

Package ‘hypeR’

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Version 2.9.2

Description An R Package for Geneset Enrichment Workflows.

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<code>.check_overlap</code>	<i>Check overlap of signature across genesets</i>
-----------------------------	---

Description

Check overlap of signature across genesets

Usage

`.check_overlap(signature, genesets, threshold = 0.05)`

Arguments

- | | |
|------------------------|-------------------------|
| <code>signature</code> | A vector of symbols |
| <code>genesets</code> | A list of genesets |
| <code>threshold</code> | Minimum percent overlap |

Value

Percent overlap

<code>.dots_multi_plot</code>	<i>Plot top enriched genesets across multiple signatures</i>
-------------------------------	--

Description

Plot top enriched genesets across multiple signatures

Usage

```
.dots_multi_plot(  
  multihyp_data,  
  top = 20,  
  abrv = 50,  
  size_by = c("genesets", "significance", "overlap", "none"),  
  pval_cutoff = 1,  
  fdr_cutoff = 1,  
  val = c("fdr", "pval"),  
  title = ""  
)
```

Arguments

<code>multihyp_data</code>	A list of hyp objects
<code>top</code>	Limit number of genesets shown
<code>abrv</code>	Abbreviation length of genesetlabels
<code>size_by</code>	Size dots by e.g. <code>c("genesets", "significance", "none")</code>
<code>pval_cutoff</code>	Filter results to be less than pval cutoff
<code>fdr_cutoff</code>	Filter results to be less than fdr cutoff
<code>val</code>	Choose significance value e.g. <code>c("fdr", "pval")</code>
<code>title</code>	Plot title

Value

A ggplot object

<code>.dots_plot</code>	<i>Plot top enriched genesets</i>
-------------------------	-----------------------------------

Description

Plot top enriched genesets

Usage

```
.dots_plot(  
  hyp_df,  
  top = 20,  
  abrv = 50,  
  size_by = c("genesets", "significance", "none"),  
  pval_cutoff = 1,  
  fdr_cutoff = 1,  
  val = c("fdr", "pval"),  
  title = ""  
)
```

Arguments

<code>hyp_df</code>	A dataframe from a hyp object
<code>top</code>	Limit number of genesets shown
<code>abrv</code>	Abbreviation length of genesetlabels
<code>size_by</code>	Size dots by e.g. <code>c("genesets", "significance", "none")</code>
<code>pval_cutoff</code>	Filter results to be less than pval cutoff
<code>fdr_cutoff</code>	Filter results to be less than fdr cutoff
<code>val</code>	Choose significance value e.g. <code>c("fdr", "pval")</code>
<code>title</code>	Plot title

Value

A ggplot object

<code>.enrichment_map</code>	<i>Plot enrichment map</i>
------------------------------	----------------------------

Description

Plot enrichment map

Usage

```
.enrichment_map(
  hyp_df,
  genesets,
  similarity_metric = c("jaccard_similarity", "overlap_similarity"),
  similarity_cutoff = 0.2,
  pval_cutoff = 1,
  fdr_cutoff = 1,
  val = c("fdr", "pval"),
  top = NULL,
  title = ""
)
```

Arguments

<code>hyp_df</code>	A dataframe from a hyp object
<code>genesets</code>	A list of genesets
<code>similarity_metric</code>	Metric to calculate geneset similarity
<code>similarity_cutoff</code>	Geneset similarity cutoff
<code>pval_cutoff</code>	Filter results to be less than pval cutoff
<code>fdr_cutoff</code>	Filter results to be less than fdr cutoff
<code>val</code>	Choose significance value shown above nodes e.g. <code>c("fdr", "pval")</code>
<code>top</code>	Limit number of pathways shown
<code>title</code>	Plot title

Value

A visNetwork object

.find_members	<i>Find geneset members</i>
---------------	-----------------------------

Description

Find geneset members

Usage

```
.find_members(id, genesets, nodes, edges)
```

Arguments

id	A vector of ids
genesets	A list of genesets (see rgsets)
nodes	A data frame of labeled nodes (see rgsets)
edges	A data frame of directed edges (see rgsets)

Value

A vector of ids

.format_str	<i>Format a string using placeholders</i>
-------------	---

Description

Format a string using placeholders

Usage

```
.format_str(string, ...)
```

Arguments

string	A an unformatted string with placeholders
...	Variables to format placeholders with

Value

A formatted string

Examples

```
## Not run:
format_str("Format with {1} and {2}", "x", "y")

## End(Not run)
```

.github_rds	Load an rds file directly from github
-------------	---------------------------------------

Description

Load an rds file directly from github

Usage

```
.github_rds(url)
```

Arguments

url A url

Value

A list

.hexa	Adjust alpha of a hex string
-------	------------------------------

Description

Adjust alpha of a hex string

Usage

```
.hexa(hex, percent = 1)
```

Arguments

hex A 6-character hex string (e.g. #000000)
percent Alpha level from 0-1

Value

A hex string

<code>.hiearchy_map</code>	<i>Plot hiearchy map</i>
----------------------------	--------------------------

