Package ‘cola’

March 13, 2024

Type Package
Title A Framework for Consensus Partitioning
Version 2.8.0
Date 2023-04-21
Depends R (>= 3.6.0)
Imports grDevices, graphics, grid, stats, utils, ComplexHeatmap (>= 2.5.4), matrixStats, GetoptLong, circlize (>= 0.4.7), GlobalOptions (>= 0.1.0), clue, parallel, RColorBrewer, cluster, skmeans, png, mclust, crayon, methods, xml2, microbenchmark, httr, knitr (>= 1.4.0), markdown (>= 1.6), digest, impute, brew, Rcpp (>= 0.11.0), BiocGenerics, eulerr, foreach, doParallel, doRNG, irlba
Suggests genefilter, mvtnorm, testthat (>= 0.3), samr, pamr, kohonen, NMF, WGCNA, Rtsne, umap, clusterProfiler, ReactomePA, DOSE, AnnotationDbi, gplots, hu6800.db, BiocManager, data.tree, dendextend, Polychrome, rmarkdown, simplifyEnrichment, cowplot, flexclust, randomForest, e1071
Description Subgroup classification is a basic task in genomic data analysis, especially for gene expression and DNA methylation data analysis. It can also be used to test the agreement to known clinical annotations, or to test whether there exist significant batch effects. The cola package provides a general framework for subgroup classification by consensus partitioning. It has the following features: 1. It modularizes the consensus partitioning processes that various methods can be easily integrated. 2. It provides rich visualizations for interpreting the results. 3. It allows running multiple methods at the same time and provides functionalities to straightforward compare results. 4. It provides a new method to extract features which are more efficient to separate subgroups. 5. It automatically generates detailed reports for the complete analysis. 6. It allows applying consensus partitioning in a hierarchical manner.

URL https://github.com/jokergoo/cola,
https://jokergoo.github.io/cola_collection/

VignetteBuilder knitr
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adjust_matrix


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adjust_matrix Remove rows with low variance and impute missing values

Description

Remove rows with low variance and impute missing values

Usage

adjust_matrix(m, sd_quantile = 0.05, max_na = 0.25, verbose = TRUE)

Arguments

m A numeric matrix.

sd_quantile Cutoff of the quantile of standard deviation. Rows with standard deviation less than it are removed.

max_na Maximum NA fraction in each row. Rows with NA fraction larger than it are removed.

verbose Whether to print messages.

Details

The function uses impute.knn to impute missing values, then uses adjust_outlier to adjust outliers and removes rows with low standard deviations.

Value

A numeric matrix.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

set.seed(123)
m = matrix(rnorm(100), nrow = 10)
m[sample(length(m), 5)] = NA
m[1, ] = 0
m
m2 = adjust_matrix(m)
m2
Adjust outliers

Usage

\texttt{adjust\_outlier}(x, q = 0.05)

Arguments

\begin{itemize}
\item \texttt{x} \hspace{1cm} A numeric vector.
\item \texttt{q} \hspace{1cm} Percentile to adjust.
\end{itemize}

Details

Values larger than percentile $1 - q$ are adjusted to the $1 - q$ percentile and values smaller than percentile $q$ are adjusted to the $q$ percentile.

Value

A numeric vector with same length as the original one.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

\begin{verbatim}
set.seed(123)
x = rnorm(40)
x[1] = 100
adjust_outlier(x)
\end{verbatim}
### all_leaves-HierarchicalPartition-method

**All leaves in the hierarchy**

#### Description

All leaves in the hierarchy

#### Usage

```r
## S4 method for signature 'HierarchicalPartition'
all_leaves(object, merge_node = merge_node_param())
```

#### Arguments

- `merge_node`: Parameters to merge sub-dendrograms, see `merge_node_param`.

#### Value

A vector of node ID.

