texttt{minReplicatesForReplace} This argument is used only when method = either 'Wald\_DESeq2' or 'LRT\_DESeq2'. See \texttt{DESeq} from DESeq2 package for more details. Default is 7.

\texttt{useT} This argument is used only when method = either 'Wald\_DESeq2' or 'LRT\_DESeq2'. See \texttt{DESeq} from DESeq2 package for more details. Default is FALSE.

\texttt{minmu} This argument is used only when method = either 'Wald\_DESeq2' or 'LRT\_DESeq2'. See \texttt{DESeq} from DESeq2 package for more details. Default is 0.5.

\texttt{parallel} whether computing (only for differential analysis with method = "Wald\_DESeq2" or "LRT\_DESeq2") is parallel (default is FALSE).

\texttt{BPPARAM} parameters for parallel computing (default is \texttt{bpparam()}).

\texttt{altHypothesis} = c('greaterAbs', 'lessAbs', 'greater', 'less'). This argument is used only when method = either 'Wald\_DESeq2' or 'LRT\_DESeq2'. See \texttt{results} from DESeq2 package for more details. Default is 'greaterAbs'.

\texttt{listValues} This argument is used only when method = either 'Wald\_DESeq2' or 'LRT\_DESeq2'. See \texttt{results} from DESeq2 package for more details. Default is c(1, -1),
cooksCutoff threshold on Cook’s distance, such that if one or more samples for a row have a distance higher, the p-value for the row is set to NA. This argument is used only when method = either 'Wald_DESeq2' or 'LRT_DESeq2'. See results from DESeq2 package for more details.

independentFiltering logical, whether independent filtering should be applied automatically. This argument is used only when method = either 'Wald_DESeq2' or 'LRT_DESeq2'. See results from DESeq2 package for more details. Default is TRUE.

alpha the significance cutoff used for optimizing the independent filtering. This argument is used only when method = either 'Wald_DESeq2' or 'LRT_DESeq2'. See results from DESeq2 package for more details. Default is 0.1.

filter the vector of filter statistics over which the independent filtering is optimized. By default the mean of normalized counts is used. This argument is used only when method = either 'Wald_DESeq2' or 'LRT_DESeq2'. See results from DESeq2 package for more details.

theta the quantiles at which to assess the number of rejections from independent filtering. This argument is used only when method = either 'Wald_DESeq2' or 'LRT_DESeq2'. See results from DESeq2 package for more details.

filterFun an optional custom function for performing independent filtering and p-value adjustment. This argument is used only when method = either 'Wald_DESeq2' or 'LRT_DESeq2'. See results from DESeq2 package for more details.

addMLE if betaPrior=TRUE was used, whether the 'unshrunken' maximum likelihood estimates (MLE) of log2 fold change should be added as a column to the results table. This argument is used only when method = either 'Wald_DESeq2' or 'LRT_DESeq2'. See results from DESeq2 package for more details. Default is FALSE.

blind logical, whether to blind the transformation to the experimental design. This argument is used only when method = either 'Wald_DESeq2' or 'LRT_DESeq2'. See vst from DESeq2 package for more details. Default is FALSE, which is different from the default of vst function.

ndups positive integer giving the number of times each distinct probe is printed on each array. This argument is used only when method = 'limma'. See lmFit from limma package for more details. Default is 1.

spacing positive integer giving the spacing between duplicate occurrences of the same probe, spacing=1 for consecutive rows. This argument is used only when method = 'limma'. See lmFit from limma package for more details. Default is 1.

block vector or factor specifying a blocking variable on the arrays. Has length equal to the number of arrays. Must be NULL if ndups > 2. This argument is used only when method = 'limma'. See lmFit from limma package for more details. Default is NULL.

correlation the inter-duplicate or inter-technical replicate correlation. The correlation value should be estimated using the duplicateCorrelation function. This argument is used only when method = 'limma'. See lmFit from limma package for more details.
weights non-negative precision weights. Can be a numeric matrix of individual weights of same size as the object expression matrix, or a numeric vector of array weights with length equal to ncol of the expression matrix, or a numeric vector of gene weights with length equal to nrow of the expression matrix. This argument is used only when method = 'limma' or 'LRT_LM'. See lmFit from limma package for more details. Default is NULL.