Description

Plot hiearchy map

Usage

```
.hiearchy_map(  
  hyp_df,  
  rgsets_obj,  
  pval_cutoff = 1,  
  fdr_cutoff = 1,  
  val = c("fdr", "pval"),  
  top = NULL,  
  title = "",  
  graph = FALSE  
)
```

Arguments

<code>hyp_df</code>	A dataframe from a hyp object
<code>rgsets_obj</code>	A relational geneset from a hyp object
<code>pval_cutoff</code>	Filter results to be less than pval cutoff
<code>fdr_cutoff</code>	Filter results to be less than fdr cutoff
<code>val</code>	Choose significance value displayed when hovering nodes e.g. <code>c("fdr", "pval")</code>
<code>top</code>	Limit number of pathways shown
<code>title</code>	Plot title
<code>graph</code>	Return an igraph object instead

Value

A visNetwork object

.hyperdb_url	<i>Get base url for hyperdb</i>
--------------	---------------------------------

Description

Get base url for hyperdb

Usage

```
.hyperdb_url(api = FALSE)
```

Value

A base url

.hyper_enrichment	<i>Overrepresentation test via hyper-geometric distribution</i>
-------------------	---

Description

Overrepresentation test via hyper-geometric distribution

Usage

```
.hyper_enrichment(  
  signature,  
  genesets,  
  background = length(unique(unlist(genesets))),  
  plotting = TRUE  
)
```

Arguments

signature	A vector of symbols
genesets	A list of genesets
background	Size of background population genes
plotting	Use true to generate plots

Value

A list of data and plots

<code>.jaccard_similarity</code>	<i>Calculate jaccard similarity of two sets</i>
----------------------------------	---

Description

Calculate jaccard similarity of two sets

Usage

```
.jaccard_similarity(a, b)
```

Arguments

a	A vector
b	A vector

Value

A numerical value

<code>.kstest</code>	<i>One-sided Kolmogorov–Smirnov test</i>
----------------------	--

Description

One-sided Kolmogorov–Smirnov test

Usage

```
.kstest(  
  n.x,  
  y,  
  weights = NULL,  
  weights.pwr = 1,  
  absolute = FALSE,  
  plotting = FALSE,  
  plot.title = ""  
)
```

Arguments

n.x	The length of a ranked list
y	A vector of positions in the ranked list
weights	Weights for weighted score (Subramanian et al.)
weights.pwr	Exponent for weights (Subramanian et al.)
absolute	Takes max-min score rather than the max deviation from null
plotting	Use true to generate plot
plot.title	Plot title

Value

A list of data and plots

.ks_enrichment	<i>Enrichment test via one-sided Kolmogorov–Smirnov test</i>
----------------	--

Description

Enrichment test via one-sided Kolmogorov–Smirnov test

Usage

```
.ks_enrichment(
  signature,
  genesets,
  weights = NULL,
  weights.pwr = 1,
  absolute = FALSE,
  plotting = TRUE
)
```

Arguments

signature	A vector of ranked symbols
genesets	A list of genesets
weights	Weights for weighted score (Subramanian et al.)
weights.pwr	Exponent for weights (Subramanian et al.)
absolute	Takes max-min score rather than the max deviation from null
plotting	Use true to generate plot

Value

A list of data and plots

<code>.overlap_similarity</code>	<i>Calculate overlap similarity of two sets</i>
----------------------------------	---

Description

Calculate overlap similarity of two sets

Usage

```
.overlap_similarity(a, b)
```

Arguments

<code>a</code>	A vector
<code>b</code>	A vector

Value

A numerical value

<code>.reverselog_trans</code>	<i>Custom reverse log transformation of continous ggplot axes</i>
--------------------------------	---

Description

Custom reverse log transformation of continous ggplot axes

Usage

```
.reverselog_trans(base = exp(1))
```

Arguments

<code>base</code>	Logarithm base
-------------------	----------------

.string_args	<i>Convert an arguments list to string format</i>
--------------	---

Description

Convert an arguments list to string format

Usage

```
.string_args(args)
```

Arguments

args	A list of keyword arguments
------	-----------------------------

Value

A string of keyword arguments

Examples

```
## Not run:  
string_args(list(x=15, y="fdr", z=TRUE))  
  
## End(Not run)
```

clean_genesets	<i>Clean labels of genesets</i>
----------------	---------------------------------

Description

Clean labels of genesets

Usage

```
clean_genesets(x)
```

Arguments

x	A vector of labels
---	--------------------

Examples

```
HALLMARK <- msigdb_download("Homo sapiens", "H")  
names(HALLMARK) <- clean_genesets(names(HALLMARK))  
head(names(HALLMARK))
```

enrichr_available	<i>Get enrichr available genesets</i>
-------------------	---------------------------------------

Description

Get enrichr available genesets

Usage

```
enrichr_available(  
  db = c("Enrichr", "YeastEnrichr", "FlyEnrichr", "WormEnrichr", "FishEnrichr")  
)
```

