# Package 'RITAN'

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<b>Description</b> Tools for comprehensive gene set enrichment and extraction of multi-resource high confidence subnetworks. RITAN facilitates bioinformatic tasks for enabling network biology research.
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# Description

as.graph

wrapper to convert a data.frame from RITAN an igraph graph object

as.graph

# Usage

```
as.graph(mat, p1 = 1, p2 = 3, ...)
```

# Arguments

mat	t matrix or data frame describing a network	
p1	[1] column of first interactor	
p2	[3] column of second interactor	
	further options passed on to igraph::graph()	

# Value

igraph object

check\_any\_net\_input 3

#### **Examples**

```
## Not run:
G <- as.graph(network_list$PID)
## End(Not run)</pre>
```

```
check_any_net_input check_any_net_input
```

# **Description**

A Quality Control function. This function applies check\_net\_input() to all available resources (default).

# Usage

```
check_any_net_input(set, resources = names(network_list))
```

# **Arguments**

set An input list of genes to check against references.

resources The collection of network resources to check within.

# Value

Logical vector indicating if the genes in "set" are within ANY of the resources.

# **Examples**

```
#' ## Check if genes in myGeneSet are annotated by any resource in "network_list" (default).
library(RITANdata)
myGeneSet <- c('BRCA1', 'RAD51C', 'VAV1', 'HRAS', 'ABCC1', 'CYP1B1', 'CYP3A5')
yorn <- check_any_net_input( myGeneSet )
print(yorn)</pre>
```

check\_net\_input

check\_net\_input

# **Description**

A Quality Control function. This function will compare an input list of genes to a network reference and report if each member of the input is present in the resource.

```
check_net_input(set, ref, check4similar = FALSE, entity1name = "p1",
  entity2name = "p1")
```

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#### **Arguments**

set An input list of genes to check against a reference.

ref A reference of network data. See readSIF().

check4similar Logical flag. If TRUE, a case-insensitive grep will be used for name matching.

For genes in families with many related members (e.g. ABC\*, FAM\*, etc.), this will not be ideal. We intend this option as a QC screening method to identify if

case, punctuaiton, etc is causing fewer than expected matches.

entity1name The column name in "ref" of the first entity. Default = "p1."

entity2name The column name in "ref" of the second entity. Default = "p2."

#### Value

Character vector of "yes/no" indicating "within-ref/not"

# **Examples**

```
## Return a "yes/no" vector indicating if each gene in myGeneSet is annotated with any term in GO
## If no match, this function can attempt to suggest closest matches (check4similar = TRUE)
library(RITANdata)
myGeneSet <- c('BRCA1', 'RAD51C', 'VAV1', 'HRAS', 'ABCC1', 'CYP1B1', 'CYP3A5')
yorn <- check_net_input( myGeneSet, network_list[["CCSB"]] )
print(yorn)

yorn <- check_net_input( myGeneSet, network_list[["PID"]] )
print(yorn)

## See check_any_net_input() for efficiently checking across all resources.</pre>
```

enrichment\_symbols

enrichment\_symbols

#### **Description**

This function is called by term\_enrichment() and term\_enrichment\_by\_subset(). The user may call it directly, but we suggest using term\_enrichment(). The function uses the resources currently loaded into the active\_genesets vector. See load\_geneset\_symbols().

#### Usage

```
enrichment_symbols(geneset, term = NULL, all_symbols = NA, ...)
```

# **Arguments**

geneset vector of gene symbols to be evaluated

term a list containing specific gene set term(s) and their corresponding gene symbols

contained in one of the annotation resources, default is all gene set terms in the

GO, ReactomePathways, KEGG\_filtered\_canonical\_pathways, and MSigDB\_Hallmarks

libraries

all\_symbols gene symbols to be evaluated, identified by gene symbol name. Default is all

protein coding genes. This parameter should be manipulated to include only the

gene symbols that pertain to the user's analysis.

