Package 'simplifyEnrichment'

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Description A new method (binary cut) is proposed to effectively cluster GO terms into groups from the semantic similarity matrix. Summaries of GO terms in each cluster are visualized by word clouds.
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all_clustering_methods

```
all_clustering_methods
```

All clustering methods

Description

All clustering methods

Usage

```
all_clustering_methods()
```

Details

The default clustering methods are:

```
kmeans see cluster_by_kmeans.
dynamicTreeCut see cluster_by_dynamicTreeCut.
mclust see cluster_by_mclust.
apcluster see cluster_by_apcluster.
hdbscan see cluster_by_hdbscan.
fast_greedy see cluster_by_igraph.
leading_eigen see cluster_by_igraph.
louvain see cluster_by_igraph.
walktrap see cluster_by_igraph.
MCL see cluster_by_igraph.
binary_cut see binary_cut.
```

Value

A vector of method names.

See Also

New methods can be added by register_clustering_methods.

```
all_clustering_methods()
```

4 cluster_by_apcluster

binary_cut	Cluster functional terms by recursively binary cutting the similarity matrix

Description

Cluster functional terms by recursively binary cutting the similarity matrix

Usage

```
binary_cut(mat, value_fun = median, partition_fun = partition_by_pam,
    cutoff = 0.85, cache = FALSE, try_all_partition_fun = FALSE)
```

Arguments

mat A similarity matrix.

value_fun Value function to calculate the score for each node in the dendrogram.

partition_fun A function to split each node into two groups. Pre-defined functions in this pack-

age are partition_by_kmeanspp, partition_by_pam and partition_by_hclust.

cutoff The cutoff for splitting the dendrogram.

cache Whether the dendrogram should be cached. Internally used.

try_all_partition_fun

Different partition_fun gives different clusterings. If the vaule of try_all_partition_fun is set to TRUE, the similarity matrix is clustered by three partitioning method: partition_by_pam, partition_by_kmeanspp and partition_by_hclust. The clustering with the highest difference score is finally selected as the final clus-

tering.

Value

A vector of cluster labels (in numeric).

Examples

cluster_by_apcluster Cluster similarity matrix by apcluster

Description

Cluster similarity matrix by apcluster

Usage

```
cluster_by_apcluster(mat, s = apcluster::negDistMat(r = 2), ...)
```

Arguments

```
mat The similarity matrix.

s Passed to the s argument in apcluster.

Other arguments passed to apcluster.
```

Value

A vector of cluster labels (in numeric).

Examples

```
\label{eq:continuous_problem} \mbox{\ensuremath{\mbox{\sc H}}} \mbox{\ensuremath{\mbox{\mbox{\sc H}}}} \mbox{\ensuremath{\mbox{\sc H}}} \mbox{\ensuremath{\mbox{\sc
```

```
cluster_by_dynamicTreeCut
```

Cluster similarity matrix by dynamicTreeCut

Description

Cluster similarity matrix by dynamicTreeCut

Usage

```
cluster_by_dynamicTreeCut(mat, minClusterSize = 5, ...)
```

Arguments

```
mat The similarity matrix.

minClusterSize Minimal number of objects in a cluster. Pass to cutreeDynamic.

Other arguments passed to cutreeDynamic.
```

Value

A vector of cluster labels (in numeric).

```
\label{eq:continuous_problem} \mbox{\ensuremath{\texttt{T}} There is no example} \\ \mbox{\ensuremath{\texttt{NULL}}}
```

6 cluster_by_igraph

cluster_by_hdbscan Clust

Cluster similarity matrix by hdbscan

Description

Cluster similarity matrix by hdbscan

Usage

```
cluster_by_hdbscan(mat, minPts = 5, ...)
```

Arguments

mat The similarity matrix.

minPts Passed to the minPts argument in hdbscan.

... Other arguments passed to hdbscan.

Value

A vector of cluster labels (in numeric).

Examples

```
\label{eq:continuous_problem} \mbox{\ensuremath{\mbox{\sc H}}} \mbox{\ensuremath{\mbox{\mbox{\sc H}}}} \mbox{\ensuremath{\mbox{\sc H}}} \mbox{\ensuremath{\mbox{\sc
```

cluster_by_igraph

Cluster similarity matrix by graph community detection methods

Description

Cluster similarity matrix by graph community detection methods

Usage

```
cluster_by_igraph(mat,
    method = c("cluster_fast_greedy",
    "cluster_leading_eigen",
    "cluster_louvain",
    "cluster_walktrap"),
    ...)
```

Arguments

mat The similarity matrix.

method The community detection method.