Arguments

db	A species
----	-----------

Value

A dataframe of available genesets

Examples

```
enrichr_available()
```

enrichr_connect	<i>Connect to the enrichr web application</i>
-----------------	---

Description

Connect to the enrichr web application

Usage

```
enrichr_connect(  
  endpoint,  
  db = c("Enrichr", "YeastEnrichr", "FlyEnrichr", "WormEnrichr", "FishEnrichr")  
)
```

Arguments

endpoint	The url endpoint to connect to
db	A species

Value

A web response

enrichr_download	<i>Download data from enrichr in the form of a named list</i>
------------------	---

Description

Download data from enrichr in the form of a named list

Usage

```
enrichr_download(  
  genesets,  
  db = c("Enrichr", "YeastEnrichr", "FlyEnrichr", "WormEnrichr", "FishEnrichr")  
)
```

Arguments

genesets	A name corresponding to available genesets
db	A species

Value

A list of genesets

Examples

```
ATLAS <- enrichr_download("Human_Gene_Atlas")
```

enrichr_gsets	<i>Download data from enrichr in the form of a gsets object</i>
---------------	---

Description

Download data from enrichr in the form of a gsets object

Usage

```
enrichr_gsets(  
  genesets,  
  db = c("Enrichr", "YeastEnrichr", "FlyEnrichr", "WormEnrichr", "FishEnrichr"),  
  clean = FALSE  
)
```

Arguments

genesets	A name corresponding to available genesets
db	A species
clean	Use true to clean labels of genesets

Value

A gsets object

Examples

```
ATLAS <- enrichr_gsets("Human_Gene_Atlas")
```

enrichr_urls	<i>Get url base for species-specific enrichr libraries</i>
--------------	--

Description

Get url base for species-specific enrichr libraries

Usage

```
enrichr_urls(  
  db = c("Enrichr", "YeastEnrichr", "FlyEnrichr", "WormEnrichr", "FishEnrichr")  
)
```

Arguments

db	A species
----	-----------

Value

A url

genesets_Server	<i>Shiny server module for geneset selection</i>
-----------------	--

Description

Shiny server module for geneset selection

Usage

```
genesets_Server(id, clean = FALSE)
```

Arguments

id	A unique namespace identifier matching to interface
clean	Use true to clean geneset names

Value

Shiny server code

genesets_UI	<i>Shiny interface module for geneset selection</i>
-------------	---

Description

Shiny interface module for geneset selection

Usage

```
genesets_UI(id)
```

Arguments

id	A unique namespace identifier
----	-------------------------------

Value

Shiny ui elements

`ggempty`*An empty ggplot*

Description

An empty ggplot

Usage

```
ggempty()
```

Value

A ggplot object

`ggeplot`*Enrichment plot implemented in ggplot*

Description

Enrichment plot implemented in ggplot

Usage

```
ggeplot(n, positions, x_axis, y_axis, title = "")
```

Arguments

<code>n</code>	The length of a ranked list
<code>positions</code>	A vector of positions in the ranked list
<code>x_axis</code>	The x-axis of a running enrichment score
<code>y_axis</code>	The y-axis of a running enrichment score
<code>title</code>	Plot title

Value

A ggplot object

ggvenn	<i>Venn diagram implemented in ggplot</i>
--------	---

Description

Venn diagram implemented in ggplot

Usage

```
ggvenn(a, b, ga, gb, title = "")
```

Arguments

- a A vector for group a
- b A vector for group b
- ga A string label for group a
- gb A string label for group b
- title Plot title

Value

A ggplot object

gsets	<i>A genesets object</i>
-------	--------------------------

Description

- A genesets object
- A genesets object

See Also

rgsets

Public fields

- genesets A named list of genesets
- name A character vector describing source of genesets
- version A character vector describing versioning

Methods

Public methods:

- `gsets$new()`
- `gsets$print()`
- `gsets$list()`
- `gsets$info()`
- `gsets$reduce()`
- `gsets$clone()`

Method `new()`: Create a gsets object

Usage:

```
gsets$new(  
  genesets,  
  name = "Custom",  
  version = "",  
  clean = FALSE,  
  quiet = FALSE  
)
```

Arguments:

`genesets` A named list of genesets

`name` A character vector describing source of genesets

`version` A character vector describing versioning

`clean` Use true to clean labels of genesets

`quiet` Use true to silence warnings

Returns: A new gsets object

Method `print()`: Print genesets information

Usage:

```
gsets$print()
```

Returns: NULL

Method `list()`: Return genesets as a list

Usage:

```
gsets$list()
```

Returns: A list of genesets

Method `info()`: Returns versioning information

Usage:

```
gsets$info()
```

Returns: A character vector with name and version

Method `reduce()`: Reduces genesets to a background distribution of symbols

Usage:

```
gsets$reduce(background)
```

Arguments:

background A character vector of symbols

Returns: A gsets object

Method clone(): The objects of this class are cloneable with this method.