proportion numeric value between 0 and 1, assumed proportion of genes which are differentially expressed. This argument is used only when method = 'limma'. See eBayes from limma package for more details. Default is 0.01.

stdev.coef.lim numeric vector of length 2, assumed lower and upper limits for the standard deviation of log2-fold-changes for differentially expressed genes. This argument is used only when method = 'limma'. See eBayes from limma package for more details. Default is c(0.1, 4).

trend logical, should an intensity-trend be allowed for the prior variance? This argument is used only when method = 'limma'. See eBayes from limma package for more details. Default is FALSE, meaning that the prior variance is constant.

robust logical, should the estimation of df.prior and var.prior be robustified against outlier sample variances? This argument is used only when method = 'limma'. See eBayes from limma package for more details. Default is FALSE.

winsor.tail.p numeric vector of length 1 or 2, giving left and right tail proportions of x to Winsorize. Used only when method = 'limma' and robust=TRUE. See eBayes from limma package for more details. Default is c(0.05,0.1)

reg a vector of regulator names (ID). By default, these are transcription (co-)factors defined by three literatures/databases, namely RegNet, TRRUST, and Marbach2016. The type (for example ENSEMBL gene ID, Entrez gene ID, or gene sym- ble/name) of names or IDs of these regulators must be the same as the type of names or IDs in the regulator-target network.

networkConstruction the method to construct this network. Possible can be: 'COEN', coexpression network; 'GRN', gene regulatory network by random forest; 'new' (default), meaning a network provided by user, rather than inferred based on the expression data.

topNetPercent numeric, what percentage of the top edges in the full network is retained. Default is 5, meaning top 5% of edges. This value must be between 0 and 100.

directed logical, whether the network is directed. Default is FALSE.

rowSample logic, if TRUE, each row represents a sample. Otherwise, each column represents a sample. Default is FALSE.

softPower numeric, a soft power to achieve scale free topology. If not provided, the parameter will be picked automatically by plotSoftPower function.

networkType network type. Allowed values are (unique abbreviations of) 'unsigned' (default), 'signed', 'signed hybrid'. See adjacency.

TOMDenom a character string specifying the TOM variant to be used. Recognized values are 'min' giving the standard TOM described in Zhang and Horvath (2005),
and 'mean' in which the min function in the denominator is replaced by mean. The 'mean' may produce better results but at this time should be considered experimental.

RsquaredCut desired minimum scale free topology fitting index $R^2$. Default is 0.85.

edgeThreshold numeric, the threshold to remove the low weighted edges. Default is NULL, which means no edges will be removed.

K integer or character. The number of features in each tree, can be either a integer number, 'sqrt', or 'all'. 'sqrt' denotes $\sqrt{\text{the number of 'reg'}}, 'all' means the number of 'reg'. Default is 'sqrt'.

nbTrees integer. The number of trees. Default is 1000.

importanceMeasure character. importanceMeasure can be '%IncMSE' or 'IncNodePurity', corresponding to type = 1 and 2 in importance function, respectively. Default is 'IncNodePurity'(decrease in node impurity), which is faster than '%IncMSE' (decrease in accuracy).

trace logical. To show the progress or not (default).

minR numeric. The minimum correlation coefficient of prediction is to control model accuracy. Default is 0.3.

enrichTest character, specifying the enrichment analysis method, which is either 'FET' (Fisher's exact test) or 'GSEA' (gene set enrichment analysis).

namedScoresCutoffs numeric, the significance cutoff for the differential analysis p value. Default is 0.05.

minSize The minimum number (default 5) of target genes.

maxSize The maximum number (default 5000) of target genes.

pvalueCutoff numeric, the significance cutoff for adjusted enrichment p value. This is used for obtaining the 'topResult' slot in the final 'Enrich' object. Default is 0.05.

qvalueCutoff numeric, the significance cutoff of enrichment q-value. Default is 0.2.

gerAltName alternative name for regulator. Default is NULL.

universe a vector of charactors. Background target genes.

nperm integer, number of permutations. The minimal possible nominal p-value is about $1/nperm$. Default is 10000.