#### Author(s)

Zuguang Gu `<z.gu@dkfz.de>`

#### Examples

```r
data(golub_cola_rh)
all_leaves(golub_cola_rh)
```

---

### all_nodes-HierarchicalPartition-method

**All nodes in the hierarchy**

#### Description

All nodes in the hierarchy

#### Usage

```r
## S4 method for signature 'HierarchicalPartition'
all_nodes(object, merge_node = merge_node_param())
```

#### Examples

```r
data(golub_cola_rh)
all_nodes(golub_cola_rh)
```
all_partition_methods

Arguments

object
A HierarchicalPartition-class object.

merge_node
Parameters to merge sub-dendrograms, see merge_node_param.

Value
A vector of node ID.

Author(s)
Zuguang Gu <z.gu@dkfz.de>

Examples

data(golub_cola_rh)
all_nodes(golub_cola_rh)

all_partition_methods
All supported partitioning methods

Description
All supported partitioning methods

Usage
all_partition_methods()

Details
New partitioning methods can be registered by register_partition_methods.

Value
A vector of supported partitioning methods.

Author(s)
Zuguang Gu <z.gu@dkfz.de>

Examples
all_partition_methods()
all_top_value_methods  *All supported top-value methods*

**Description**

All supported top-value methods

**Usage**

```r
all_top_value_methods()
```

**Details**

New top-value methods can be registered by `register_top_value_methods`.

**Value**

A vector of supported top-value methods.

**Author(s)**

Zuguang Gu <z.gu@dkfz.de>

**Examples**

```r
all_top_value_methods()
```

---

**aPAC  *Adapted PAC scores***

**Description**

Adapted PAC scores

**Usage**

```r
aPAC(consensus_mat)
```

**Arguments**

- `consensus_mat`  A consensus matrix.
Details

For the consensus values x, it is transformed to 1 - x if x < 0.5. After the transformation, for any pair of samples in the consensus matrix, if they are always in the same group or always in different groups, the value x is both to 1. Thus, if the consensus matrix shows stable partitions, values x will be all close to 1. Reflected in the CDF of x, the curve is shifted to the right and the area under CDF curve should be very small.

An aPAC value less than 0.05 is considered as the stable partition, which can be thought the proportion of ambiguous partitioning is less than 0.05.

Value

A numeric value.

Examples

data(golub_cola)
aPAC(get_consensus(golub_cola[1, 1], k = 2))
aPAC(get_consensus(golub_cola[1, 1], k = 3))
aPAC(get_consensus(golub_cola[1, 1], k = 4))
aPAC(get_consensus(golub_cola[1, 1], k = 5))
aPAC(get_consensus(golub_cola[1, 1], k = 6))

---

ATC

<table>
<thead>
<tr>
<th>Ability to correlate to other rows</th>
</tr>
</thead>
</table>

Description

Ability to correlate to other rows

Usage

ATC(mat, cor_fun = stats::cor, min_cor = 0, power = 1, k_neighbours = -1, group = NULL, mc.cores = 1, cores = mc.cores, ...)

Arguments

<table>
<thead>
<tr>
<th>mat</th>
<th>A numeric matrix. ATC score is calculated by rows.</th>
</tr>
</thead>
<tbody>
<tr>
<td>cor_fun</td>
<td>A function which calculates correlations.</td>
</tr>
<tr>
<td>min_cor</td>
<td>Cutoff for the minimal absolute correlation.</td>
</tr>
<tr>
<td>power</td>
<td>Power on the correlation values.</td>
</tr>
<tr>
<td>k_neighbours</td>
<td>Nearest k neighbours.</td>
</tr>
<tr>
<td>mc.cores</td>
<td>Number of cores. This argument will be removed in future versions.</td>
</tr>
<tr>
<td>cores</td>
<td>Number of cores.</td>
</tr>
<tr>
<td>group</td>
<td>A categorical variable. If it is specified, the correlation is only calculated for the rows in the same group as current row.</td>
</tr>
<tr>
<td>...</td>
<td>Pass to cor_fun.</td>
</tr>
</tbody>
</table>
Details

For a given row in a matrix, the ATC score is the area above the curve of the cumulative density distribution of the absolute correlation to all other rows. Formally, if $F_i(X)$ is the cumulative distribution function of $X$ where $X$ is the absolute correlation for row $i$ with power power (i.e. $x = cor^\text{power}$), $\text{ATC}_i = 1 - \int_{\text{min_cor}}^1 F_i(X)$.