Usage:

```
gsets$clone(deep = FALSE)
```

Arguments:

deep Whether to make a deep clone.

Examples

```
genesets <- list("GSET1" = c("GENE1", "GENE2", "GENE3"),
                 "GSET2" = c("GENE4", "GENE5", "GENE6"),
                 "GSET3" = c("GENE7", "GENE8", "GENE9"))

gsets_obj <- gsets$new(genesets, name="example", version="v1.0")
print(gsets_obj)
```

hyp	A hyp object
-----	--------------

Description

A hyp object
A hyp object

See Also

multihyp

Public fields

data A dataframe returned by hypeR()
plots A list of plots returned by hypeR()
args A list of arguments passed to hypeR()
info Exported information for reproducibility

Methods

Public methods:

- `hyp$new()`
- `hyp$print()`
- `hyp$as.data.frame()`
- `hyp$clone()`

Method `new()`: Create a hyp object

Usage:

```
hyp$new(data, plots = NULL, args = NULL, info = NULL)
```

Arguments:

`data` A dataframe returned by `hypeR()`

`plots` A list of plots returned by `hypeR()`

`args` A list of arguments passed to `hypeR()`

`info` Exported information for reproducibility

Returns: A new hyp object

Method `print()`: Print hyp object

Usage:

```
hyp$print()
```

Returns: NULL

Method `as.data.frame()`: Extract dataframe from hyp object

Usage:

```
hyp$as.data.frame()
```

Returns: NULL

Method `clone()`: The objects of this class are cloneable with this method.

Usage:

```
hyp$clone(deep = FALSE)
```

Arguments:

`deep` Whether to make a deep clone.

Examples

```
data <- data.frame(replicate(5, sample(0:1, 10, rep=TRUE)))
args <- list("arg_1"=1, "arg_2"=2, "arg_3"=3)
hyp_obj <- hyp$new(data, args=args)
```

hypeR*Calculate enrichment of one or more signatures*

Description

Calculate enrichment of one or more signatures

Usage

```
hypeR(  
  signature,  
  genesets,  
  test = c("hypergeometric", "kstest"),  
  background = 23467,  
  power = 1,  
  absolute = FALSE,  
  pval = 1,  
  fdr = 1,  
  plotting = FALSE,  
  quiet = TRUE  
)
```

Arguments

signature	A vector of symbols
genesets	A gsets/rgsets object or a named list of genesets
test	Choose an enrichment type e.g. c("hypergeometric", "kstest")
background	Size or character vector of background population genes
power	Exponent for weights (kstest only)
absolute	Takes max-min score rather than the max deviation from null (kstest only)
pval	Filter results to be less than pval cutoff
fdr	Filter results to be less than fdr cutoff
plotting	Use true to generate plots for each geneset test (may slow performance)
quiet	Use true to suppress logs and warnings

Value

A hyp object

Examples

```
genesets <- msigdb_gsets("Homo sapiens", "C2", "CP:KEGG_LEGACY")

signature <- c("IDH3B", "DLST", "PCK2", "CS", "PDHB", "PCK1", "PDHA1", "LOC642502",
               "PDHA2", "LOC283398", "FH", "SDHD", "OGDH", "SDHB", "IDH3A", "SDHC",
               "IDH2", "IDH1", "OGDHL", "PC", "SDHA", "SUCLG1", "SUCLA2", "SUCLG2")

hyp_obj <- hyperR(signature, genesets, background=2522)
```

hyperdb_available	<i>Check available data to download from hyperdb</i>
-------------------	--

Description

Check available data to download from hyperdb

Usage

```
hyperdb_available()
```

Examples

```
hyperdb_available()
```

hyperdb_gsets	<i>Download data from hyperdb</i>
---------------	-----------------------------------

Description

Download data from hyperdb

Usage

```
hyperdb_gsets(source, gsets)
```

Arguments

source	A source identifier
gsets	A genesets identifier

Value

A list

Examples

```
KEGG <- hyperdb_rgsets("KEGG", "KEGG_v92.0.rds")
```

hyperdb_rgsets

Download data from hyperdb in the form of a rgsets object

Description

Download data from hyperdb in the form of a rgsets object

Usage

```
hyperdb_rgsets(rgsets, version)
```

Arguments

rgsets	A name corresponding to an available relational genesets object
version	A version number

Value

An rgsets object

Examples

```
REACTOME <- hyperdb_rgsets("REACTOME", "70.0")
```

hyp_dots

Visualize hyp/multihyp objects as a dots plot

Description

Visualize hyp/multihyp objects as a dots plot

Usage

```
hyp_dots(
  hyp_obj,
  top = 20,
  abrv = 50,
  size_by = c("genesets", "significance", "none"),
  pval = 1,
  fdr = 1,
  val = c("fdr", "pval"),
  title = "",
  merge = FALSE
)
```

Arguments

hyp_obj	A hyp or multihyp object
top	Limit number of genesets shown
abbrv	Abbreviation length of geneset labels
size_by	Size dots by e.g. c("genesets", "significance", "none")
pval	Filter results to be less than pval cutoff
fdr	Filter results to be less than fdr cutoff
val	Choose significance value for plot e.g. c("fdr", "pval")
title	Plot title
merge	Use true to merge a multihyp object into one plot

