# Package 'fgsea'

April 5, 2020

```
Description The package implements an algorithm for fast gene set enrichment
      analysis. Using the fast algorithm allows to make more permutations and get
      more fine grained p-values, which allows to use accurate stantard approaches to
     multiple hypothesis correction.
biocViews GeneExpression, DifferentialExpression, GeneSetEnrichment,
     Pathways
SystemRequirements C++11
Depends R (>= 3.3), Rcpp
Imports data.table, BiocParallel, stats, ggplot2 (>= 2.2.0),
      gridExtra, grid, fastmatch, Matrix, utils
Suggests testthat, knitr, rmarkdown, reactome.db, AnnotationDbi,
      parallel, org.Mm.eg.db, limma, GEOquery
License MIT + file LICENCE
LazyData true
LinkingTo Rcpp, BH
RoxygenNote 6.1.1
Encoding UTF-8
VignetteBuilder knitr
URL https://github.com/ctlab/fgsea/
BugReports https://github.com/ctlab/fgsea/issues
git_url https://git.bioconductor.org/packages/fgsea
git_branch RELEASE_3_10
git_last_commit 22c00b6
git_last_commit_date 2019-10-29
Date/Publication 2020-04-04
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```

Title Fast Gene Set Enrichment Analysis

**Version** 1.12.0 **Date** 2019-10-25

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# **Description**

Takes  $O(k \log k)$  time, where k is a size of 'selectedSize'.

# Usage

```
calcGseaStat(stats, selectedStats, gseaParam = 1,
  returnAllExtremes = FALSE, returnLeadingEdge = FALSE)
```

# **Arguments**

stats Named numeric vector with gene-level statistics sorted in decreasing order (or-

der is not checked).

selectedStats Indexes of selected genes in the 'stats' array.

gseaParam GSEA weight parameter (0 is unweighted, suggested value is 1).

returnAllExtremes

If TRUE return not only the most extreme point, but all of them. Can be used

for enrichment plot

returnLeadingEdge

If TRUE return also leading edge genes.

# Value

Value of GSEA statistic if both returnAllExtremes and returnLeadingEdge are FALSE. Otherwise returns list with the following elements:

- res value of GSEA statistic
- tops vector of top peak values of cumulative enrichment statistic for each gene;

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- bottoms vector of bottom peak values of cumulative enrichment statistic for each gene;
- leadingGene vector with indexes of leading edge genes that drive the enrichment, see http: //software.broadinstitute.org/gsea/doc/GSEAUserGuideTEXT.htm#\_Running\_a\_Leading.

## **Examples**

```
data(exampleRanks)
data(examplePathways)
ranks <- sort(exampleRanks, decreasing=TRUE)
es <- calcGseaStat(ranks, na.omit(match(examplePathways[[1]], names(ranks))))</pre>
```

calcGseaStatBatchCpp

Calculates GSEA statistic valus for all gene sets in 'selectedStats' list.

# Description

Takes O(n + mKlogK) time, where n is the number of genes, m is the number of gene sets, and k is the mean gene set size.

# Usage

```
calcGseaStatBatchCpp(stats, selectedGenes, geneRanks)
```

#### **Arguments**

stats Numeric vector of gene-level statistics sorted in decreasing order

 $selected Genes \quad List \ of \ integer \ vector \ with \ integer \ gene \ IDs \ (from \ 1 \ to \ n)$ 

geneRanks Integer vector of gene ranks

# Value

Numeric vector of GSEA statistics of the same length as 'selectedGenes' list

collapse Pathways

Collapse list of enriched pathways to independent ones.

## **Description**

Collapse list of enriched pathways to independent ones.

# Usage