By default the ATC scores are calculated by Pearson correlation, to use Spearman correlation, you can register a new top-value method by:

```r
register_top_value_methods(
  "ATC_spearman" = function(m) ATC(m, method = "spearman")
)
```

Similarly, to use a robust correlation method, e.g. `bicor` function, you can do like:

```r
register_top_value_methods(
  "ATC_bicor" = function(m) ATC(m, cor_fun = WGCNA::bicor)
)
```

If the number of rows exceeds 30000, it internally uses `ATC_approx`.

Value

A vector of numeric values with the same order as rows in the input matrix.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

See Also

https://jokergoo.github.io/cola_supplementary/suppl_1_ATC/suppl_1_ATC.html

Examples

```r
set.seed(12345)
nr1 = 100
mat1 = matrix(rnorm(100*nr1), nrow = nr1)
nr2 = 10
require(mvtnorm)
sigma = matrix(0.8, nrow = nr2, ncol = nr2); diag(sigma) = 1
mat2 = t(rmvnorm(100, mean = rep(0, nr2), sigma = sigma))
nr3 = 50
sigma = matrix(0.5, nrow = nr3, ncol = nr3); diag(sigma) = 1
mat3 = t(rmvnorm(100, mean = rep(0, nr3), sigma = sigma))
mat = rbind(mat1, mat2, mat3)
ATC_score = ATC(mat)
plot(ATC_score, pch = 16, col = c(rep(1, nr1), rep(2, nr2), rep(3, nr3)))
```
ATC_approx

---

**ATC_approx**

*Ability to correlate to other rows - an approximated method*

---

**Description**

Ability to correlate to other rows - an approximated method

**Usage**

```
ATC_approx(mat, cor_fun = stats::cor, min_cor = 0, power = 1, k_neighbours = -1, 
            mc.cores = 1, cores = mc.cores, n_sampling = c(1000, 500), 
            group = NULL, ...)
```

**Arguments**

- **mat**: A numeric matrix. ATC score is calculated by rows.
- **cor_fun**: A function which calculates correlations on matrix rows.
- **min_cor**: Cutoff for the minimal absolute correlation.
- **power**: Power on the correlation values.
- **k_neighbours**: Nearest k neighbours. Note when this argument is set, there won’t be subset sampling for calculating correlations, which means, it will calculate correlation to all other rows.
- **mc.cores**: Number of cores. This argument will be removed in future versions.
- **cores**: Number of cores.
- **n_sampling**: When there are too many rows in the matrix, to get the cumulative distribution of how one row correlates other rows, actually we don’t need to use all the rows in the matrix, e.g. 1000 rows can already give a very nice estimation.
- **group**: A categorical variable. If it is specified, the correlation is only calculated for the rows in the same group as current row.
- **...**: Pass to `cor_fun`.

**Details**

For a matrix with huge number of rows. It is not possible to calculate correlation to all other rows, thus the correlation is only calculated for a randomly sampled subset of other rows.

With small numbers of rows of the matrix, ATC should be used which calculates the "exact" ATC value, but the value of ATC and ATC_approx should be very similar.

**Examples**

```r
# There is no example
NULL
```
cola

Description
A bottle of cola

Usage
cola()

Details
Simply serve you a bottle of cola.
The ASCII art is from http://ascii.co.uk/art/coke.

Value
No value is returned.

Author(s)
Zuguang Gu <z.gu@dkfz.de>

Examples
for(i in 1:10) cola()

cola_opt

Description
Global parameters

Usage
cola_opt(..., RESET = FALSE, READ.ONLY = NULL, LOCAL = FALSE, ADD = FALSE)

Arguments
... Arguments for the parameters, see "details" section.
RESET Whether to reset to default values.
READ.ONLY Please ignore.
LOCAL Please ignore.
ADD Please ignore.
Details

There are following global parameters:

- **group_diff**: Used in `get_signatures,ConsensusPartition-method` to globally control the minimal difference between subgroups.
- **fdr_cutoff**: Used in `get_signatures,ConsensusPartition-method` to globally control the cutoff of FDR for the differential signature tests.
- **color_set_2**: Colors for the predicted subgroups.
- **help**: Whether to print help messages.
- **message**: Whether to print messages.