... additional arguments are not used

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#### **Details**

Outputs a data frame containing the gene set name, a hypergeometric-test p value, the number of genes from the input gene list that occur in the gene set, the number of genes in the gene set, the gene symbols for the genes in the input gene list, and the q value.

#### Value

results matrix of input gene list compared to active gene sets. Q value is calculated using entire group of active gene sets.

# **Examples**

```
require(RITANdata)
myGeneSet <- c('BRCA1','RAD51C','VAV1','HRAS','ABCC1','CYP1B1','CYP3A5')</pre>
## Not run:
## We suggest using term_enrichment() instead. E.g.:
e <- enrichment_symbols(myGeneSet, 'GO')</pre>
## End(Not run)
## But, you may use enrichment_symbols() directly for an individual term:
load_geneset_symbols('GO')
e <- enrichment_symbols(myGeneSet, 'DNA_repair')</pre>
print(e)
## Not run:
## Gene set enrichment using intersection of gene symbols
    provided in geneset parameter and all protein coding genes.
enrichment_symbols(geneset = vac1.day@vs31.de.genes)
## choose which terms to evaluate
t <- active_genesets[1:5]
## Test enrichment of that set of terms
enrichment_symbols(geneset = vac1.day@vs31.de.genes, term = t)
## End(Not run)
```

geneset\_overlap

geneset\_overlap

# **Description**

Return assymetric matrix of the fraction of genes shared between sets. E.G. The fraction of the first set that is "covered" by or "overlaps" the second set.

```
geneset_overlap(s1, s2 = s1, s.size = unlist(lapply(s1, length)))
```

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# **Arguments**

s1	The first geneset
s2	the second geneset
s.size	Denominator used in each comparison. The default is to determint the lengths of elements in "s1"

#### Value

results matrix of input gene list compared to active gene sets. Q value is calculated using entire group of active gene sets.

# **Examples**

```
 require(RITANdata) \\ r <- geneset\_overlap( geneset\_list$MSigDB\_Hallmarks, geneset\_list$NetPath\_Gene\_regulation ) \\ heatmap(r, col = rev(gray(seq(0,1,length.out = 15))) ) \\ summary(c(r))
```

icon\_test

icon\_test

# **Description**

"icon" is an abbreviation for the "interconnectivity" of a network or graph.

#### Usage

```
icon_test(n1 = NULL, n2 = NULL, s = 100, verbose = TRUE, ...)
```

# **Arguments**

n1	[NULL] the first network. See network_overlap().
n2	[NULL] the second network. See network_overlap().
S	[100] teh number of random permutations to make.
verbose	[TRUE] If optional text describing what the algorithm is doing should be shown in the console.
	Additional argumetns are passed on to the specific test performed

#### **Details**

This function handles different inputs and directs them to the appropriate "icon" testing method. Depending on the values given to "n1" and "n2," a different specific test is performed.

Note that the specific functions called make use of the "param" attribute of each input. These parameters are populated by network\_overlap() so that the permutation reflects the exact procedure that was done to generate "n1" and/or "n2."

#### Value

metrics and significance of the network overlap

#### **Examples**

```
## Not run:
icon_test( n1=n, s=10)
## End(Not run)
```

# **Description**

The character array returned is, by default, all human protein coding gene symbols. This variable defines the "universe of possible genes" for use in enrichment. Users should load a different "universe" or filter this one down to the most appropriate setting for their current study. For example, if running RNA-Seq, genes are in the universie if they are detected in any sample.