Other arguments passed to the corresponding community detection function, see

Details.

cluster_by_kmeans 7

Details

The symmetric similarity matrix can be treated as an adjacency matrix and constructed as a graph/network with the similarity values as the weight of hte edges. Thus, clustering the similarity matrix can be treated as detecting clusters/modules/communities from the graph.

Four methods implemented in igraph package can be used here:

```
cluster_fast_greedy uses cluster_fast_greedy.
cluster_leading_eigen uses cluster_leading_eigen.
cluster_louvain uses cluster_louvain.
cluster_walktrap uses cluster_walktrap.
```

Value

A vector of cluster labels (in numeric).

Examples

```
\hbox{\tt\# There is no example}\\ \hbox{\tt NULL}
```

cluster_by_kmeans

Cluster similarity matrix by k-means clustering

Description

Cluster similarity matrix by k-means clustering

Usage

```
cluster_by_kmeans(mat, \max_k = \max(2, \min(\text{round}(\text{nrow}(\text{mat})/5), 100)), \ldots)
```

Arguments

```
mat The similarity matrix.

max_k maximal k for k-means clustering.

Other arguments passed to kmeans.
```

Details

The best number of k for k-means clustering is identified according to the "elbow" or "knee" method on the distribution of within-cluster sum of squares (WSS) at each k.

Value

A vector of cluster labels (in numeric).

```
# There is no example NULL
```

8 cluster_by_mclust

cluster_by_MCL

Cluster similarity matrix by MCL

Description

Cluster similarity matrix by MCL

Usage

```
cluster_by_MCL(mat, addLoops = FALSE, ...)
```

Arguments

mat The similarity matrix.

addLoops Passed to the addLoops argument in mcl.

... Other arguments passed to mcl.

Value

A vector of cluster labels (in numeric).

Examples

```
\# There is no example NULL
```

cluster_by_mclust

Cluster similarity matrix by mclust

Description

Cluster similarity matrix by mclust

Usage

```
cluster_by_mclust(mat, G = seq_len(max(2, min(round(nrow(mat)/5), 100))), ...)
```

Arguments

mat The similarity matrix.

G Passed to the G argument in Mclust.
... Other arguments passed to Mclust.

Value

A vector of cluster labels (in numeric).

cluster_terms 9

Examples

```
# There is no example NULL
```

cluster_terms

Cluster functional terms

Description

Cluster functional terms

Usage

```
cluster_terms(mat, method = "binary_cut", control = list(), catch_error = FALSE,
    verbose = TRUE)
```

Arguments

mat A similarity matrix.

method Method for clustering the matrix.

control A list of parameters passed to the corresponding clustering function.

catch_error Internally used.

verbose Whether to print messages.

Details

The following methods are the default:

```
kmeans see cluster_by_kmeans.
dynamicTreeCut see cluster_by_dynamicTreeCut.
mclust see cluster_by_mclust.
apcluster see cluster_by_apcluster.
hdbscan see cluster_by_hdbscan.
fast_greedy see cluster_by_igraph.
leading_eigen see cluster_by_igraph.
louvain see cluster_by_igraph.
walktrap see cluster_by_igraph.
MCL see cluster_by_MCL.
binary_cut see binary_cut.
```

Also the user-defined methods in all_clustering_methods can be used here.

New clustering methods can be registered by register_clustering_methods.

Please note it is better to directly call cluster_terms for clustering while not the individual cluster_by_* functions because cluster_terms does additional cluster label adjustment.

10 cmp_make_clusters

Value

A numeric vector of cluster labels (in numeric).

If catch_error is set to TRUE and if the clustering produces an error, the function returns a try-error object.

Examples

```
# There is no example NULL
```

cmp_make_clusters

Apply various clustering methods

Description

Apply various clustering methods

Usage

```
cmp_make_clusters(mat, method = setdiff(all_clustering_methods(), "mclust"),
    verbose = TRUE)
```

Arguments

mat The similarity matrix.

method Which methods to compare. All available methods are in all_clustering_methods.

A value of all takes all available methods. By default mclust is excluded be-

cause its long runtime.

verbose Whether to print messages.

Details

The function compares following default clustering methods by default:

```
kmeans see cluster_by_kmeans.
dynamicTreeCut see cluster_by_dynamicTreeCut.
mclust see cluster_by_mclust. By default it is not included.
apcluster see cluster_by_apcluster.
hdbscan see cluster_by_hdbscan.
fast_greedy see cluster_by_igraph.
leading_eigen see cluster_by_igraph.
louvain see cluster_by_igraph.
walktrap see cluster_by_igraph.
MCL see cluster_by_MCL.
binary_cut see binary_cut.
```

Also the user-defined methods in all_clustering_methods are also compared.

cmp_make_plot 11

Value

A list of cluster label vectors for different clustering methods.