Examples

```r
cola_opt
cola_opt$group_diff = 0.2  # e.g. for methylation datasets
cola_opt$fdr_cutoff = 0.1  # e.g. for methylation datasets
cola_opt
cola_opt(RESET = TRUE)
```

Description

Make HTML report from the ConsensusPartition object

Usage

```r
## S4 method for signature 'ConsensusPartition'
cola_report(object, output_dir = getwd(),
  title = qq("cola Report for Consensus Partitioning (@{object@top_value_method}:@{object@partition_method})"),
  env = parent.frame())
```

Arguments

- **object**: A `ConsensusPartition-class` object.
- **output_dir**: The output directory where the report is saved.
- **title**: Title of the report.
- **env**: Where the objects in the report are found, internally used.

Details

It generates report for a specific combination of top-value method and partitioning method.
Value

No value is returned.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

See Also

cola_report, ConsensusPartitionList-method

Examples

# There is no example
NULL
Value

No value is returned.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

```r
if(FALSE) {
# the following code is runnable
data(golub_cola)
cola_report(golub_cola[c("SD", "MAD"), c("hclust", "skmeans")], output_dir = "~/test_cola_cl_report")
}
```

Description

Method dispatch page for `cola_report`.

Dispatch

`cola_report` can be dispatched on following classes:

- `cola_report,HierarchicalPartition-method, HierarchicalPartition-class` class method
- `cola_report,ConsensusPartition-method, ConsensusPartition-class` class method
- `cola_report,ConsensusPartitionList-method, ConsensusPartitionList-class` class method

Examples

# no example
NULL
Description

Make HTML report from the HierarchicalPartition object

Usage

```r
## S4 method for signature 'HierarchicalPartition'
cola_report(object, output_dir = getwd(), mc.cores = 1, cores = mc.cores,
title = qq("cola Report for Hierarchical Partitioning"),
env = parent.frame())
```

Arguments

- `output_dir`: The output directory where the report is put.
- `mc.cores`: Multiple cores to use. This argument will be removed in future versions.
- `cores`: Number of cores, or a cluster object returned by `makeCluster`.
- `title`: Title of the report.
- `env`: Where the objects in the report are found, internally used.

Details

This function generates a HTML report which contains all plots for all nodes in the partition hierarchy.

Value

No value is returned.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

```r
if(FALSE) {
  # the following code is runnable
  data(golub_cola_rh)
cola_report(golub_cola_rh, output_dir = "~/test_cola_rh_report")
}
```
Example ConsensusPartitionList object

Description
Example ConsensusPartitionList object

Usage
data(cola_rl)

Details
Following code was used to generate cola_rl:

```r
set.seed(123)
m = cbind(rbind(matrix(rnorm(20*20, mean = 1, sd = 0.5), nr = 20),
              matrix(rnorm(20*20, mean = 0, sd = 0.5), nr = 20),
              matrix(rnorm(20*20, mean = 0, sd = 0.5), nr = 20)),
           rbind(matrix(rnorm(20*20, mean = 0, sd = 0.5), nr = 20),
                  matrix(rnorm(20*20, mean = 1, sd = 0.5), nr = 20),
                  matrix(rnorm(20*20, mean = 0, sd = 0.5), nr = 20)),
           rbind(matrix(rnorm(20*20, mean = 0.5, sd = 0.5), nr = 20),
                  matrix(rnorm(20*20, mean = 0.5, sd = 0.5), nr = 20),
                  matrix(rnorm(20*20, mean = 1, sd = 0.5), nr = 20))) + matrix(rnorm(60*60, sd = 0.5), nr = 60)
cola_rl = run_all_consensus_partition_methods(data = m, cores = 6)
```