Value an object of RegenrichSet class.

Examples

# library(RegEnrich)
data("Lyme_GSE63085")
data("TFs")

data = log2(Lyme_GSE63085$FPKM + 1)
colData = Lyme_GSE63085$sampleInfo
# Take first 2000 rows for example
data1 = data[seq(2000),]
design = model.matrix(~0 + patientID + week, data = colData)

# Initializing a 'RegenrichSet' object
object = RegenrichSet(expr = data1,
                      colData = colData,
                      method = 'limma',
                      minMeanExpr = 0,
                      design = design,
                      contrast = c(rep(0, ncol(design) - 1), 1),
                      networkConstruction = 'COEN',
                      enrichTest = 'FET')

---

**RegenrichSet-class**  
*RegenrichSet class*

**Description**

The RegenrichSet is the fundamental class that RegEnrich package is working with.

**Slots**

- **assayRaw** matrix, the initial raw expression data.
- **colData** DataFrame object, indicating sample information. Each row represent a sample and each column represent a feature of samples.
- **assays** SimpleList object, containing the expression data after filtering (and after Variance Stabilizing Transformation, i.e. VST, if the differential analysis method is 'Wald\_DESeq2' or 'LRT\_DESeq2').
- **elementMetadata** DataFrame object, a slot for saving results by differential expression analysis, containing at least three columns: 'gene', 'p' and 'logFC'.
- **topNetwork** TopNetwork object, a slot for saving top network edges. After regulator-target network inference, a TopNetwork-class object is assigned to this slot, containing only top ranked edges in the full network. Default is NULL.
- **resEnrich** Enrich object, a slot for saving enrichment analysis either by Fisher's exact test (FET) or gene set enrichment analysis (GSEA).
- **resScore** Score object, a slot for saving regulator ranking results. It contains five components, which are 'reg' (regulator), 'negLogPDEA' (-log10(p values of differential expression analysis)), 'negLogPEnrich' (-log10(p values of enrichment analysis)), 'logFC' (log2 fold changes), and 'score' (RegEnrich ranking score).
- **paramsIn** list. The parameters used in the whole RegEnrich analysis. This slot can be updated by respecifying arguments in each step of RegEnrich analysis.
paramsOut: a list of four elements: DeaMethod (differential expression method), networkType (regulator-target network construction method), percent (what percentage of edges from the full network is used), and enrichTest (enrichment method). By default, each element is NULL.

network: TopNetwork object, a slot for saving a full network.

---

**regenrich_diffExpr**  
**Differential expression analysis step**

**Description**

This is the first step of RegEnrich analysis. Differential expression analysis by this function needs to be performed on a 'RegenrichSet' object.

**Usage**

```r
regenrich_diffExpr(object, ...) # S4 method for signature 'RegenrichSet'
regenrich_diffExpr(object, ...)```

**Arguments**

- object: a 'RegenrichSet' object, which is initialized by `RegenrichSet` function.
- ...: arguments for differential analysis. After constructing a 'RegenrichSet' object, all arguments for RegEnrich analysis have been initialized and stored in 'paramsIn' slot, while the arguments for differential analysis can be re-specified here.

These arguments include 'method', 'minMeanExpr', 'design', 'reduced', 'contrast', 'coef', 'name', 'fitType', 'sfType', 'betaPrior', 'minReplicatesForReplace', 'useT', 'minmu', 'parallel', 'BPPARAM', 'altHypothesis', 'listValues', 'cooksCutoff', 'independentFiltering', 'alpha', 'filter', 'theta', 'filterFun', 'addMLE', 'blind', 'ndups', 'spacing', 'block', 'correlation', 'weights', 'proportion', 'stdev.coef.lim', 'trend', 'robust', and 'winsor.tail.p'.

See `RegenrichSet` function for more details about these arguments.

**Value**

This function returns a 'RegenrichSet' object with an updated 'resDEA' slot, which is a 'DeaSet' object, and an updated 'paramsIn' slot. See `newDeaSet` function for more details about 'DeaSet' class. If an argument not in the above list is specified in the regenrich_diffExpr function, a warning or error will be raised.

**See Also**

Initialization of a 'RegenrichSet' object `RegenrichSet`, and next step `regenrich_network`. 
Examples