#### Usage

```
load_all_protein_coding_symbols(file = "ftp://ftp.ebi.ac.uk/pub/databases/genenames/new/tsv/loc
col_name = "symbol")
```

# **Arguments**

file file name of a table containing gene symbols col\_name column name within "file" that contains symbols

#### Value

A unique list of gene symbols from the current protein coding set at the EBI

```
load_geneset_symbols load_geneset_symbols
```

# Description

For most applications, this function is used internally by term\_enrichment(). Users may call this function directly in some cases to force FDR adjustment to be across multiple resources. See Vignette for more details.

```
load_geneset_symbols(gmt = NA, gmt_dir = "", verbose = TRUE)
```

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#### **Arguments**

gmt Either 1) name of pre-loaded resource (i.e. names(geneset\_list)) or 2) gmt file

containing annotation resources for enrichment annotation

gmt\_dir location of gmt file named in gmt parameter

verbose print results to screen

#### **Details**

load\_geneset\_symbols allows the user to specify an annotation resource (e.g. Gene Ontology terms) to use in enrichment analysis. The expectation is that the annotation resource contains of at least one set of genes in the form of a list. The RITAN package comes with 15 pre-loaded annotation resources. The default active annotation resources are GO, ReactomePathways, KEGG\_filtered\_canonical\_pathways, and MSigDB\_Hallmarks.

The result of calling this function is to set the variable "active\_genesets" which will be used by further functions.

#### Value

R list object named active\_genesets

#### **Examples**

```
## Load generic GO-slim terms
require(RITANdata)
load_geneset_symbols("GO_slim_generic")
print(length(active_genesets))
print(head(active_genesets[[1]]))
## Not run:
## load the default set of resources into "active_genesets"
load_geneset_symbols()
## Use only the Reactome Pathways annotation resource.
load_geneset_symbols(gmt="ReactomePathways")
## Suppresses output message describing the annotation resource and size.
load_geneset_symbols(gmt="ReactomePathways", verbose=FALSE)
## To list the available resources within RITAN:
print(names(geneset_list))
## You can also load your own data
load_geneset_symbols(gmt="myFile.gmt")
## End(Not run)
```

network\_overlap

network\_overlap

#### **Description**

network\_overlap

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#### Usage

```
network_overlap(gene_list = NA, resources = c("PID", "TFe", "dPPI", "CCSB",
   "STRING"), minStringScore = 700, minHumanNetScore = 0.4, minScore = 0,
   verbose = TRUE, dedup = TRUE, directed_net = FALSE,
   include_neighbors = FALSE, STRING_cache_directory = NA,
   STRING_species = 9606, STRING_version = "10")
```

#### **Arguments**

gene\_list A list of genes to use. The function will identify edges across resources for or

among these genes; identify the induced subnetwork around the gene\_list.

resources Name of network resource(s) to use.

minStringScore If STRING is among the resources, only edges of at least the indicated score

will be included.

minHumanNetScore

If HumanNet is among the resources, only edges of at least the indicated score

will be included.

minScore Same as above, but used for any other networks where "score" is provided

verbose If TRUE (default), the function will update the user on what it is doing and how

many edges are identified for each resource.

dedup If TRUE (Default = TRUE), remove edges reported by multiple resources. The

edge type will be a semi-colon delimited list of the resources that had reported

the interaction.

directed\_net Logical indicating if the network resources should be interpreted as directed.

include\_neighbors

Logical to include 1st neighbors of "gene\_list" (genes not in gene\_list, but di-

rectly connected to them) in the induced subnetwork.

STRING\_cache\_directory

A directry where STRING data files are cached to speed up subsequent queries; no need to re-download. If NA (the default), caches STRING data in your Rpackages directory. If "", uses a temporary directory that is cleared when the

R-session closes.

STRING\_species Sepcies taxon ID (number) to use in searching STRING data. (Default = 9606)

STRING\_version Version of the STRING database (Default = "10")

#### Value

Data table describing the induced subnetwork for "gene\_list" across the requested resources.

```
## Get interactions among a list of genes from the PID: Pathway Interaction Database
require(RITANdata)
myGeneSet <- c('BRCA1','RAD51C','VAV1','HRAS','ABCC1','CYP1B1','CYP3A5')
sif <- network_overlap( myGeneSet, resources = 'PID')
print(sif)

## Not run:
## Get the PPI network induced by genes within myGeneSet
## Use 4 seperate resources, but trim STRING to only include more confident interactions</pre>
```

```
sif <- network_overlap( myGeneSet, c('dPPI', 'PID', 'CCSB', 'STRING'), minStringScore = 500 )
## End(Not run)</pre>
```

```
plot.term_enrichment plot.term_enrichment
```