```
collapsePathways(fgseaRes, pathways, stats, pval.threshold = 0.05,
   nperm = 10/pval.threshold, gseaParam = 1)
```

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#### **Arguments**

fgseaRes Table with results of running fgsea(), should be filtered by p-value, for example

by selecting ones with padj < 0.01.

pathways List of pathways, should contain all the pathways present in 'fgseaRes'.

stats Gene-level statistic values used for ranking, the same as in 'fgsea()'.

pval.threshold Two pathways are considered dependent when p-value of enrichment of one

pathways on background of another is greater then 'pval.threshold'.

nperm Number of permutations to test for independence, should be several times greater

than '1/pval.threhold'. Default value: '10/pval.threshold'.

gseaParam GSEA parameter, same as for 'fgsea()'

#### Value

Named list with two elments: 'mainPathways' containing IDs of pathways not reducable to each other, and 'parentPathways' with vector describing for all the pathways to which ones they can be reduced. For pathways from 'mainPathways' vector 'parentPathways' contains 'NA' values.

# **Examples**

examplePathways Example list of mouse Reactome pathways.

#### **Description**

The list was obtained by selecting all the pathways from 'reactome.db' package that contain mouse genes. The exact script is available as system.file("gen\_reactome\_pathways.R", package="fgsea")

exampleRanks Example vector of gene-level statistics obtained for Th1 polarization.

# **Description**

The data were obtained by doing differential expression between Naive and Th1-activated states for GEO dataset GSE14308. The exact script is available as system.file("gen\_gene\_ranks.R", package="fgsea")

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fgsea	Runs preranked gene set enrichment analysis.
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# **Description**

The function takes about  $O(nk^{3/2})$  time, where n is number of permutations and k is a maximal size of the pathways. That means that setting 'maxSize' parameter with a value of ~500 is strongly recommended.

# Usage

```
fgsea(pathways, stats, nperm, minSize = 1, maxSize = Inf, nproc = 0,
  gseaParam = 1, BPPARAM = NULL)
```

# **Arguments**

pathways	List of gene sets to check.
stats	Named vector of gene-level stats. Names should be the same as in 'pathways'
nperm	Number of permutations to do. Minimial possible nominal p-value is about 1/nperm
minSize	Minimal size of a gene set to test. All pathways below the threshold are excluded.
maxSize	Maximal size of a gene set to test. All pathways above the threshold are excluded.
nproc	If not equal to zero sets BPPARAM to use nproc workers (default = 0).
gseaParam	GSEA parameter value, all gene-level statis are raised to the power of 'gsea-Param' before calculation of GSEA enrichment scores.
BPPARAM	Parallelization parameter used in bplapply. Can be used to specify cluster to run. If not initialized explicitly or by setting 'nproc' default value 'bpparam()' is used.

# Value

A table with GSEA results. Each row corresponds to a tested pathway. The columns are the following:

- pathway name of the pathway as in 'names(pathway)';
- pval an enrichment p-value;
- padj a BH-adjusted p-value;
- ES enrichment score, same as in Broad GSEA implementation;
- NES enrichment score normalized to mean enrichment of random samples of the same size;
- nMoreExtreme' a number of times a random gene set had a more extreme enrichment score value;
- size size of the pathway after removing genes not present in 'names(stats)'.
- leadingEdge vector with indexes of leading edge genes that drive the enrichment, see http: //software.broadinstitute.org/gsea/doc/GSEAUserGuideTEXT.htm#\_Running\_a\_Leading.

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## **Examples**

```
data(examplePathways)
data(exampleRanks)
fgseaRes <- fgsea(examplePathways, exampleRanks, nperm=10000, maxSize=500)
# Testing only one pathway is implemented in a more efficient manner
fgseaRes1 <- fgsea(examplePathways[1], exampleRanks, nperm=10000)</pre>
```

fgseaLabel

Runs label-permuring gene set enrichment analysis.

# **Description**

Runs label-permuring gene set enrichment analysis.

# Usage

```
fgseaLabel(pathways, mat, labels, nperm, minSize = 1, maxSize = Inf,
    nproc = 0, gseaParam = 1, BPPARAM = NULL)
```

# **Arguments**

pathways	List of gene sets to check.
mat	Gene expression matrix. Row name should be the same as in 'pathways'
labels	Numeric vector of labels for the correlation score of the same length as the number of columns in 'mat'
nperm	Number of permutations to do. Minimial possible nominal p-value is about 1/nperm
minSize	Minimal size of a gene set to test. All pathways below the threshold are excluded.
maxSize	Maximal size of a gene set to test. All pathways above the threshold are excluded.
nproc	If not equal to zero sets BPPARAM to use nproc workers (default = 0).
gseaParam	GSEA parameter value, all gene-level statis are raised to the power of 'gsea-Param' before calculation of GSEA enrichment scores.
BPPARAM	Parallelization parameter used in bplapply. Can be used to specify cluster to run. If not initialized explicitly or by setting 'nproc' default value 'bpparam()' is used.

#### Value

A table with GSEA results. Each row corresponds to a tested pathway. The columns are the following:

- pathway name of the pathway as in 'names(pathway)';
- pval an enrichment p-value;
- padj a BH-adjusted p-value;
- ES enrichment score, same as in Broad GSEA implementation;
- NES enrichment score normalized to mean enrichment of random samples of the same size;

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• nMoreExtreme • – a number of times a random gene set had a more extreme enrichment score value;

- size size of the pathway after removing genes not present in 'names(stats)'.
- leadingEdge vector with indexes of leading edge genes that drive the enrichment, see http://software.broadinstitute.org/gsea/doc/GSEAUserGuideTEXT.htm#\_Running\_a\_Leading.

## **Examples**