```r
# library(RegEnrich)
data("Lyme_GSE63085")
data("TFs")

data = log2(Lyme_GSE63085$FPKM + 1)
colData = Lyme_GSE63085$sampleInfo

# Take first 2000 rows for example
data1 = data[seq(2000), ]
design = model.matrix(~0 + patientID + week, data = colData)

# Initializing a 'RegenrichSet' object
object = RegenrichSet(expr = data1,
colData = colData,
method = 'limma', minMeanExpr = 0,
design = design,
contrast = c(rep(0, ncol(design) - 1), 1),
networkConstruction = 'COEN',
enrichTest = 'FET')

# Using the predefined parameters in the previous step
(object = regenrich_diffExpr(object))

# re-specifying parameter 'minMeanExpr'
print(slot(object, 'paramsIn')$minMeanExpr)
(object = regenrich_diffExpr(object, minMeanExpr = 1))
print(slot(object, 'paramsIn')$minMeanExpr)

# Unrecognized argument 'unrecognizedArg' (Error)
# object = regenrich_diffExpr(object, minMeanExpr = 1,
# unrecognizedArg = 23)

# Argument not for differential expression analysis (Warning)
# print(slot(object, 'paramsIn')$networkConstruction)
# (object = regenrich_diffExpr(object, minMeanExpr = 1,
# networkConstruction = 'GRN'))
# print(slot(object, 'paramsIn')$networkConstruction) # not changed
```

---

### Description

As the third step of RegEnrich analysis, enrichment analysis is followed by differential expression analysis (regenrich_diffExpr), and regulator-target network inference (regenrich_network).
Usage

regenrich_enrich(object, ...)

## S4 method for signature 'RegenrichSet'
regenrich_enrich(object, ...)

Arguments

- **object**: a `RegenrichSet` object, to which `regenrich_diffExpr`, and `regenrich_network`, functions have been already applied.
- **...**: arguments for enrichment analysis. After constructing a `RegenrichSet` object using `RegenrichSet` function, all arguments for RegEnrich analysis have been initialized and stored in `paramsIn` slot. The arguments for enrichment analysis can be re-specified here.

These arguments include 'enrichTest', 'namedScoresCutoffs', 'minSize', 'maxSize', 'pvalueCutoff', 'qvalueCutoff', 'regAltName', 'universe', 'minSize', ' maxSize', 'pvalueCutoff', and 'nperm'.

See `RegenrichSet` function for more details about these arguments.

Value

This function returns a `RegenrichSet` object with an updated `resEnrich` slots, which is `Enrich` objects, and an updated `paramsIn` slot. See `Enrich-class` function for more details about `Enrich` class.

See Also

Previous step `regenrich_network`, and next step `regenrich_rankScore`.

Examples

```r
# library(RegEnrich)
data("Lyme_GSE63085")
data("TFs")
data = log2(Lyme_GSE63085$FPKM + 1)
colData = Lyme_GSE63085$sampleInfo

# Take first 2000 rows for example
data1 = data[seq(2000), ]
design = model.matrix(~0 + patientID + week, data = colData)

# Initializing a 'RegenrichSet' object
object = RegenrichSet(expr = data1, colData = colData, method = 'limma', minMeanExpr = 0, design = design)
```
# Differential expression analysis
object = regenrich_diffExpr(object)

# Network inference using 'COEN' method
object = regenrich_network(object)

# Enrichment analysis by Fisher's exact test (FET)
(object = regenrich_enrich(object))

# Enrichment analysis by Fisher's exact test (GSEA)
(object = regenrich_enrich(object, enrichTest = "GSEA"))

---

**Description**

As the second step of RegEnrich analysis, network inference is followed by differential expression analysis (regenrich_diffExpr).

Provide a network to ‘RegenrichSet’ object.

**Usage**

regenrich_network(object, ...)