# Description

plot.term\_enrichment

# Usage

```
## S3 method for class 'term_enrichment'
plot(x = NA, min_q = 0.05, max_terms = 25,
    extend_mar = c(0, 10, 0, 0), ...)
```

# **Arguments**

x	data frame returned by term_enrichment
min_q	Only q-values more significant than this threshold will be plotted. Default $= 0.05$ .
max_terms	Up to max_terms will be plotted. Default = 25.
extend_mar	Term names can be long. We attempt to keep them readable by extending the left-hand-side margins automatically. Default = $c(0,10,0,0)$ added to par()\$mar.
	Additional arguments are passed on to plot()

#### Value

silent return from plot

# **Examples**

```
require(RITANdata)
e <- term_enrichment(vac1.day0vs31.de.genes, resources = 'GO_slim_generic')
plot(e, min_q = .1)</pre>
```

```
plot.term\_enrichment\_by\_subset \\ plot.term\_enrichment\_by\_subset
```

# Description

```
plot.term_enrichment_by_subset
```

#### Usage

```
## S3 method for class 'term_enrichment_by_subset'
plot(x, show_values = TRUE,
   annotation_matrix = NA, low = "white", high = "#2166AC",
   return_ggplot_object = FALSE, label_size_x = 16, label_angle_x = -30,
   label_size_y = 9, wrap_y_labels = 20, grid_line_color = "white",
   mid = 0, cap = NA, annotation_palates = c("Reds", "Greens", "Purples",
   "Greys", "BuPu", "RdPu", "BrBG", "PiYG", "Spectral"),
   annotation_legend_x = -0.3, ...)
```

#### **Arguments**

```
data frame returned by term_enrichment_by_subset
show values
                  True or False, plot values on the heatmap
annotation_matrix
                  a matrix() of group-levle characteristics - same number of columns as "m"
                  color for low end of range
low
high
                  color for high end of range
return_ggplot_object
                  logical flag (default FALSE) that if TRUE, the ggplot object for the plot is re-
                  turned
                  size of text for x label. Default lable_size_x=16
label_size_x
                  angle for text for x label. Default is -30 degrees
label_angle_x
label_size_y
                  size of text for y label. Default label_size_y=9
wrap_y_labels
                  Number of characters to wrap row labels
grid_line_color
                  color o grid lines between cells. Default is white.
                  sets lower threshold for color scale
mid
                  Clip numeric values to this maximum threshold
cap
annotation_palates
                  Color palates (RColorBrewer) used for each row of the annotation matrix
annotation_legend_x
                  offset for placing the legend
                  further areguments are not used at this time. If the user wants to modify the plot,
                  use return_ggplot_object = TRUE.
```

# Value

silent return, unless return\_ggplot\_object==TRUE. Then, the ggplot object for the plot is returned.

```
## Create list of gene sets to evaluate.
## This example is from a vaccine study where we pre-generated differentially expressed genes.
## This object will be passed to the groups parameter.
require(RITANdata)
vac1.de.genes <- list(vac1.day0vs31.de.genes, vac1.day0vs56.de.genes)
names(vac1.de.genes) <- c("Day0vs31", "Day0vs56")
print(str(vac1.de.genes))</pre>
```

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```
## Not run:
## Run term_enrichment_by_subset on the two results.
## This function usually takes a few seconds to a minute to run.
m <- term_enrichment_by_subset(groups = vac1.de.genes, q_value_threshold = .9)
summary(m)
plot( m, label_size_y = 4, show_values = FALSE )
## End(Not run)</pre>
```

readGMT

readGMT

# **Description**

Created for simplification of reading .gmt files into RITAN.

# Usage