Examples

cmp_make_plot

Make plots for comparing clustering methods

Description

Make plots for comparing clustering methods

Usage

```
cmp_make_plot(mat, clt, plot_type = c("mixed", "heatmap"), nrow = 3)
```

Arguments

mat A similarity matrix.

clt A list of clusterings from cmp_make_clusters.

plot_type What type of plots to make. See Details.

nrow Number of rows of the layout when plot_type is set to heatmap.

Details

If plot_type is the default value mixed, a figure with three panels generated:

- A heatmap of the similarity matrix with different classifications as row annotations.
- A heatmap of the pair-wise concordance of the classifications of every two clustering methods.
- Barplots of the difference scores for each method (calculated by difference_score), the number of clusters (total clusters and the clusters with size >= 5) and the mean similarity of the terms that are in the same clusters.

If plot_type is heatmap. There are heatmaps for the similarity matrix under clusterings from different methods. The last panel is a table with the number of clusters under different clusterings.

Value

No value is returned.

Examples

compare_clustering_methods

Compare clustering methods

Description

Compare clustering methods

Usage

```
compare_clustering_methods(mat, method = setdiff(all_clustering_methods(), "mclust"),
    plot_type = c("mixed", "heatmap"), nrow = 3, verbose = TRUE)
```

Arguments

mat The similarity matrix.

method Which methods to compare. All available methods are in all_clustering_methods.

A value of all takes all available methods. By default mclust is excluded be-

cause its long runtime.

plot_type See explanation in cmp_make_plot.

nrow Number of rows of the layout when plot_type is set to heatmap.

verbose Whether to print messages.

Details

The function compares following clustering methods by default:

```
kmeans see cluster_by_kmeans.
dynamicTreeCut see cluster_by_dynamicTreeCut.
mclust see cluster_by_mclust. By default it is not included.
apcluster see cluster_by_apcluster.
hdbscan see cluster_by_hdbscan.
fast_greedy see cluster_by_igraph.
leading_eigen see cluster_by_igraph.
louvain see cluster_by_igraph.
walktrap see cluster_by_igraph.
MCL see cluster_by_MCL.
```

count_word 13

```
binary_cut see binary_cut.
```

This function is basically a wrapper function. It calls the following two functions:

- cmp_make_clusters: applies clustering with different methods.
- cmp_make_plot: makes the plots.

Value

No value is returned.

Examples

count_word

Calculate word frequency

Description

Calculate word frequency

Usage

```
count_word(go_id, term = NULL, exclude_words = NULL)
```

Arguments

go_id A vector of GO IDs.

term The corresponding names or description of terms if the input are not GO terms.

exclude_words The words that should be excluded.

Details

The input can be simply set with a vector of GO id to go_id argument that the GO names are automatically extracted. If the input are not GO terms, users need to provide a vector of long names/descriptions by term argument.

If the input is GO id, the following words are excluded: c("via", "protein", "factor", "side", "type", "specific") They are analyzed by simplifyEnrichment:::all_GO_word_count().

The text preprocessing followings the instructions from http://www.sthda.com/english/wiki/word-cloud-generator-in-r-one-killer-function-to-do-everything-you-need.

Value

A data frame with words and frequencies.

14 dend_node_apply

Examples

```
go_id = random_GO(500)
head(count_word(go_id))
```

dend_node_apply

Apply functions on every node in a dendrogram

Description

Apply functions on every node in a dendrogram

Usage

```
dend_node_apply(dend, fun)
```

Arguments

dend A dendrogram.

fun A self-defined function.

Details

The function returns a vector or a list as the same length as the number of nodes in the dendrogram.

The self-defined function can have one single argument which is the sub-dendrogram at a certain node. E.g. to get the number of members at every node:

```
dend_node_apply(dend, function(d) attr(d, "members"))
```

The self-defined function can have a second argument, which is the index of current sub-dendrogram in the complete dendrogram. E.g. dend[[1]] is the first child node of the complete dendrogram and dend[[c(1,2)]] is the second child node of dend[[1]], et al. This makes that at a certain node, it is possible to get information of its child nodes and parent nodes.

```
dend_node_apply(dend, function(d, index) {
    dend[[c(index, 1)]] # is the first child node of d, or simply d[[1]]
    dend[[index[-length(index)]]] # is the parent node of d
    ...
})
```

Note for the top node, the value of index is NULL.