```
library(limma)
library(GEOquery)
es <- getGEO("GSE19429", AnnotGPL = TRUE)[[1]]
exprs(es) <- normalizeBetweenArrays(log2(exprs(es)+1), method="quantile")
es <- es[!grepl("///", fData(es)$`Gene ID`), ]
es <- es[fData(es)$`Gene ID` != "", ]
es <- es[order(apply(exprs(es), 1, mean), decreasing=TRUE), ]
es <- es[!duplicated(fData(es)$`Gene ID`), ]
rownames(es) <- fData(es)$`Gene ID`

pathways <- reactomePathways(rownames(es))
mat <- exprs(es)
labels <- as.numeric(as.factor(gsub(" .*", "", es$title)))
fgseaRes <- fgseaLabel(pathways, mat, labels, nperm = 1000, minSize = 15, maxSize = 500)</pre>
```

fgseaMultilevel

Runs preranked gene set enrichment analysis.

# **Description**

This feature is based on the adaptive multilevel splitting Monte Carlo approach. This allows us to exceed the results of simple sampling and calculate arbitrarily small P-values.

# Usage

```
fgseaMultilevel(pathways, stats, sampleSize = 101, minSize = 1,
  maxSize = Inf, absEps = 0, nproc = 0, BPPARAM = NULL)
```

# Arguments

pathways	List of gene sets to check.
stats	Named vector of gene-level stats. Names should be the same as in 'pathways'
sampleSize	The size of a random set of genes which in turn has size = pathwaySize
minSize	Minimal size of a gene set to test. All pathways below the threshold are excluded.
maxSize	Maximal size of a gene set to test. All pathways above the threshold are excluded.
absEps	This parameter sets the boundary for calculating the p value.
nproc	If not equal to zero sets BPPARAM to use nproc workers (default = 0).
BPPARAM	Parallelization parameter used in bplapply. Can be used to specify cluster to run. If not initialized explicitly or by setting 'nproc' default value 'bpparam()' is used.

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#### Value

A table with GSEA results. Each row corresponds to a tested pathway. The columns are the following

- pathway name of the pathway as in 'names(pathway)';
- pval an enrichment p-value;
- padj a BH-adjusted p-value;
- log2err the expected error for the standard deviation of the P-value logarithm.
- ES enrichment score, same as in Broad GSEA implementation;
- NES enrichment score normalized to mean enrichment of random samples of the same size;
- size size of the pathway after removing genes not present in 'names(stats)'.
- leadingEdge vector with indexes of leading edge genes that drive the enrichment, see http: //software.broadinstitute.org/gsea/doc/GSEAUserGuideTEXT.htm#\_Running\_a\_Leading.

#### **Examples**

```
data(examplePathways)
data(exampleRanks)
fgseaMultilevelRes <- fgseaMultilevel(examplePathways, exampleRanks, maxSize=500)</pre>
```

fgseaSimpleImpl Runs preranked gene set enrichment analysis for preprocessed input data.

# **Description**

Runs preranked gene set enrichment analysis for preprocessed input data.

# Usage