Value

A ggplot object

Examples

```
genesets <- msigdb_gsets("Homo sapiens", "C2", "CP:KEGG_LEGACY")

signature <- c("IDH3B", "DLST", "PCK2", "CS", "PDHB", "PCK1", "PDHA1", "LOC642502",
               "PDHA2", "LOC283398", "FH", "SDHD", "OGDH", "SDHB", "IDH3A", "SDHC",
               "IDH2", "IDH1", "OGDHL", "PC", "SDHA", "SUCLG1", "SUCLA2", "SUCLG2")

hyp_obj <- hypeR(signature, genesets, background=2522)

hyp_dots(hyp_obj, val="fdr")
```

hyp_emap

Visualize hyp/multihyp objects as an enrichment map

Description

Visualize hyp/multihyp objects as an enrichment map

Usage

```
hyp_emap(
  hyp_obj,
  similarity_metric = c("jaccard_similarity", "overlap_similarity"),
  similarity_cutoff = 0.2,
  pval = 1,
  fdr = 1,
  val = c("fdr", "pval"),
  top = NULL,
  title = ""
)
```

Arguments

hyp_obj	A hyp or multihyp object
similarity_metric	Metric to calculate geneset similarity
similarity_cutoff	Geneset similarity cutoff
pval	Filter results to be less than pval cutoff
fdr	Filter results to be less than fdr cutoff
val	Choose significance value shown above nodes e.g. c("fdr", "pval")
top	Limit number of pathways shown
title	Plot title

Value

A visNetwork object

Examples

```
genesets <- msigdb_gsets("Homo sapiens", "C2", "CP:KEGG_LEGACY")

signature <- c("IDH3B", "DLST", "PCK2", "CS", "PDHB", "PCK1", "PDHA1", "LOC642502",
               "PDHA2", "LOC283398", "FH", "SDHD", "OGDH", "SDHB", "IDH3A", "SDHC",
               "IDH2", "IDH1", "OGDHL", "PC", "SDHA", "SUCLG1", "SUCLA2", "SUCLG2")

hyp_obj <- hypeR(signature, genesets, background=2522)

hyp_emap(hyp_obj, top=30, val="fdr")
```

hyp_hmap

Visualize hyp/multihyp objects as a hierarchy map

Description

Visualize hyp/multihyp objects as a hierarchy map

Usage

```
hyp_hmap(
  hyp_obj,
  pval = 1,
  fdr = 1,
  val = c("fdr", "pval"),
  top = NULL,
  title = "",
  graph = FALSE
)
```

Arguments

hyp_obj	A hyp or multihyp object
pval	Filter results to be less than pval cutoff
fdr	Filter results to be less than fdr cutoff
val	Choose significance value displayed when hovering nodes e.g. c("fdr", "pval")
top	Limit number of pathways shown
title	Plot title
graph	Return an igraph object instead

Value

A visNetwork object

Examples

```
genesets <- hyperdb_rgsets("REACTOME", "70.0")

signature <- c("IDH3B", "DLST", "PCK2", "CS", "PDHB", "PCK1", "PDHA1", "LOC642502",
               "PDHA2", "LOC283398", "FH", "SDHD", "OGDH", "SDHB", "IDH3A", "SDHC",
               "IDH2", "IDH1", "OGDHL", "PC", "SDHA", "SUCLG1", "SUCLA2", "SUCLG2")

hyp_obj <- hypeR(signature, genesets, background=2522)

hyp_hmap(hyp_obj, top=60)
```

hyp_show

Convert a hyp object to a reactable table

Description

Convert a hyp object to a reactable table

Usage

```
hyp_show(hyp_obj, simple = FALSE)
```

Arguments

hyp_obj	A hyp object
simple	Use true to only include essential columns

Value

A reactable table

Examples

```
genesets <- msigdb_gsets("Homo sapiens", "C2", "CP:KEGG_LEGACY")

signature <- c("IDH3B", "DLST", "PCK2", "CS", "PDHB", "PCK1", "PDHA1", "LOC642502",
               "PDHA2", "LOC283398", "FH", "SDHD", "OGDH", "SDHB", "IDH3A", "SDHC",
               "IDH2", "IDH1", "OGDHL", "PC", "SDHA", "SUCLG1", "SUCLA2", "SUCLG2")

hyp_obj <- hypeR(signature, genesets, background=2522)

hyp_show(hyp_obj)
```

hyp_to_excel

Export hyp/multihyp object to excel

Description

Export hyp/multihyp object to excel

Usage

```
hyp_to_excel(hyp_obj, file_path, cols = NULL, versioning = TRUE)
```

Arguments

hyp_obj	A hyp or multihyp object
file_path	A file path
cols	Dataframe columns to include
versioning	Add sheet with versioning information