Value

A vector or a list, depends on whether fun returns a scalar or more complex values.

difference_score 15

Examples

```
mat = matrix(rnorm(100), 10)
dend = as.dendrogram(hclust(dist(mat)))
# number of members on every node
dend_node_apply(dend, function(d) attr(d, "members"))
# the depth on every node
dend_node_apply(dend, function(d, index) length(index))
```

 ${\tt difference_score}$

Difference score

Description

Difference score

Usage

```
difference_score(mat, cl)
```

Arguments

mat The similarity matrix.

cl Cluster labels.

Details

This function measures the different between the similarity values for the terms that belong to the same clusters and in different clusters. The difference score is the Kolmogorov-Smirnov statistic between the two distributions.

Value

A numeric scalar.

16 edit_node

DO_similarity

Calculate Disease Ontology (DO) semantic similarity matrix

Description

Calculate Disease Ontology (DO) semantic similarity matrix

Usage

```
DO_similarity(do_id, measure = "Rel")
```

Arguments

do_id A vector of DO IDs.

measure Semantic measurement for the DO similarity, pass to doSim.

Details

This function is basically a wrapper on doSim.

Value

A symmetric matrix.

Examples

```
require(DOSE)
do_id = random_DO(10)
DO_similarity(do_id)
```

edit_node

Modify nodes in a dendrogram

Description

Modify nodes in a dendrogram

Usage

```
edit_node(dend, fun = function(d, index) d)
```

Arguments

dend A dendrogram.

fun A self-defined function.

GO_similarity 17

Details

if fun only has one argument, it is basically the same as dendrapply, but it can have a second argument which is the index of the node in the dendrogram, which makes it possible to get information of child nodes and parent nodes for a specific node.

As an example, we first assign random values to every node in the dendrogram:

```
mat = matrix(rnorm(100), 10)
dend = as.dendrogram(hclust(dist(mat)))
dend = edit_node(dend, function(d) {attr(d, 'score') = runif(1); d})
```

Then for every node, we take the maximal absolute difference to all its child nodes and parent node as the attribute abs_diff

```
dend = edit_node(dend, function(d, index) {
    n = length(index)
    s = attr(d, "score")
    if(is.null(index)) { # d is the top node
        s_children = sapply(d, function(x) attr(x, "score"))
        s_parent = NULL
    } else if(is.leaf(d)) { # d is the leaf
        s_{children} = NULL
        s_parent = attr(dend[[index[-n]]], "score")
    } else {
        s_children = sapply(d, function(x) attr(x, "score"))
        s_parent = attr(dend[[index[-n]]], "score")
    abs_diff = max(abs(s - c(s_children, s_parent)))
    attr(d, "abs_diff") = abs_diff
    return(d)
})
```

Value

A dendrogram object.

Examples

```
# There is no example NULL
```

GO_similarity

Calculate Gene Ontology (GO) semantic similarity matrix

Description

Calculate Gene Ontology (GO) semantic similarity matrix

Usage

```
GO_similarity(go_id, ont, db = 'org.Hs.eg.db', measure = "Rel")
```

18 guess_ont

Arguments

go_id A vector of GO IDs.

ont GO ontology. Value should be one of "BP", "CC" or "MF". If it is not specified,

the function automatically identifies it by random sampling 10 IDs from go_id

(see guess_ont).

db Annotation database. It should be from https://bioconductor.org/packages/

3.10/BiocViews.html#___OrgDb

measure Semantic measurement for the GO similarity, pass to termSim.

Details

This function is basically a wrapper on ${\tt termSim}$.

Value

A symmetric matrix.

Examples

```
go_id = random_GO(100)
mat = GO_similarity(go_id)
```

guess_ont

Guess the ontology of the input GO IDs

Description

Guess the ontology of the input GO IDs

Usage

```
guess_ont(go_id, db = 'org.Hs.eg.db')
```

Arguments

go_id A vector of GO IDs.

db Annotation database. It should be from https://bioconductor.org/packages/

3.10/BiocViews.html#___OrgDb

Details

10 GO IDs are randomly sampled and checked.

Value

A single character scalar of "BP", "CC" or "MF".

If there are more than one ontologies detected. It returns NULL.

Examples

```
go_id = random_GO(100)
guess_ont(go_id)
```

heightDetails.word_cloud

Height for word_cloud grob

Description

Height for word_cloud grob

Usage

```
## S3 method for class 'word_cloud'
heightDetails(x)
```

Arguments

Χ

The word_cloud grob returned by word_cloud_grob.

Value

A unit object.