Author(s)
Zuguang Gu <z.gu@dkfz.de>

Examples
data(cola_rl)
cola_rl

Collect subgroups from ConsensusPartition object

Description
Collect subgroups from ConsensusPartition object
Usage

```r
## S4 method for signature 'ConsensusPartition'
collect_classes(object, internal = FALSE,
    show_row_names = FALSE, row_names_gp = gpar(fontsize = 8),
    anno = object@anno, anno_col = object@anno_col)
```

Arguments

- `internal`: Used internally.
- `show_row_names`: Whether to show row names in the heatmap (which is the column name in the original matrix).
- `row_names_gp`: Graphics parameters for row names.
- `anno`: A data frame of annotations for the original matrix columns. By default it uses the annotations specified in `consensus_partition` or `run_all_consensus_partition_methods`.
- `anno_col`: A list of colors (color is defined as a named vector) for the annotations. If `anno` is a data frame, `anno_col` should be a named list where names correspond to the column names in `anno`.

Details

The percent membership matrix and the subgroup labels for each k are plotted in the heatmaps. Same row in all heatmaps corresponds to the same column in the original matrix.

Value

No value is returned.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

```r
data(golub_cola)
collect_classes(golub_cola["ATC", "skmeans"])
```
Usage

```r
## S4 method for signature 'ConsensusPartitionList'
collect_classes(object, k, show_column_names = FALSE,
column_names_gp = gpar(fontsize = 8),
anno = get_anno(object), anno_col = get_anno_col(object),
simplify = FALSE, ...)
```

Arguments

- **object**: A `ConsensusPartitionList-class` object returned by `run_all_consensus_partition_methods`.
- **k**: Number of subgroups.
- **show_column_names**: Whether to show column names in the heatmap (which is the column name in the original matrix).
- **column_names_gp**: Graphics parameters for column names.
- **anno**: A data frame of annotations for the original matrix columns. By default it uses the annotations specified in `run_all_consensus_partition_methods`.
- **anno_col**: A list of colors (color is defined as a named vector) for the annotations. If `anno` is a data frame, `anno_col` should be a named list where names correspond to the column names in `anno`.
- **simplify**: Internally used.
- **...**: Pass to `drawHeatmapList-method`.

Details

There are following panels in the plot:

- a heatmap showing partitions predicted from all methods where the top annotation is the consensus partition summarized from partitions from all methods, weighted by mean silhouette scores in every single method.
- a row barplot annotation showing the mean silhouette scores for different methods.

The row clustering is applied on the dissimilarity matrix calculated by `cl_dissimilarity` with the comembership method.

The brightness of the color corresponds to the silhouette scores for the consensus partition in each method.

Value

No value is returned.

Author(s)

Zuguang Gu <z.gu@dkfz.de>
Examples

data(golub_cola)
collect_classes(golub_cola, k = 3)

collect_classes-HierarchicalPartition-method

Collect classes from HierarchicalPartition object

Description

Collect classes from HierarchicalPartition object

Usage

## S4 method for signature 'HierarchicalPartition'
collect_classes(object, merge_node = merge_node_param(),
    show_row_names = FALSE, row_names_gp = gpar(fontsize = 8),
    anno = get_anno(object[1]), anno_col = get_anno_col(object[1]), ...)

collect_classes-dispatch

Method dispatch page for collect_classes

Description

Method dispatch page for collect_classes.

Dispatch

collect_classes can be dispatched on following classes:

• collect_classes,HierarchicalPartition-method,HierarchicalPartition-class class method
• collect_classes,ConsensusPartitionList-method,ConsensusPartitionList-class class method
• collect_classes,ConsensusPartition-method,ConsensusPartition-class class method

Examples

# no example
NULL
Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>object</td>
<td>A HierarchicalPartition-class object.</td>
</tr>
<tr>
<td>merge_node</td>
<td>Parameters to merge sub-dendrograms, see merge_node_param.</td>
</tr>
<tr>
<td>show_row_names</td>
<td>Whether to show the row names.</td>
</tr>
<tr>
<td>row_names_gp</td>
<td>Graphic parameters for row names.</td>
</tr>
<tr>
<td>anno</td>
<td>A data frame of annotations for the original matrix columns. By default it uses the annotations specified in hierarchical_partition.</td>
</tr>
<tr>
<td>anno_col</td>
<td>A list of colors (color is defined as a named vector) for the annotations. If anno is a data frame, anno_col should be a named list where names correspond to the column names in anno.</td>
</tr>
<tr>
<td>...</td>
<td>Other arguments.</td>
</tr>
</tbody>
</table>

Details

The function plots the hierarchy of the classes.

