

Package ‘gsEasy’

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 gsEasy-package

Gene Set Enrichment Analysis in R

Description

R-interface to C++ implementation of the rank/score based GSEA test described by Subramanian et al 2005.

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References

Subramanian, A, Tamayo, P, Mootha, VK, Mukherjee, S, Ebert, BL, Gillette, MA, Paulovich, A, Pomeroy, SL, Golub, TR, Lander, ES, Mesirov, JP (2005). Gene set enrichment analysis: a knowledge-based approach for interpreting genome-wide expression profiles. Proc. Natl. Acad. Sci. U.S.A., 102, 43:15545-50, doi: 10.1073/pnas.0506580102.

Ashburner et al. Gene ontology: tool for the unification of biology (2000) Nat Genet 25(1):25-9

 get_GO_gene_sets

Create list of gene sets defined by GO term annotation

Description

Note, this function takes several minutes to execute.

Usage

```
get_GO_gene_sets(
  GO_annotation_file,
  GO_file = "http://purl.obolibrary.org/obo/go.obo",
  min_genes = 15,
  max_genes = 500,
  verbose = TRUE
)
```

Arguments

GO_annotation_file

File path of annotation file, which should contain a column of genes and a column of terms. Can be downloaded from at http://geneontology.org/gene-associations/gene_association.goa_human.gz.

GO_file

File path of gene ontology.

min_genes	Minimum number of genes in gene sets.
max_genes	Maximum number of genes in gene sets.
verbose	Print progress.

Value

List of character vectors of term IDs.

get_ontological_gene_sets	<i>Create list of gene sets defined by ontological annotation</i>
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Description

Create list of gene sets defined by ontological annotation

Usage

```
get_ontological_gene_sets(ontology, gene, term, min_genes = 1, max_genes = 500)
```

Arguments

ontology	ontology_index object.
gene	Character vector of genes.
term	Character vector of term IDs annotated to corresponding genes.
min_genes	Minimum number of genes in gene sets.
max_genes	Maximum number of genes in gene sets.

Value

List of character vectors of term IDs.

GO_gene_sets	<i>GO term gene sets</i>
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Description

List of gene sets annotated by each GO term

Format

List of character vectors of genes per GO term, and named by term ID.

Details

Based on gene-GO term annotations downloaded from geneontology.org. Only contains gene sets for terms with up to 500 genes.

gset	<i>Gene set enrichment test</i>
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Description

Gene set enrichment test

Usage

```
gset(
  S,
  N = NULL,
  r = NULL,
  p = 1,
  min_its = 200,
  max_its = 1e+05,
  significance_threshold = 0.05,
  log_dismiss = -10,
  raw_score = FALSE
)
```

Arguments

S	Ranks of gene set
N	Integer value. Only required if r is not specified.
r	Rank/correlation scores. If S is character, then r must be named by gene or be a character vector of the gene names in rank order (necessarily containing S).
p	Weighting of ranking/correlations, see Subramanian et. al 2005.
min_its	Minimum number of null permutations to compare.
max_its	Maximum number of null permutations to compare.
significance_threshold	Maximum p-value of significant result.
log_dismiss	Threshold log probability of returning a significant result, below which function returns current p-value.
raw_score	Logical value determining whether to return the raw value of the gene set enrichment score.

Value

Numeric value - p-value of enrichment.

Examples

```
gset(S=1:5 * 2, N=1000)
gset(S=letters[1:3], r=letters)
```

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