Examples

```
\# There is no example NULL
```

ht_clusters

Visualize the similarity matrix and the clustering

Description

Visualize the similarity matrix and the clustering

Usage

```
ht_clusters(mat, cl, dend = NULL, col = c("white", "red"),
    draw_word_cloud = is_GO_id(rownames(mat)[1]) || !is.null(term),
    term = NULL, min_term = 5,
    order_by_size = FALSE, cluster_slices = FALSE,
    exclude_words = character(0), max_words = 10,
    word_cloud_grob_param = list(), fontsize_range = c(4, 16),
    column_title = NULL, ht_list = NULL, use_raster = TRUE, ...)
```

20 ht_clusters

Arguments

mat

cl Cluster labels inferred from the similarity matrix, e.g. from cluster_terms or binary_cut.

dend Used internally.

col A vector of colors that map from 0 to the 95th percentile of the similarity

values.

A similarity matrix.

draw_word_cloud

Whether to draw the word clouds.

term The full name or the description of the corresponding GO IDs.

min_term Minimal number of functional terms in a cluster. All the clusters with size less

than min_term are all merged into one separated cluster in the heatmap.

order_by_size Whether to reorder clusters by their sizes. The cluster that is merged from small

clusters (size < min_term) is always put to the bottom of the heatmap.

cluster_slices Whether to cluster slices.

exclude_words Words that are excluded in the word cloud.

max_words Maximal number of words visualized in the word cloud.

word_cloud_grob_param

A list of graphic parameters passed to word_cloud_grob.

fontsize_range The range of the font size. The value should be a numeric vector with length

two. The minimal font size is mapped to word frequency value of 1 and the maximal font size is mapped to the maximal word frequency. The font size

interlopation is linear.

ht_list A list of additional heatmaps added to the left of the similarity heatmap.

use_raster Whether to write the heatmap as a raster image.

... Other arguments passed to draw, HeatmapList-method.

Value

A HeatmapList-class object.

partition_by_hclust 21

Description

Partition by hclust

Usage

```
partition_by_hclust(mat)
```

Arguments

mat

The similarity matrix.

Details

The "ward.D2" clusering method was used.

This function is used to set to the partition_fun argument in binary_cut.

Examples

```
\label{eq:continuous_problem} \mbox{\ensuremath{\mbox{\sc H}}} \mbox{\ensuremath{\mbox{\mbox{\sc H}}}} \mbox{\ensuremath{\mbox{\sc H}}} \mbox{\ensuremath{\mbox{\sc
```

partition_by_kmeans

Partition by kmeans

Description

Partition by kmeans

Usage

```
partition_by_kmeans(mat, n_repeats = 10)
```

Arguments

mat The similarity matrix.

 $n_repeats$ Number of repeated runs of k-means.

Details

Since k-means clustering brings randomness, this function performs k-means clustering several times and uses the final consensus partitioning.

This function is used to set to the partition_fun argument in binary_cut.

22 partition_by_pam

Examples

```
\label{eq:total_problem} \mbox{\ensuremath{\texttt{#}} There is no example} \\ \mbox{\ensuremath{\texttt{NULL}}}
```

```
partition_by_kmeanspp Partition by kmeans++
```

Description

Partition by kmeans++

Usage

```
partition_by_kmeanspp(mat)
```

Arguments

mat

The similarity matrix.

Details

This function is used to set to the partition_fun argument in binary_cut.

Examples

```
\label{eq:total_problem} \mbox{\ensuremath{\texttt{#}} There is no example} \\ \mbox{\ensuremath{\texttt{NULL}}} \mbox{\ensuremath{\texttt{NULL}}} \mbox{\ensuremath{\texttt{NULL}}} \mbox{\ensuremath{\texttt{*}}} \mbox{\ensure
```

partition_by_pam

Partition by PAM

Description

Partition by PAM

Usage

```
partition_by_pam(mat)
```

Arguments

mat

The similarity matrix.

Details

The clustering is performed by pam with setting pamonce argument to 5.

This function is used to set to the partition_fun argument in binary_cut.

plot_binary_cut 23

Examples

```
# There is no example NULL
```

plot_binary_cut

Visualize the process of binary cut

Description

Visualize the process of binary cut

Usage

```
plot_binary_cut(mat, value_fun = median, cutoff = 0.85,
    partition_fun = partition_by_pam, dend = NULL, dend_width = unit(3, "cm"),
    depth = NULL, show_heatmap_legend = TRUE, ...)
```

Arguments

mat The similarity matrix.

value_fun Value function to calculate the score for each node in the dendrogram.

cutoff The cutoff for splitting the dendrogram.

partition_fun A function to split each node into two groups. Pre-defined functions in this pack-

age are partition_by_kmeanspp, partition_by_pam and partition_by_hclust.

dend A dendrogram object, used internally.depth Depth of the recursive binary cut process.

dend_width Width of the dendrogram.

show_heatmap_legend

Whether to show the heatmap legend.