Value

No value is returned.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

data(golub_cola_rh)
collect_classes(golub_cola_rh)
collect_classes(golub_cola_rh, merge_node = merge_node_param(depth = 2))
Details

Plots by `plot_ecdf, collect_classes, ConsensusPartition-method, consensus_heatmap, membership_heatmap` and `get_signatures` are arranged in one single page, for all available k.

Value

No value is returned.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

See Also

`collect_plots, ConsensusPartitionList-method` collects plots for the `ConsensusPartitionList-class` object.

Examples

```r
data(golub_cola)
collect_plots(golub_cola["ATC", "skmeans"])
```

Description

Collect plots from ConsensusPartitionList object

Usage

```r
## S4 method for signature 'ConsensusPartitionList'
collect_plots(object, k = 2, fun = consensus_heatmap,
              top_value_method = object@top_value_method,
              partition_method = object@partition_method,
              verbose = TRUE, mc.cores = 1, cores = mc.cores, ...)
```

Arguments

- `object` A `ConsensusPartitionList-class` object from `run_all_consensus_partition_methods`.
- `k` Number of subgroups.
- `fun` Function used to generate plots. Valid functions are `consensus_heatmap, plot_ecdf, membership_heatmap, get_signatures` and `dimension_reduction`.
- `top_value_method` A vector of top-value methods.
**partition_method**
A vector of partitioning methods.

**verbose**
Whether to print message.

**mc.cores**
Number of cores. This argument will be removed in figure versions.

**cores**
Number of cores, or a cluster object returned by `makeCluster`.

... other Arguments passed to corresponding fun.

### Details
Plots for all combinations of top-value methods and partitioning methods are arranged in one single page.
This function makes it easy to directly compare results from multiple methods.

### Value
No value is returned.

### Author(s)
Zuguang Gu <z.gu@dkfz.de>

### See Also
- `collect_plots,ConsensusPartition-method` collects plots for a single `ConsensusPartition-class` object.

### Examples
```r
data(golub_cola)
collect_plots(cola_rl, k = 3)
collect_plots(cola_rl, k = 3, fun = membership_heatmap)
collect_plots(cola_rl, k = 3, fun = get_signatures)
```

---

**Description**
Method dispatch page for `collect_plots`.

**Dispatch**
`collect_plots` can be dispatched on following classes:

- `collect_plots,ConsensusPartition-method,ConsensusPartition-class` class method
- `collect_plots,ConsensusPartitionList-method,ConsensusPartitionList-class` class method
**Examples**

```r
# no example
NULL
```

**Description**

Draw and compare statistics for a single method

**Usage**

```r
## S4 method for signature 'ConsensusPartition'
collect_stats(object, ...)
```

**Arguments**

- `object` A `ConsensusPartition-class` object.
- `...` Other arguments.

**Details**

It is identical to `select_partition_number,ConsensusPartition-method`.

**Examples**

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

**collect_stats-ConsensusPartitionList-method**

*Draw and compare statistics for multiple methods*

**Description**

Draw and compare statistics for multiple methods

**Usage**

```r
## S4 method for signature 'ConsensusPartitionList'
collect_stats(object, k, layout_nrow = 2, all_stats = FALSE, ...)
```
Arguments

- **object**: A ConsensusPartitionList-class object.
- **k**: Number of subgroups.
- **layout_nrow**: Number of rows in the layout.
- **all_stats**: Whether to show all statistics that were calculated. Used internally.
- **...**: Other arguments.

Details

It draws heatmaps for statistics for multiple methods in parallel, so that users can compare which combination of methods gives the best results with given the number of subgroups.

Examples

```r
data(golub_cola)
collect_stats(golub_cola, k = 3)
```

---

**Description**

Method dispatch page for collect_stats.