Examples

```
genesets <- msigdb_gsets("Homo sapiens", "C2", "CP:KEGG_LEGACY")

signature <- c("IDH3B", "DLST", "PCK2", "CS", "PDHB", "PCK1", "PDHA1", "LOC642502",
               "PDHA2", "LOC283398", "FH", "SDHD", "OGDH", "SDHB", "IDH3A", "SDHC",
               "IDH2", "IDH1", "OGDHL", "PC", "SDHA", "SUCLG1", "SUCLA2", "SUCLG2")

hyp_obj <- hypeR(signature, genesets, background=2522)

hyp_to_excel(hyp_obj, file_path="pathways.xlsx")
```

hyp_to_graph	<i>Convert a hyp object to an igraph object</i>
--------------	---

Description

Convert a hyp object to an igraph object

Usage

```
hyp_to_graph(hyp_obj)
```

Arguments

hyp_obj A hyp object

Value

An igraph object

Examples

```
genesets <- hyperdb_rgsets("REACTOME", "70.0")

signature <- c("IDH3B", "DLST", "PCK2", "CS", "PDHB", "PCK1", "PDHA1", "LOC642502",
               "PDHA2", "LOC283398", "FH", "SDHD", "OGDH", "SDHB", "IDH3A", "SDHC",
               "IDH2", "IDH1", "OGDHL", "PC", "SDHA", "SUCLG1", "SUCLA2", "SUCLG2")

hyp_obj <- hyper(signature, genesets, background=2522)

ig <- hyp_to_graph(hyp_obj)
```

hyp_to_rmd	<i>Export hyp object to rmarkdown</i>
------------	---------------------------------------

Description

Export hyp object to rmarkdown

Usage

```

hyp_to_rmd(
  hyp_obj,
  file_path,
  title = "Workflow Report",
  subtitle = "",
  author = "",
  header = "Results",
  versioning = TRUE,
  show_dots = TRUE,
  show_emaps = TRUE,
  show_hmaps = FALSE,
  show_tables = TRUE,
  hyp_dots_args = list(top = 15, val = "fdr"),
  hyp_emap_args = list(top = 25, val = "fdr", similarity_metric = "jaccard_similarity",
    similarity_cutoff = 0.2),
  hyp_hmap_args = list(top = 25, val = "fdr"),
  custom_rmd_config = NULL,
  custom_pre_content = NULL,
  custom_post_content = NULL,
  session_info = FALSE
)

```

Arguments

hyp_obj	A hyp object, multihyp object, or list of multihyp objects
file_path	A file path
title	Title of markdown report
subtitle	Subtitle of markdown report
author	Authors of markdown report
header	Header name of tabset section
versioning	Add versioning information
show_dots	Option to show dots plots in tabs
show_emaps	Option to show enrichment maps in tabs
show_hmaps	Option to show hierarchy maps in tabs
show_tables	Option to show table in tabs
hyp_dots_args	A list of keyword arguments passed to hyp_dots
hyp_emap_args	A list of keyword arguments passed to hyp_emap
hyp_hmap_args	A list of keyword arguments passed to hyp_hmap
custom_rmd_config	Replace configuration section of markdown report
custom_pre_content	Insert custom content before tabset section

custom_post_content Insert custom content after tabset section

session_info Use true to include session info

hyp_to_table *Export hyp/multihyp object to table*

Description

Export hyp/multihyp object to table

Usage

```
hyp_to_table(hyp_obj, file_path, sep = "\t", cols = NULL, versioning = TRUE)
```

Arguments

hyp_obj A hyp or multihyp object

file_path A file path for hyp objects and directory for multihyp objects

sep The field separator string

cols Dataframe columns to include

versioning Add header with versioning information

Examples

```
genesets <- msigdb_gsets("Homo sapiens", "C2", "CP:KEGG_LEGACY")

signature <- c("IDH3B", "DLST", "PCK2", "CS", "PDHB", "PCK1", "PDHA1", "LOC642502",
               "PDHA2", "LOC283398", "FH", "SDHD", "OGDH", "SDHB", "IDH3A", "SDHC",
               "IDH2", "IDH1", "OGDHL", "PC", "SDHA", "SUCLG1", "SUCLA2", "SUCLG2")

hyp_obj <- hypeR(signature, genesets, background=2522)

hyp_to_table(hyp_obj, file_path="pathways.txt")
```

limma	<i>Differential Expression</i>
-------	--------------------------------

Description

A differential expression table

Usage

```
limma
```

Format

A data frame

msigdb_available	<i>Get msigdb available genesets</i>
------------------	--------------------------------------

Description

Get msigdb available genesets

Usage

```
msigdb_available(species = "Homo sapiens")
```

Arguments

species	A species to determine gene symbols (refer to ?msigdb::msigdb for available species)
---------	--

Value

A dataframe of available genesets

Examples

```
msigdb_available("Homo sapiens")
```

msigdb_check_species	<i>Check if species is available</i>
----------------------	--------------------------------------