... Other arguments.

Details

After the functions which performs clustering are executed, such as simplifyGO or binary_cut, the dendrogram is temporarily saved and plot_binary_cut directly uses this dendrogram. So, if the partition function brings randomness, it makes sure the clustering is the same as the one made by e.g. simplifyGO.

24 random_GO

random_DO

Generate random Disease Ontology (DO) IDs

Description

Generate random Disease Ontology (DO) IDs

Usage

```
random_DO(n)
```

Arguments

n

Number of DO IDs.

Details

DO. db package should be installed.

Value

A vector of DO IDs.

Examples

```
random_DO(100)
```

random_G0

Generate random GO IDs

Description

Generate random GO IDs

Usage

```
random_GO(n, ont = "BP", db = 'org.Hs.eg.db')
```

Arguments

n Number of GO IDs.

ont GO ontology. Value should be one of "BP", "CC" or "MF".

db Annotation database. It should be from https://bioconductor.org/packages/

3.10/BiocViews.html#___OrgDb

Value

A vector of GO IDs.

Examples

```
random_G0(100)
```

```
register_clustering_methods
```

Register new clustering methods

Description

Register new clustering methods

Usage

```
register_clustering_methods(...)
```

Arguments

... A named list of clustering functions, see Details.

Details

The user-defined functions should accept at least one argument which is the input matrix. The second optional argument should always be ... so that parameters for the clustering function can be passed by control argument from cluster_terms, simplifyGO or simplifyEnrichment. If users forget to add ..., it is added internally.

Please note, the user-defined function should automatically identify the optimized number of clusters.

The function should return a vector of cluster labels. Internally it is converted to numeric labels.

Value

No value is returned.

```
register_clustering_methods(
# assume there are 5 groups
random = function(mat, ...) sample(5, nrow(mat), replace = TRUE)
)
all_clustering_methods()
remove_clustering_methods("random")
```

```
remove_clustering_methods
```

Remove clustering methods

Description

Remove clustering methods

Usage

```
remove_clustering_methods(method)
```

Arguments

method

A vector of method names.

Value

No value is returned.

Examples

```
# There is no example NULL
```

```
reset_clustering_methods
```

Reset to default clustering methods

Description

Reset to default clustering methods

Usage

```
reset_clustering_methods()
```

Details

The default methods are:

```
kmeans see cluster_by_kmeans.
dynamicTreeCut see cluster_by_dynamicTreeCut.
mclust see cluster_by_mclust.
apcluster see cluster_by_apcluster.
hdbscan see cluster_by_hdbscan.
fast_greedy see cluster_by_igraph.
```

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```
leading_eigen see cluster_by_igraph.
louvain see cluster_by_igraph.
walktrap see cluster_by_igraph.
MCL see cluster_by_MCL.
binary_cut see binary_cut.
```

Value

No value is returned.

Examples

```
all_clustering_methods()
remove_clustering_methods(c("kmeans", "mclust"))
all_clustering_methods()
reset_clustering_methods()
all_clustering_methods()
```

scale_fontsize

Scale font size

Description

Scale font size

Usage

```
scale_fontsize(x, rg = c(1, 30), fs = c(4, 16))
```

Arguments

x A numeric vector.

rg The range.

fs Range of the font size.

Value

A numeric vector.

Detaisl

It is a linear interpolation.

```
x = runif(10, min = 1, max = 20)
# scale x to fontsize 4 to 16.
scale_fontsize(x)
```

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-			
se	lect	cutoff	

Select the cutoff for binary cut

Description

Select the cutoff for binary cut

Usage

```
select\_cutoff(mat, cutoff = seq(0.6, 0.98, by = 0.01), verbose = TRUE, ...)
```

Arguments

```
mat A similarity matrix.

cutoff A list of cutoffs to test. Note the range of the cutoff values should be inside [0.5, 1].

verbose Whether to print messages.

Pass to binary_cut.
```

Details

Binary cut is applied to each of the cutoff and the clustering results are evaluated by following metrics:

- difference score, calculated by difference_score.
- number of clusters.
- block mean, which is the mean similarity in the blocks in the diagonal of the heatmap.

Examples

simplifyEnrichment

Simplify functional enrichment results

Description

Simplify functional enrichment results

Usage