```
fgseaSimpleImpl(pathwayScores, pathwaysSizes, pathwaysFiltered,
  leadingEdges, permPerProc, seeds, toKeepLength, stats, BPPARAM)
```

#### **Arguments**

pathwayScores Vector with enrichment scores for the 'pathways'.

pathwaysSizes Vector of path sizes.

pathwaysFiltered

Filtered pathways.

leadingEdges Leading edge genes.

permPerProc Parallelization parameter for permutations.

seeds Seed vector

toKeepLength Number of 'pathways' that meet the condition for 'minSize' and 'maxSize'.

stats Named vector of gene-level stats. Names should be the same as in 'pathways'

BPPARAM Parallelization parameter used in bplapply. Can be used to specify cluster to

run. If not initialized explicitly or by setting 'nproc' default value 'bpparam()'

is used.

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#### Value

A table with GSEA results. Each row corresponds to a tested pathway. The columns are the following:

- pathway name of the pathway as in 'names(pathway)';
- pval an enrichment p-value;
- padj a BH-adjusted p-value;
- ES enrichment score, same as in Broad GSEA implementation;
- NES enrichment score normalized to mean enrichment of random samples of the same size;
- nMoreExtreme' a number of times a random gene set had a more extreme enrichment score value;
- size size of the pathway after removing genes not present in 'names(stats)'.
- leadingEdge vector with indexes of leading edge genes that drive the enrichment, see http: //software.broadinstitute.org/gsea/doc/GSEAUserGuideTEXT.htm#\_Running\_a\_Leading.

gmtPathways

Returns a list of pathways from a GMT file.

#### **Description**

Returns a list of pathways from a GMT file.

## Usage

```
gmtPathways(gmt.file)
```

# Arguments

```
gmt.file Path to a GMT file.
```

#### Value

A list of vectors with gene sets.

# **Examples**

```
pathways <- gmtPathways(system.file(
   "extdata", "mouse.reactome.gmt", package="fgsea"))</pre>
```

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multilevelError	Calculates the expected error for the standard deviation of the P-value
	logarithm.

#### **Description**

Calculates the expected error for the standard deviation of the P-value logarithm.

#### Usage

```
multilevelError(pval, sampleSize)
```

# **Arguments**

pval P-value

sampleSize equivavlent to sampleSize in fgseaMultilevel

#### Value

The value of the expected error

## **Examples**

```
expectedError <- multilevelError(pval=1e-10, sampleSize=1001)</pre>
```

multilevelImpl

Calculates P-values for preprocessed data.

# Description

Calculates P-values for preprocessed data.

# Usage

```
multilevelImpl(multilevelPathwaysList, stats, sampleSize, seed, absEps,
    sign = FALSE, BPPARAM = NULL)
```

# **Arguments**

 $\verb|multilevelPath| ways \verb|List|$ 

List of pathways for which P-values will be calculated.

stats Named vector of gene-level stats. Names should be the same as in 'pathways' sampleSize The size of a random set of genes which in turn has size = pathwaySize

seed 'seed' parameter from 'fgseaMultilevel'

absEps This parameter sets the boundary for calculating the p value.

sign This option will be used in future implementations.

BPPARAM Parallelization parameter used in bplapply. Can be used to specify cluster to

run. If not initialized explicitly or by setting 'nproc' default value 'bpparam()'

is used.

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#### Value

List of P-values.

plotEnrichment

Plots GSEA enrichment plot.

# Description

Plots GSEA enrichment plot.

# Usage

```
plotEnrichment(pathway, stats, gseaParam = 1, ticksSize = 0.2)
```

# **Arguments**

pathway Gene set to plot.
stats Gene-level statistics.
gseaParam GSEA parameter.

ticksSize width of vertical line corresponding to a gene (default: 0.2)

# Value

ggplot object with the enrichment plot.

# **Examples**

plotGseaTable

Plots table of enrichment graphs using ggplot and gridExtra.

# Description

Plots table of enrichment graphs using ggplot and gridExtra.

# Usage

```
plotGseaTable(pathways, stats, fgseaRes, gseaParam = 1,
   colwidths = c(5, 3, 0.8, 1.2, 1.2), render = TRUE)
```

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## **Arguments**

pathways Pathways to plot table, as in 'fgsea' function. stats Gene-level stats, as in 'fgsea' function.

fgseaRes Table with fgsea results.

gseaParam GSEA-like parameter. Adjusts displayed statistic values, values closer to 0 flat-

ten plots. Default = 1, value of 0.5 is a good choice too.

colwidths Vector of five elements corresponding to column width for grid.arrange. If col-

umn width is set to zero, the column is not drawn.

render If true, the plot is rendered to the current device. Otherwise, the grob is returned.

Default is true.

#### Value

TableGrob object returned by grid.arrange.

## **Examples**

reactomePathways

Returns a list of Reactome pathways for given Entrez gene IDs

# Description

Returns a list of Reactome pathways for given Entrez gene IDs

#### Usage

```
reactomePathways(genes)
```

#### **Arguments**

genes Entrez IDs of query genes.

## Value

A list of vectors with gene sets.

# **Examples**

```
data(exampleRanks)
pathways <- reactomePathways(names(exampleRanks))</pre>
```

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