Description

Check if species is available

Usage

```
msigdb_check_species(species = "")
```

Arguments

species A species

Examples

```
## Not run:
msigdb_check_species("Homo sapiens")

## End(Not run)
```

msigdb_download	<i>Download data from msigdb in the form of a named list</i>
-----------------	--

Description

Download data from msigdb in the form of a named list

Usage

```
msigdb_download(species, collection, subcollection = NULL)
```

Arguments

species A species to determine gene symbols (refer to ?msigdbr::msigdbr for available species)

collection Geneset collection (refer to ?msigdbr::msigdbr_collections for available categories)

subcollection Geneset subcollection (refer to ?msigdbr::msigdbr_collections for available sub-categories)

Value

A list of genesets

Examples

```
HALLMARK_HUMAN <- msigdb_download("Homo sapiens", "H")
HALLMARK_MOUSE <- msigdb_download("Mus musculus", "MH")
```

msigdb_gsets	<i>Download data from msigdb in the form of a gsets object</i>
--------------	--

Description

Download data from msigdb in the form of a gsets object

Usage

```
msigdb_gsets(species, collection, subcollection = NULL, clean = FALSE)
```

Arguments

species	A species to determine gene symbols (refer to ?msigdb::msigdb_species for available species)
collection	Geneset collection (refer to ?msigdb::msigdb_collections for available categories)
subcollection	Geneset subcollection (refer to ?msigdb::msigdb_collections for available sub-categories)
clean	Use true to clean labels of genesets

Value

A gsets object

Examples

```
HALLMARK_HUMAN <- msigdb_gsets("Homo sapiens", "H")
HALLMARK_MOUSE <- msigdb_gsets("Mus musculus", "MH")
```

msigdb_info	<i>Print msigdb gsets information</i>
-------------	---------------------------------------

Description

Print msigdb gsets information

Usage

```
msigdb_info()
```

Examples

```
msigdb_info()
```

msigdb_species	<i>Get msigdb available species</i>
----------------	-------------------------------------

Description

Get msigdb available species

Usage

```
msigdb_species()
```

Value

A character vector of species

Examples

```
msigdb_species()
```

msigdb_version	<i>Get msigdb package version number</i>
----------------	--

Description

Get msigdb package version number

Usage

msigdb_version()

Value

Version number

Examples

msigdb_version()

multihyp	<i>A multihyp object</i>
----------	--------------------------

Description

A multihyp object
A multihyp object

See Also

hyp

Public fields

data A list of hyp objects

Methods

Public methods:

- `multihyp$new()`
- `multihyp$print()`
- `multihyp$as.list()`
- `multihyp$clone()`

Method new(): Create a multihyp object

Usage:

```
multihyp$new(data)
```

Arguments:

data A list of hyp objects

Returns: A new multihyp object

Method print(): Print multihyp object

Usage:

```
multihyp$print()
```

Returns: NULL

Method as.list(): Print multihyp object

Usage:

```
multihyp$as.list()
```

Returns: A list of hyp objects as dataframes

Method clone(): The objects of this class are cloneable with this method.

Usage:

```
multihyp$clone(deep = FALSE)
```

Arguments:

deep Whether to make a deep clone.

Examples

```
data <- data.frame(replicate(5,sample(0:1,10,rep=TRUE)))
args <- list("arg_1"=1, "arg_2"=2, "arg_3"=3)
hyp_obj <- hyp$new(data, args=args)
data <- list("hyp_1"=hyp_obj, "hyp_2"=hyp_obj,"hyp_3"=hyp_obj)
multihyp_obj <- multihyp$new(data)
```

pvector

A push/pop capable vector

Description

A push/pop capable vector

A push/pop capable vector

Public fields

values A vector of values

Methods

Public methods:

- `pvector$new()`
- `pvector$print()`
- `pvector$length()`
- `pvector$pop()`
- `pvector$push()`
- `pvector$clone()`

Method `new()`: Create a pvector

Usage:

`pvector$new(values = c())`

Arguments:

`values` A vector of values

Returns: A new pvector

Method `print()`: Print pvector

Usage:

`pvector$print()`

Returns: NULL

Method `length()`: Get length of pvector

Usage:

`pvector$length()`

Returns: An integer

Method `pop()`: Pop vector

Usage:

`pvector$pop()`

Returns: Popped value

Method `push()`: Push values

Usage:

`pvector$push(pushed.values)`

Arguments:

`pushed.values` A vector of values

Returns: NULL

Method `clone()`: The objects of this class are cloneable with this method.