```
readGMT(f = NA)
```

# **Arguments**

f

GMT file name. Please provide a full path if the file is not in the current working directory.

# Value

A list() where the name of each entry is the term (first column of GMT file) and the value is a chr array of genes associated with the term.

readSIF

|--|--|--|

# Description

This function reads a data table into R; the data table describes network interactions. It is named for the Simple Interaction Format (SIF), but can read any data table if the users identifies which columns contain the pertinent data (see below).

#### Usage

```
readSIF(file = NA, header = FALSE, sep = "\t^n, as.is = TRUE, p1 = 1, p2 = 2, et = 3, score = NA, ...)
```

# **Arguments**

file	location of file
header	indicator of presense of header on file
sep	file delimiter - used by read.table()
as.is	logical (default TRUE)
p1	Column number for the 1st entity. Default = 1.
p2	Column number for the 2nd entity. Default = $2$ .
et	Column number for the edge type. Default = 3. Optionally, it may be a string label to be used as the edge type for all interactions from the input file.
score	Column number for edge scores or weights. Default = NA (no score read).
	Other options to read.table().

#### **Details**

The SIF file format is a 3-column format, with an optional 4th column: <entity-1><tab><edge-type><tab><entity-2><tab><score>

Entities may be genes, proteins, metabolites, etc. The edge type typically conveys the type of relationship that exists between the two entities, such as physical interaciton, phosphorylation, or activation.

#### Value

Returns a data.frame with 3 (or 4) columns of data.

resource\_reduce Merge terms across resources to reduce the number of redundant and semi-redundant terms

**Description** 

resource\_reduce Merge terms across resources to reduce the number of redundant and semi-redundant terms

#### Usage

```
resource_reduce(genesets = NULL, min_overlap = 0.8, verbose = TRUE)
```

#### **Arguments**

genesets the input genesets to consider. May be from one or multiple resources.

min\_overlap terms that share at least this fraction of genes will be merged

verbose if TRUE, print status and summary output

mutual\_overlap [TO DO; FEATURE NOT IMPLEMENTED] if TRUE, the overlap must be re-

ciprocal in order to merge

#### Value

the list of terms, after merging to reduce redundaat and semi-redundant terms

# **Examples**

```
require(RITANdata)
r <- resource_reduce( geneset_list$DisGeNet )</pre>
```

```
show\_active\_genesets\_hist \\ show\_active\_genesets\_hist
```

# **Description**

function to plot distribution of size of active\_genesets object

# Usage

```
show_active_genesets_hist(nbins = 50, ...)
```

#### **Arguments**

nbins Number of bins to include in histogram
... further argumants are passed on to plot()

#### Value

NULL. The plot is shown.

# **Examples**

```
require(RITANdata)
load_geneset_symbols('GO_slim_generic')
show_active_genesets_hist()

## Not run:
## Show the distribution of geneset sizes for the default set of geneset resources
load_geneset_symbols()
show_active_genesets_hist()

## Show the distribution of geneset sizes for a specific resource
load_geneset_symbols(gmt="ReactomePathways")
show_active_genesets_hist()

## End(Not run)

summary.term_enrichment

summary.term_enrichment
```

# Description

summary.term\_enrichment

# Usage

```
## S3 method for class 'term_enrichment'
summary(object, ...)
```

#### **Arguments**

```
object data frame returned by term_enrichment()
... Further arguments are passed on to head()
```

# Value

the data.frame of top enrichment results

```
require(RITANdata)
e <- term_enrichment( vac1.day0vs31.de.genes, "MSigDB_Hallmarks" )
summary(e, n=3)</pre>
```

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```
summary.term_enrichment_by_subset
summary.term_enrichment_by_subset
```

# Description

```
summary.term_enrichment_by_subset
```

#### Usage

```
## S3 method for class 'term_enrichment_by_subset'
summary(object, verbose = TRUE, ...)
```

# **Arguments**

object data frame returned by term\_enrichment\_by\_subset()
verbose if TRUE (default), print a header describing the data type