```r
## S4 method for signature 'RegenrichSet'
regenrich_network(object, ...)
```

```r
regenrich_network(object) <- value
## S4 replacement method for signature 'RegenrichSet,TopNetwork'
regenrich_network(object) <- value
```

```r
## S4 replacement method for signature 'RegenrichSet,data.frame'
regenrich_network(object) <- value
```

**Arguments**

- `object` a ‘RegenrichSet’ object, to which `regenrich_diffExpr` function has been already applied.
arguments for network inference. After constructing a ‘RegenrichSet’ object using `RegenrichSet` function, all arguments for RegEnrich analysis have been initialized and stored in ‘paramsIn’ slot. The arguments for network inference can be re-specified here.


See `RegenrichSet` function for more details about these arguments.

Value

This function returns a ‘RegenrichSet’ object with an updated ‘network’ and ‘topNetP’ slots, which are ‘TopNetwork’ objects, and an updated ‘paramsIn’ slot. See `TopNetwork-class` class for more details.

See Also

Previous step `regenrich_diffExpr`, and next step `regenrich_enrich`. User defined network `regenrich_network<-`

Examples

```r
# library(RegEnrich)
data("Lyme_GSE63085")
data("TFs")

data = log2(Lyme_GSE63085$FPKM + 1)
colData = Lyme_GSE63085$sampleInfo

# Take first 2000 rows for example
data1 = data[seq(2000), ]

design = model.matrix(~0 + patientID + week, data = colData)

# Initializing a 'RegenrichSet' object
object = RegenrichSet(expr = data1,
    colData = colData,
    method = 'limma', minMeanExpr = 0,
    design = design,
    contrast = c(rep(0, ncol(design) - 1), 1),
    networkConstruction = 'COEN',
    enrichTest = 'FET')
```
# Differential expression analysis
(object = regenrich_diffExpr(object))

# Network inference using 'COEN' method
(object = regenrich_network(object))

---

## Description

As the fourth step of RegEnrich analysis, regulator ranking is followed by differential expression analysis (regenrich_diffExpr), regulator-target network inference (regenrich_network), and enrichment analysis (regenrich_enrich).

## Usage

```r
regenrich_rankScore(object)
```

## Arguments

- `object` a `RegenrichSet` object, to which `regenrich_diffExpr`, `regenrich_network`, and `regenrich_enrich` functions all have been already applied.

## Value

This function returns a `RegenrichSet` object with an updated `resScore` slots, which is a `regEnrichScore` (also `data.frame`) object, and an updated `paramsIn` slot. In the `regEnrichScore` object there are five columns, which are `reg` (regulator), `negLogPDEA` (-log10(p values of differential expression analysis)), `negLogPEnrich` (-log10(p values of enrichment analysis)), `logFC` (log2 fold changes), and `score` (RegEnrich ranking score).

## See Also

Previous step `regenrich_enrich`.

## Examples

```r
# library(RegEnrich)
data("Lyme_GSE63085")
data("TFs")

data = log2(Lyme_GSE63085$FPKM + 1)
```
colData = Lyme_GSE63085$sampleInfo

# Take first 2000 rows for example
data1 = data[seq(2000),]

design = model.matrix(~0 + patientID + week, data = colData)

# Initializing a 'RegenrichSet' object
object = RegenrichSet(expr = data1,  
                       colData = colData,  
                       method = 'limma', minMeanExpr = 0,  
                       design = design,  
                       contrast = c(rep(0, ncol(design) - 1), 1),  
                       networkConstruction = 'COEN',  
                       enrichTest = 'FET')

# Differential expression analysis
object = regenrich_diffExpr(object)

# Network inference using 'COEN' method
object = regenrich_network(object)

# Enrichment analysis by Fisher's exact test (FET)
object = regenrich_enrich(object)

# Regulators ranking
(object = regenrich_rankScore(object))

---

**results_expr**

**Result accessor functions**

**Description**

- results_expr accesses raw expression data.
- results_DEA accesses results from differential expression analysis.
- results_topNet accesses results from network inference.
- results_enrich accesses results from FET/GSEA enrichment analysis.
- results_score accesses results from regulator scoring and ranking.

**Usage**