```
simplifyEnrichment(mat, method = "binary_cut", control = list(),
   plot = TRUE, term = NULL, verbose = TRUE,
   column_title = qq("@{nrow(mat)} terms clustered by '@{method}'"),
   ht_list = NULL, ...)
```

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Arguments

A similarity matrix. mat method Method for clustering the matrix. See cluster_terms. control A list of parameters for controlling the clustering method, passed to cluster_terms. plot Whether to make the heatmap. The full name or the description of the corresponding terms. term column_title Column title for the heatmap. verbose Whether to print messages. ht_list A list of additional heatmaps added to the left of the similarity heatmap.

... Arguments passed to ht_clusters.

Details

The usage is the same as simplifyGO, except you need to manually provide the term names by term argument if you want to draw the word clouds.

Examples

```
# There is no example NULL
```

simplifyG0

Simplify Gene Ontology (GO) enrichment results

Description

Simplify Gene Ontology (GO) enrichment results

Usage

```
simplifyGO(mat, method = "binary_cut", control = list(),
   plot = TRUE, term = NULL, verbose = TRUE,
   column_title = qq("@{nrow(mat)} GO terms clustered by '@{method}'"),
   ht_list = NULL, ...)
```

Arguments

mat A GO similarity matrix. method Method for clustering the matrix. See cluster_terms. control A list of parameters for controlling the clustering method, passed to cluster_terms. Whether to make the heatmap. plot term The full name or the description of the corresponding GO IDs. The values are automatically extracted if it is not provided. Column title for the heatmap. column_title verbose Whether to print messages. ht_list A list of additional heatmaps added to the left of the similarity heatmap. Arguments passed to ht_clusters. . . .

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Details

This is basically a wrapper function that it first runs cluster_terms to cluster GO terms and then runs ht_clusters to visualize the clustering.

The arguments in simplifyGO passed to ht_clusters are:

draw_word_cloud Whether to draw the word clouds.

min_term Minimal number of GO terms in a cluster. All the clusters with size less than min_term are all merged into one single cluster in the heatmap.

order_by_size Whether to reorder GO clusters by their sizes. The cluster that is merged from small clusters (size < min_term) is always put to the bottom of the heatmap.

exclude_words Words that are excluded in the word cloud.

max_words Maximal number of words visualized in the word cloud.

word_cloud_grob_param A list of graphic parameters passed to word_cloud_grob.

fontsize_range The range of the font size. The value should be a numeric vector with length two. The minimal font size is mapped to word frequency value of 1 and the maximal font size is mapped to the maximal word frequency. The font size interlopation is linear.

Value

A data frame with three columns: GO IDs, GO term names and cluster labels.

Examples

```
set.seed(123)
go_id = random_GO(500)
mat = GO_similarity(go_id)
df = simplifyGO(mat, word_cloud_grob_param = list(max_width = 80))
head(df)
```

subset_enrichResult

Subset method of the enrichResult class

Description

Subset method of the enrichResult class

Usage

```
subset_enrichResult(x, i)
```

Arguments

- A enrichResult object from 'clusterProfiler' or other related packages.
- i Row indices.

Value

Still a enrichResult object but with the selected subset of rows.

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Examples

```
\hbox{\tt\# There is no example}\\ \hbox{\tt NULL}
```

term_similarity

Similarity between terms based on the overlap of genes

Description

Similarity between terms based on the overlap of genes

Usage

```
term_similarity(gl, method = c("kappa", "jaccard", "dice", "overlap"))
```

Arguments

gl A list of genes that are in the terms.

method The similarity measurement.

Details

The definition of the four similarity measurements can be found at https://simplifyenrichment.github.io/supplementary/suppl1_coefficient_definition/suppl1_coefficient_definition.html.

Value

A symmetric matrix.

Examples

```
\label{eq:continuous_problem} \mbox{\ensuremath{\texttt{T}} There is no example} \\ \mbox{\ensuremath{\texttt{NULL}}}
```

term_similarity_from_enrichResult

Similarity between terms in the enrichResult class

Description

Similarity between terms in the enrichResult class

Usage

```
term\_similarity\_from\_enrichResult(x, ...)
```

Arguments

x A enrichResult object from 'clusterProfiler' or other related packages.

... Pass to term_similarity.

Details

The object is normally from the 'clusterProfiler', 'DOSE', 'meshes' or 'ReactomePA' package.

Value

A symmetric matrix.

Examples

```
\# There is no example NULL
```

```
term_similarity_from_gmt
```

Similarity between terms from a gmt file

Description

Similarity between terms from a gmt file

Usage

```
term_similarity_from_gmt(term_id, gmt, extract_term_id = NULL, ...)
```

Arguments

```
term_id A vector of terms.

gmt The path of the gmt file.

extract_term_id

If the term ID in contained in the first column only as a substring, setting a function to extract this substring.

... Pass to term_similarity.
```

Value

A symmetric matrix.