... Further arguments are passed on to head()

#### Value

the data.frame of top enrichment results

# **Examples**

```
require(RITANdata)
vac1.de.genes <- list(vac1.day0vs31.de.genes, vac1.day0vs56.de.genes)
names(vac1.de.genes) <- c("Day0vs31", "Day0vs56")
e <- term_enrichment_by_subset(vac1.de.genes, "MSigDB_Hallmarks", q_value_threshold = 0.1 )
summary(e)</pre>
```

term\_enrichment

term enrichment

# **Description**

term\_enrichment evaluates the input gene list for enrichment within each of the annotation resources. This differs from the enrichment\_symbols function which evaluates the gene list for enrichment against all of the annotation resources grouped together.

```
term_enrichment(geneset, resources = resources.default,
  report_resources_separately = FALSE, verbose = TRUE, all_symbols = NA,
  filter_to_intersection = FALSE, ...)
```

#### **Arguments**

vector of gene symbols to be evaluated geneset

list containing the reference gene sets to test for enrichment resources

report\_resources\_separately

logical (default FALSE) flag to report enrichments seperately for each requested resource, or to combine them and produce FDR adjustment across the combined

verbose print the top results for each annotation resource

the background/global set of gene symbols (study dependent; we provide all all\_symbols

protien coding genes as a default)

filter\_to\_intersection

[FALSE] should the background and foreground genesets be subsetted to one

another?

further arguments are passed on to enrichment\_symbols()

#### Value

results matrix of input gene list compared to active gene sets. Q value is calculated within each of the active gene sets.

#### **Examples**

```
## Check if there is enrichment for any "Hallmark" functions within a input set of genes
require(RITANdata)
myGeneSet <- c('BRCA1', 'RAD51C', 'VAV1', 'HRAS', 'ABCC1', 'CYP1B1', 'CYP3A5')</pre>
e <- term_enrichment(myGeneSet, "MSigDB_Hallmarks")</pre>
print( e[1:2, -6] )
## Not run:
term_enrichment(geneset = vac1.day@vs31.de.genes)
term_enrichment(geneset = vac1.day@vs31.de.genes, resources = "MSigDB_Hallmarks")
vac1.day0v31.enrichment <- term_enrichment(geneset = vac1.day0vs31.de.genes, verbose = FALSE)</pre>
## End(Not run)
```

term\_enrichment\_by\_subset

term\_enrichment\_by\_subset

# **Description**

Run enrichment simultaneously across a group of prioritized gene lists. For example, in a time course dataset, one may have a different list of genes that are differentially expressed at each time point. This function facilitates rapid evaluation of term enrichment across time point comparisons. Alternatively, one may have a different list of differentially expressed genes by drug treatment, environmental condition, ect.

```
term_enrichment_by_subset(groups = NA, resources = resources.default,
 q_value_threshold = 0.01, verbose = TRUE, display_type = "q",
 phred = TRUE, ...)
```

#### **Arguments**

groups A list() of genes for enrichment. Each entry in the list() is an input set of genes.

Enrichment is performed for each of these entries.

resources character vector for which resources to use in enrichment

q\_value\_threshold

minimum q-value (FDR adjusted p-value) in any group for the term to be in-

cluded in results

verbose print additional status updates on what the function is doing

display\_type Flag for which data type will be returned. One of "q" (default) for q-values, "p"

for unadjusted p-values, or "n" for the number of genes overlapping the term.

phred Logical flag (default TRUE) to return the -log10 of p/q values
... Further arguments are passed on to enrichment\_symbols()

#### Value

Returns a term-by-study matrix of enrichment values (value determined by "display\_type")