Usage:

`pvector$clone(deep = FALSE)`

Arguments:

`deep` Whether to make a deep clone.

rctbl_build	<i>Reactable builder for hyp or mhyp objects</i>
-------------	--

Description

Reactable builder for hyp or mhyp objects

Usage

```
rctbl_build(obj, ...)
```

Arguments

obj	A hyp or multihyp object
...	Arguments passed to table generators

Examples

```
genesets <- msigdb_gsets("Homo sapiens", "C2", "CP:KEGG_LEGACY")$genesets[1:5]

experiment <- list("S1"=c("IDH3B", "DLST", "PCK2", "CS", "PDHB", "PCK1", "PDHA1", "LOC642502"),
                  "S2"=c("PDHA2", "LOC283398", "FH", "SDHD", "OGDH", "SDHB", "IDH3A", "SDHC"))

mhyp_obj <- hypeR(experiment, genesets, background=2522)

rctbl_build(mhyp_obj)
```

rctbl_hyp	<i>Reactable table for hyp objects</i>
-----------	--

Description

Reactable table for hyp objects

Usage

```
rctbl_hyp(
  hyp,
  type = c("inner", "outer"),
  show_emaps = FALSE,
  show_hmaps = FALSE,
  hyp_emap_args = list(top = 25, val = "fdr"),
  hyp_hmap_args = list(top = 25, val = "fdr")
)
```

Arguments

hyp	A hyp object
type	Use style class for outer or inner tables
show_emaps	Option to show enrichment maps in tabs
show_hmaps	Option to show hiearchy maps in tabs
hyp_emap_args	A list of keyword arguments passed to hyp_emap
hyp_hmap_args	A list of keyword arguments passed to hyp_hmap

Examples

```
genesets <- msigdb_gsets("Homo sapiens", "C2", "CP:KEGG_LEGACY")$genesets[1:5]

signature <- c("IDH3B", "DLST", "PCK2", "CS", "PDHB", "PCK1", "PDHA1", "LOC642502",
               "PDHA2", "LOC283398", "FH", "SDHD", "OGDH", "SDHB", "IDH3A", "SDHC",
               "IDH2", "IDH1", "OGDHL", "PC", "SDHA", "SUCLG1", "SUCLA2", "SUCLG2")

hyp_obj <- hypeR(signature, genesets, background=2522)

rctbl_hyp(hyp_obj)
```

rctbl_mhyp	<i>Reactable table for multihyp objects</i>
------------	---

Description

Reactable table for multihyp objects

Usage

```
rctbl_mhyp(
  mhyp,
  show_emaps = FALSE,
  show_hmaps = FALSE,
  hyp_emap_args = list(top = 25, val = "fdr"),
  hyp_hmap_args = list(top = 25, val = "fdr")
)
```

Arguments

mhyp	A multihyp object
show_emaps	Option to show enrichment maps in tabs
show_hmaps	Option to show hiearchy maps in tabs
hyp_emap_args	A list of keyword arguments passed to hyp_emap
hyp_hmap_args	A list of keyword arguments passed to hyp_hmap

Examples

```
genesets <- msigdb_gsets("Homo sapiens", "C2", "CP:KEGG_LEGACY")$genesets[1:5]

experiment <- list("S1"=c("IDH3B", "DLST", "PCK2", "CS", "PDHB", "PCK1", "PDHA1", "LOC642502"),
                  "S2"=c("PDHA2", "LOC283398", "FH", "SDHD", "OGDH", "SDHB", "IDH3A", "SDHC"))

mhyp_obj <- hypeR(experiment, genesets, background=2522)

rctbl_mhyp(mhyp_obj)
```

rgsets	<i>A relational genesets object</i>
--------	-------------------------------------

Description

A relational genesets object

A relational genesets object

See Also

gsets

Public fields

genesets A list of genesets where list names refers to geneset labels and values are geneset members represented as a vector

nodes A data frame of labeled nodes

edges A data frame of directed edges

name A character vector describing source of genesets

version A character vector describing versioning

Methods**Public methods:**

- `rgsets$new()`
- `rgsets$print()`
- `rgsets$info()`
- `rgsets$reduce()`
- `rgsets$subset()`
- `rgsets$clone()`

Method `new()`: Create a rgsets object

Usage:

```
rgsets$new(
  genesets,
  nodes,
  edges,
  name = "Custom",
  version = "",
  quiet = FALSE
)
```

Arguments:

genesets A list of genesets where list names refers to geneset labels and values are geneset members represented as a vector

nodes A data frame of labeled nodes

edges A data frame of directed edges

name A character vector describing source of genesets

version A character vector describing versioning

quiet Use true to silence warnings

Returns: A new rgsets object

Method `print()`: Print relational genesets information

Usage:

```
rgsets$print()
```

Returns: NULL

Method `info()`: Returns versioning information

Usage:

```
rgsets$info()
```

Returns: A character vector with name and version

Method `reduce()`: Reduces genesets to a background distribution of symbols

Usage:

```
rgsets$reduce(background)
```

Arguments:

background A character vector of symbols

Returns: A rgsets object

Method `subset()`: Subsets genesets on a character vector of labels

Usage:

```
rgsets$subset(labels)
```

Arguments:

labels A character vector of genesets

Returns: A rgsets object

Method `clone()`: The objects of this class are cloneable with this method.

Usage:

```
rgsets$clone(deep = FALSE)
```

Arguments:

deep Whether to make a deep clone.

wgcna

Co-expression Modules

Description

A list of co-expression modules

Usage

```
wgcna
```

Format

A nested list of character vectors

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