```r
results_expr(object)
results_DEA(object)
results_topNet(object)
```
results_expr

results_enrich(object)

results_score(object)

Arguments

object RegenrichSet object.

Value

results_expr returns an expression matrix.
results_DEA returns a list result of differential analysis.
results_topNet returns a TopNetwork object.
results_enrich returns an Enrich object by either FET or GSEA method.
results_score returns a data frame of summarized ranking scores of regulators.

Examples

# library(RegEnrich)
data("Lyme_GSE63085")
data("TFs")

data = log2(Lyme_GSE63085$FPKM + 1)
colData = Lyme_GSE63085$sampleInfo

# Take first 2000 rows for example
data1 = data[seq(2000), ]
design = model.matrix(~0 + patientID + week, data = colData)

# Initializing a 'RegenrichSet' object
object = RegenrichSet(expr = data1,
colData = colData,
method = 'limma', minMeanExpr = 0,
design = design,
contrast = c(rep(0, ncol(design) - 1), 1),
networkConstruction = 'COEN',
enrichTest = 'FET')

# Differential expression analysis
object = regenrich_diffExpr(object)
results_expr(object)
results_DEA(object)

# Network inference using 'COEN' method
object = regenrich_network(object)
results_topNet(object)

# Enrichment analysis by Fisher's exact test (FET)
Score-class

object = regenrich_enrich(object)
results_enrich(object)

# Regulators ranking
object = regenrich_rankScore(object)
results_score(object)

---

Score-class

Score class

Description

‘Score’ class inherits tibble (“tbl”). The objects of ‘Score’ class are to store information of regulator ranking scores.

Usage

newScore(
  reg = character(),
  negLogPDEA = numeric(),
  negLogPEnrich = numeric(),
  logFC = numeric(),
  score = numeric()
)

Arguments

reg character, regulator IDs.

negLogPDEA numeric, -log(p_DEA).

negLogPEnrich numeric, -log(p_Enrich).

logFC numeric, log2 fold change.

score numeric, RegEnrich ranking score.

Value

newScore function returns a Score object.

Slots

names character vector, containing "reg", "negLogPDEA", "negLogPEnrich", "logFC", and "score".

.Data a list of length 5, each elements corresponds to the names slots.

.row.names character, regulators corresponding to .Data slot.

.S3Class character vector, containing "tbl_dfc", "tbl", "data.frame", indicating the classes that ‘Score’ class inherits.
**Examples**

```r
ewScore()
newScore(letters[1:5], 1:5, 1:5, -2:2, seq(2, 1, len = 5))
```

---

**Description**

methods of generic function "show"

**Usage**

```r
## S4 method for signature 'DeaSet'
show(object)

## S4 method for signature 'TopNetwork'
show(object)

## S4 method for signature 'Enrich'
show(object)

## S4 method for signature 'Score'
show(object)

## S4 method for signature 'RegenrichSet'
show(object)
```

**Arguments**

- `object` one object of either DeaSet, TopNetwork, Enrich, Score, or RegenrichSet class.

**Value**

show returns an invisible original object.

**Examples**

```r
x = newScore(letters[1:5], 1:5, 1:5, -2:2, seq(2, 1, len = 5))
show(x)
```
**Description**

The transcription factors and co-factors in humans are considered the regulators in RegEnrich. And these regulators are obtained from (Han et al. 2015; Marbach et al. 2016; and Liu et al. 2015).

**Usage**

data(TFs)

**Format**

An object of 2-column data.frame; The first column is ENSEMBL ID of gene regulators. The second column is gene name of gene regulators. The row name of this data frame is identical to the ENSEMBL ID column.

**References**


---

**TopNetwork-class**

**Description**

The 'TopNetwork' object is to store either a full network (the percentage of top edges is 100 between 0 to 10).

**Slots**

element tibble, the pool of targets in the network.
set tibble, the pool of valid regulators.

elementset tibble, regulator-target edges with edge weights. and the elements are regulators of the targets indicated by the element name.

directed logical, whether the network is directed.

networkConstruction character, by which method this network is constructed. Either 'COEN' (coexpression network using WGCNA), or 'GRN' (gene regulatory network using random forest), or 'new' (a network provided by the user).

percent numeric, what percentage of the top edges are remained. The value must be between 0 (excluding) and 100 (including).

active character, which data table is activated, the default is "elementset".
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