

Package ‘escape’

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Title Easy single cell analysis platform for enrichment

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Description A bridging R package to facilitate gene set enrichment analysis (GSEA) in the context of single-cell RNA sequencing. Using raw count information, Seurat objects, or SingleCellExperiment format, users can perform and visualize GSEA across individual cells.

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GeneSetEnrichment, Sequencing, GeneSignaling, Pathways

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SingleCellExperiment, limma, ggridges, msigdb, stats,
BiocParallel, Matrix

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dittoSeq (>= 1.1.2)

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enrichIt	<i>Calculate gene set enrichment scores for single-cell data</i>
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Description

This function allows users to input both the single-cell RNA-sequencing counts and any gene set pathways either from the stored data or from other sources. The enrichment calculation itself uses the `gsva` R package and the poisson distribution for RNA.

Usage

```
enrichIt(obj, gene.sets = NULL, groups = 1000, cores = 2)
```

Arguments

<code>obj</code>	The count matrix, Seurat, or SingleCellExperiment object.
<code>gene.sets</code>	Gene sets from getGeneSets to use for the enrichment analysis. Alternatively a simple base R list where the names of the list elements correspond to the name of the gene set and the elements themselves are simple vectors of gene names representing the gene set.
<code>groups</code>	The number of cells to separate the enrichment calculation.
<code>cores</code>	The number of cores to use for parallelization.

Value

Data frame of normalized enrichment scores (NES)

Author(s)

Nick Borcharding, Jared Andrews

See Also

[getGeneSets](#) to collect gene sets.

Examples

```
# download HALLMARK gene set collection
GS <- getGeneSets(library = "H")
GS <- GS[c(1:2)] #Reduce list size for example
seurat_ex <- suppressWarnings(SeuratObject::pbmc_small)
ES <- enrichIt(obj = seurat_ex, gene.sets = GS)

# alternatively, construct your own list of gene sets
myGS <- list(Housekeeping = c("ACTA1", "ACTN1", "GAPDH"),
  Cancer = c("TP53", "BRCA2", "ERBB2", "MYC"))
```

getGeneSets

Get a collection of gene sets to perform enrichment on

Description

This function allows users to select libraries and specific gene.sets to form a GeneSetCollection that is a list of gene sets.

Usage

```
getGeneSets(species = "Homo sapiens", library = NULL, gene.sets = NULL)
```

Arguments

species	The scientific name of the species of interest in order to get correcent gene nomenclature
library	Individual collection(s) of gene sets, e.g. c("H", "C5"). See https://www.gsea-msigdb.org/gsea/msigdb/collections.jsp for all MSigDB collections.
gene.sets	Select gene sets or pathways, using specific names, example: pathways = c("HALLMARK_TNFA_SIGNA... Will only be honored if library is set, too.

Value

A GeneSetCollection object containing the requested GeneSet objects.

Author(s)

Nick Borcharding, Jared Andrews

Examples

```
GS <- getGeneSets(library = "H")
```

getSignificance	<i>Perform significance testing between groups and enrichment scores.</i>
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Description

This functions takes the enrichment scores and performs statistical testing to evaluate the difference by group selected. The function can perform 3 tests: 1) linear model based on the limma package, 2) Welch's T test, and 3) one-way ANOVA. The output includes adjusted p-values based on the Benjamini Hochberg method.

Usage

```
getSignificance(enriched, group = NULL, fit = "linear.model")
```

Arguments

enriched	The output of enrichIt .
group	The parameter to group for the comparison, should a column of the enriched input
fit	The test used for significance, either linear.model, ANOVA, or T.test

Value

Data frame of test statistics

See Also

[enrichIt](#) for generating enrichment scores.

Examples

```
ES2 <- readRDS(url(
  "https://ncborcherding.github.io/vignettes/escape_enrichment_results.rds"))
output <- getSignificance(ES2, group = "Type", fit = "linear.model")
```

masterPCAPlot	<i>Visualize the components of the PCA analysis of the enrichment results</i>
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Description

Graph the major gene set contributors to the [pcaEnrichment](#).

Usage

```
masterPCAPlot(enriched, PCx, PCy, top.contribution = 10)
```

Arguments

enriched	The output of enrichIt .
PCx	The principal component graphed on the x-axis.
PCy	The principal component graphed on the y-axis.
top.contribution	The number of gene sets to graph, organized by PCA contribution.

Value

ggplot2 object summarizing the PCA for the enrichment scores

See Also

[enrichIt](#) for generating enrichment scores.