```
\# There is no example NULL
```

```
term_similarity_from_KEGG
```

Similarity between KEGG terms

Description

Similarity between KEGG terms

Usage

```
term_similarity_from_KEGG(term_id, ...)
```

Arguments

```
term_id A vector of KEGG IDs, e.g., hsa001.
... Pass to term_similarity.
```

Value

A symmetric matrix.

Examples

```
\hbox{\tt\# There is no example}\\ \hbox{\tt NULL}
```

```
{\tt term\_similarity\_from\_MSigDB}
```

Similarity between MSigDB terms

Description

Similarity between MSigDB terms

Usage

```
term_similarity_from_MSigDB(term_id, category = NULL, subcategory = NULL, ...)
```

Arguments

```
term_id A vector of MSigDB gene set names.
category E.g., 'C1', 'C2', pass to msigdbr.
subcategory E.g., 'CGP', 'BP', pass to msigdbr.
... Pass to term_similarity.
```

Value

A symmetric matrix.

Examples

```
\label{eq:continuous_problem} \mbox{\ensuremath{\mbox{\sc H}}} \mbox{\ensuremath{\mbox{\mbox{\sc H}}}} \mbox{\ensuremath{\mbox{\sc H}}} \mbox{\ensuremath{\mbox{\sc
```

```
{\tt term\_similarity\_from\_Reactome}
```

Similarity between Reactome terms

Description

Similarity between Reactome terms

Usage

```
term_similarity_from_Reactome(term_id, ...)
```

Arguments

```
term_id A vector of Reactome IDs.
... Pass to term_similarity.
```

Value

A symmetric matrix.

Examples

```
\label{eq:total_problem} \mbox{\ensuremath{\texttt{#}} There is no example} \\ \mbox{\ensuremath{\texttt{NULL}}}
```

```
widthDetails.word_cloud
```

Width for word_cloud grob

Description

Width for word_cloud grob

Usage

```
## S3 method for class 'word_cloud'
widthDetails(x)
```

Arguments

x The word_cloud grob returned by word_cloud_grob.

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Value

```
A unit object.
```

Examples

```
\label{eq:continuous_problem} \mbox{\ensuremath{\mbox{\sc H}}} \mbox{\ensuremath{\mbox{\mbox{\sc H}}}} \mbox{\ensuremath{\mbox{\sc H}}} \mbox{\ensuremath{\mbox{\sc
```

word_cloud_grob

A simple grob for the word cloud

Description

A simple grob for the word cloud

Usage

```
word_cloud_grob(text, fontsize,
    line_space = unit(4, "pt"), word_space = unit(4, "pt"), max_width = unit(80, "mm"),
    col = function(fs) circlize::rand_color(length(fs), luminosity = "dark"),
    test = FALSE)
```

Arguments

text	A vector of words.
fontsize	The corresponding font size. With the frequency of the words known, scale_fontsize can be used to linearly interpolate frequencies to font sizes.
line_space	Space between lines. The value can be a unit object or a numeric scalar which is measured in mm.
word_space	Space between words. The value can be a unit object or a numeric scalar which is measured in mm.
max_width	The maximal width of the viewport to put the word cloud. The value can be a unit object or a numeric scalar which is measured in mm. Note this might be larger than the final width of the returned grob object.
col	Colors for the words. The value can be a vector, in numeric or character, which should have the same length as text. Or it is a self-defined function that takes the font size vector as the only argument. The function should return a color vector. See Examples.
test	Internally used. It basically adds borders to the words and the viewport.

Value

A grob object. The width and height of the grob can be get by grobWidth and grobHeight.

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```
# very old R versions do not have strrep() function
if(!exists("strrep")) {
   strrep = function(x, i) paste(rep(x, i), collapse = "")
words = sapply(1:30, function(x) strrep(sample(letters, 1), sample(3:10, 1)))
require(grid)
gb = word_cloud_grob(words, fontsize = runif(30, min = 5, max = 30),
   max_width = 100)
grid.newpage(); grid.draw(gb)
# color as a single scalar
gb = word_cloud_grob(words, fontsize = runif(30, min = 5, max = 30),
   max_width = 100, col = 1)
grid.newpage(); grid.draw(gb)
# color as a vector
gb = word_cloud_grob(words, fontsize = runif(30, min = 5, max = 30),
    max_width = 100, col = 1:30)
grid.newpage(); grid.draw(gb)
# color as a function
require(circlize)
col_fun = colorRamp2(c(5, 17, 30), c("blue", "black", "red"))
gb = word\_cloud\_grob(words, fontsize = runif(30, min = 5, max = 30),
    max_width = 100, col = function(fs) col_fun(fs))
grid.newpage(); grid.draw(gb)
```

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