**Dispatch**

collect_stats can be dispatched on following classes:

- `collect_stats,ConsensusPartitionList-method,ConsensusPartitionList-class` class method
- `collect_stats,ConsensusPartition-method,ConsensusPartition-class` class method

**Examples**

```r
# no example
NULL
```
Description

Column names of the matrix

Usage

```r
## S4 method for signature 'ConsensusPartition'
colnames(x)
```

Arguments

- `x`: A `ConsensusPartition-class` object.

Examples

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

Description

Column names of the matrix

Usage

```r
## S4 method for signature 'ConsensusPartitionList'
colnames(x)
```

Arguments

- `x`: A `ConsensusPartitionList-class` object.

Examples

```r
# There is no example
NULL
```
Description

Method dispatch page for colnames.

Dispatch

colnames can be dispatched on following classes:

- `colnames,ConsensusPartition-method`, `ConsensusPartition-class` class method
- `colnames,ConsensusPartitionList-method`, `ConsensusPartitionList-class` class method
- `colnames,DownSamplingConsensusPartition-method`, `DownSamplingConsensusPartition-class` class method
- `colnames,HierarchicalPartition-method`, `HierarchicalPartition-class` class method

Examples

```
# no example
NULL
```

---

### colnames-DownSamplingConsensusPartition-method

**Column names of the matrix**

Description

Column names of the matrix

Usage

```
## S4 method for signature 'DownSamplingConsensusPartition'
colnames(x)
```

Arguments

- `x` A `DownSamplingConsensusPartition-class` object.

Examples

```
# There is no example
NULL
```
**colnames-HierarchicalPartition-method**

*Column names of the matrix*

**Description**

Column names of the matrix

**Usage**

```r
## S4 method for signature 'HierarchicalPartition'
colnames(x)
```

**Arguments**

- `x` A `HierarchicalPartition-class` object.

**Examples**

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

---

**compare_partitions-ConsensusPartition-method**

*Compare two partitionings*

**Description**

Compare two partitionings

**Usage**

```r
## S4 method for signature 'ConsensusPartition'
compare_partitions(object, object2, output_file, k1 = 2, k2 = 2,
                   dimension_reduction_method = "UMAP",
                   id_mapping = guess_id_mapping(rownames(object), "org.Hs.eg.db", FALSE),
                   row_km1 = ifelse(k1 == 2, 2, 1),
                   row_km2 = ifelse(k1 ==2 & k2 == 2, 2, 1),
                   row_km3 = ifelse(k2 == 2, 2, 1))
```
**Arguments**

- **object** A `ConsensusPartition` object.
- **object2** A `ConsensusPartition` object.
- **output_file** The path of the output HTML file. If it is not specified, the report will be opened in the web browser.
- **k1** Number of subgroups in `object`.
- **k2** Number of subgroups in `object2`.
- **dimension_reduction_method** Which dimension reduction method to use.
- **id_mapping** Pass to `functional_enrichment`.
- **row_km1** Number of k-means groups, see Details.
- **row_km2** Number of k-means groups, see Details.
- **row_km3** Number of k-means groups, see Details.

**Details**

The function produces a HTML report which includes comparisons between two partitioning results.

In the report, there are three heatmaps which visualize A) the signature genes specific in the first partition, B) the signature genes both in the two partitionings and C) the signatures genes specific in the second partition. Argument `row_km1`, `row_km2` and `row_km3` control how many k-means groups should be applied on the three heatmaps.

**Examples**

```r
## Not run:
data(golub_cola)
require(hu6800.db)
x = hu6800ENTREZID
mapped_probes = mappedkeys(x)
id_mapping = unlist(as.list(x[mapped_probes]))
compare_partitions(golub_cola["ATC:skmeans"], golub_cola["SD:kmeans"],
                   id_mapping = id_mapping)
## End(Not run)
```

**Description**

Compare Signatures from Different k
Usage

```r
## S4 method for signature 'ConsensusPartition'
compare_signatures(object, k = object@k, verbose = interactive(), ...)
```

Arguments

- **object**: A `ConsensusPartition-class` object.
- **k**: Number of subgroups. Value should be a vector.
- **verbose**: Whether to print message.
- **...**: Other arguments passed to `get_signatures,ConsensusPartition-method`.