Examples

```
ES2 <- readRDS(url(
  "https://ncborcherding.github.io/vignettes/escape_enrichment_results.rds"))
masterPCAPlot(ES2, PCx = "PC1", PCy = "PC2", top.contribution = 10)
```

pcaEnrichment

Density plot of the principal components

Description

Density plot of the principal components

Usage

```
pcaEnrichment(
  PCAout,
  PCx,
  PCy,
  colors = c("#0348A6", "#7AC5FF", "#C6FDEC", "#FFB433", "#FF4B20"),
  contours = TRUE,
  facet = NULL
)
```

Arguments

PCAout	The output of performPCA
PCx	The principal component graphed on the x-axis
PCy	The principal component graphed on the y-axis
colors	The color palette for the density plot
contours	Binary classifier to add contours to the density plot
facet	A parameter to separate the graph

Value

ggplot2 object of the results of PCA for the enrichment scores

See Also

[performPCA](#) for generating PCA results.

Examples

```
ES2 <- readRDS(url(
  "https://ncborcherding.github.io/vignettes/escape_enrichment_results.rds"))
PCA <- performPCA(enriched = ES2, groups = c("Type", "Cluster"))
pcaEnrichment(PCA, PCx = "PC1", PCy = "PC2", contours = TRUE)
```

performPCA

Calculate Principal Components for the Enrichment Scores

Description

Using all or selected enrichment scores of individual single-cells, this function will calculate principal components using scaled values and attach to the output columns to use to graph later.

Usage

```
performPCA(enriched, groups)
```

Arguments

enriched	The output of enrichIt .
groups	The column headers to use in future graphing functions.

Value

Data frame of principal components

Author(s)

Nick Borchering

Examples

```
ES2 <- readRDS(url(
  "https://ncborcherding.github.io/vignettes/escape_enrichment_results.rds"))
PCA <- performPCA(enriched = ES2, groups = c("Type", "Cluster"))
```

ridgeEnrichment *Generate a ridge plot to examine enrichment distributions*

Description

This function allows to the user to examine the distribution of enrichment across groups by generating a ridge plot.

Usage

```
ridgeEnrichment(  
  enriched,  
  group = "cluster",  
  gene.set = NULL,  
  scale.bracket = NULL,  
  facet = NULL,  
  add.rug = FALSE,  
  colors = c("#0348A6", "#7AC5FF", "#C6FDEC", "#FFB433", "#FF4B20")  
)
```

Arguments

enriched	The output of enrichIt
group	The parameter to group, displayed on the y-axis.
gene.set	The gene set to graph on the x-axis.
scale.bracket	This will filter the enrichment scores to remove extreme outliers. Values entered (1 or 2 numbers) will be the filtering parameter using z-scores of the selected gene.set. If only 1 value is given, a secondary bracket is automatically selected as the inverse of the number.
facet	A parameter to separate the graph.
add.rug	Binary classifier to add a rug plot to the x-axis.
colors	The color palette for the ridge plot.

Value

ggplot2 object with ridge-based distributions of selected gene.set

See Also

[enrichIt](#) for generating enrichment scores.

Examples

```
ES2 <- readRDS(url(
  "https://ncborcherding.github.io/vignettes/escape_enrichment_results.rds"))
ridgeEnrichment(ES2, gene.set = "HALLMARK_DNA_REPAIR", group = "cluster",
  facet = "Type", add.rug = TRUE)
```

splitEnrichment

Generate a split violin plot examine enrichment distributions

Description

This function allows to the user to examine the distribution of enrichment across groups by generating a split violin plot.

Usage

```
splitEnrichment(
  enriched,
  x.axis = NULL,
  scale.bracket = NULL,
  split = NULL,
  gene.set = NULL,
  colors = c("#0348A6", "#7AC5FF", "#C6FDEC", "#FFB433", "#FF4B20")
)
```

Arguments

enriched	The output of enrichIt
x.axis	Optional parameter for seperation.
scale.bracket	This will filter the enrichment scores to remove extreme outliers. Values entered (1 or 2 numbers) will be the filtering parameter using z-scores of the selected gene.set. If only 1 value is given, a seocndary bracket is autommatically selected as the inverse of the number.
split	The parameter to split, must be binary.
gene.set	The gene set to graph on the y-axis.
colors	The color palette for the ridge plot.

Value

ggplot2 object violin-based distributions of selected gene.set

See Also

[enrichIt](#) for generating enrichment scores.

Examples

```
ES2 <- readRDS(url(
  "https://ncborcherding.github.io/vignettes/escape_enrichment_results.rds"))
splitEnrichment(ES2, x.axis = "cluster", split = "Type",
  gene.set = "HALLMARK_DNA_REPAIR")
```

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