#### **Examples**

```
## Create list of gene sets to evaluate.
## This example is from a vaccine study where we pre-generated differentially expressed genes.
## This object will be passed to the groups parameter.
require(RITANdata)
vac1.de.genes <- list(vac1.day0vs31.de.genes, vac1.day0vs56.de.genes)
names(vac1.de.genes) <- c("Day0vs31", "Day0vs56")
print(str(vac1.de.genes))

## Not run:
## Run term_enrichment_by_subset on the two results.
## This function usually takes a few seconds to a minute to run.
m <- term_enrichment_by_subset(groups = vac1.de.genes, q_value_threshold = .9)
summary(m)
plot( m, label_size_y = 4, show_values = FALSE )

## End(Not run)</pre>
```

vac1.day0vs31.de.genes

This dataset is included as an example in the package:

# **Description**

This dataset is included as an example in the package:

#### Usage

```
vac1.day0vs31.de.genes
```

# Format

An object of class character of length 669.

#### Value

differentially expressed genes at 31 days post-vaccination with vaccine1

#### References

```
https://www.ncbi.nlm.nih.gov/pubmed/26755593
```

# **Examples**

```
## Not run:
  #data("vac1.day0vs31.de.genes")
  te <- term_enrichment(geneset = vac1.day0vs31.de.genes)
## End(Not run)</pre>
```

```
vac1.day0vs56.de.genes
```

This dataset is included as an example in the package:

# **Description**

This dataset is included as an example in the package:

# Usage

```
vac1.day0vs56.de.genes
```

# **Format**

An object of class character of length 471.

# Value

differentially expressed genes at 56 days post-vaccination with vaccine1

#### References

```
https://www.ncbi.nlm.nih.gov/pubmed/26755593
```

```
## Not run:
  #data("vac1.day0vs56.de.genes")
  te <- term_enrichment(geneset = vac1.day0vs56.de.genes)
## End(Not run)</pre>
```

```
vac2.day0vs31.de.genes
```

This dataset is included as an example in the package:

# Description

This dataset is included as an example in the package:

# Usage

```
vac2.day0vs31.de.genes
```

#### **Format**

An object of class character of length 522.

# Value

differentially expressed genes at 31 days post-vaccination with vaccine2

# References

```
https://www.ncbi.nlm.nih.gov/pubmed/26755593
```

# **Examples**

```
## Not run:
  #data("vac2.day0vs31.de.genes")
  te <- term_enrichment(geneset = vac2.day0vs31.de.genes)
## End(Not run)</pre>
```

```
vac2.day0vs56.de.genes
```

This dataset is included as an example in the package:

# Description

This dataset is included as an example in the package:

# Usage

```
vac2.day0vs56.de.genes
```

# Format

An object of class character of length 660.

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#### Value

differentially expressed genes at 56 days post-vaccination with vaccine2

#### References

```
https://www.ncbi.nlm.nih.gov/pubmed/26755593
```

# **Examples**

```
## Not run:
  #data("vac2.day0vs56.de.genes")
  te <- term_enrichment(geneset = vac2.day0vs56.de.genes)
## End(Not run)</pre>
```

writeGMT

writeGMT

# **Description**

Created for future use and simplification of writing .gmt files from the package.

# Usage

```
writeGMT(s, file = NA, link = rep("", length(s)))
```

# **Arguments**

s list of gene sets in current R session. Each entry will become a row in the GMT file.

file file name to write to

link default is "". This is the second column of a GMT file and is usually a hyperlink or note about the origin of the term

# Value

Nothing is returned. A file is written.

22 write\_simple\_table

```
write_simple_table write_simple_table
```

# **Description**

This is a simple wrapper around "write.table" that writes a tab-delimited table with column names, no quoting, and no row names.

# Usage

```
write_simple_table(d = NULL, f = NULL, ...)
```

# Arguments

d R data objectf file path... further options passed on to write.table

# Value

```
invisible (nothing is returned)
```

```
## Not run:
simple wrapper around write.table for writing a tab-delimieted, no row names, tab-seperated file
## End(Not run)
```

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