Details

It plots an Euler diagram showing the overlap of signatures from different k.

Examples

```r
data(golub_cola)
res = golub_cola["ATC", "skmeans"]
compare_signatures(res)
```
**compare_signatures-HierarchicalPartition-method**

*Compare Signatures from Different Nodes*

**Description**

Compare Signatures from Different Nodes

**Usage**

```r
## S4 method for signature 'HierarchicalPartition'
compare_signatures(object, merge_node = merge_node_param(),
                   method = c("euler", "upset"), upset_max_comb_sets = 20,
                   verbose = interactive(), ...)
```

**Arguments**

- `merge_node`: Parameters to merge sub-dendrograms, see `merge_node_param`.
- `method`: Method to visualize.
- `upset_max_comb_sets`: Maximal number of combination sets to show.
- `verbose`: Whether to print message.
- `...`: Other arguments passed to `get_signatures,HierarchicalPartition-method`.

**Details**

It plots an Euler diagram or a UpSet plot showing the overlap of signatures from different nodes. On each node, the number of subgroups is inferred by `suggest_best_k,ConsensusPartition-method`.

**Examples**

```r
data(golub_cola_rh)
compare_signatures(golub_cola_rh)
```

---

**concordance**

*Concordance to the consensus partition*

**Description**

Concordance to the consensus partition

**Usage**

```r
concordance(membership_each, class)
```
Arguments

membership_each
A matrix which contains partitions in every single runs where columns correspond to runs. The object can be get from get_membership(..., each = TRUE).

class
Consensus subgroup labels.

Details

Note subgroup labels in membership_each should already be adjusted to the consensus labels, i.e. by relabel_class.
The concordance score is the mean proportion of samples having the same subgroup labels as the consensus labels among individual partition runs.

Value

A numeric value.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

data(golub_cola)
membership_each = get_membership(golub_cola["SD", "kmeans"], each = TRUE, k = 3)
consensus_classes = get_classes(golub_cola["SD", "kmeans"], k = 3)$class
cordance(membership_each, consensus_classes)

Description

Adjust parameters for default ATC method

Usage

cfg_ATC(cor_fun = stats::cor, min_cor = 0, power = 1, k_neighbours = -1, group = NULL, cores = 1, ...)

Arguments

cor_fun
A function that calculates correlations from a matrix (on matrix rows).

min_cor
Cutoff for the minimal absolute correlation.

power
Power on the correlation values.

k_neighbours
Number of the closest neighbours to use.
group A categorical variable.
cores Number of cores.
... Other arguments passed to ATC.

Details
This function changes the default parameters for ATC method. All the arguments in this function all pass to ATC.

Examples

# use Spearman correlation
config_ATC(cor_fun = function(m) stats::cor(m, method = "spearman"))
# use knn
config_ATC(k_neighbours = 100)

Description
The ConsensusPartition class

Methods
The ConsensusPartition-class has following methods:

consensus_partition: constructor method, run consensus partitioning with a specified top-value method and a partitioning method.
select_partition_number,ConsensusPartition-method: make a list of plots for selecting optimized number of subgroups.
consensus_heatmap,ConsensusPartition-method: make heatmap of the consensus matrix.
membership_heatmap,ConsensusPartition-method: make heatmap of the membership for individual partitions.
get_signatures,ConsensusPartition-method: get the signature rows and make heatmap.
dimension_reduction,ConsensusPartition-method: make dimension reduction plots.
collect_plots,ConsensusPartition-method: make heatmaps for consensus matrix and membership matrix with different number of subgroups.
collect_classes,ConsensusPartition-method: make heatmap with subgroups with different numbers.
get_param,ConsensusPartition-method: get parameters for the consensus clustering.
get_matrix,ConsensusPartition-method: get the original matrix.
get_consensus,ConsensusPartition-method: get the consensus matrix.
get_membership, ConsensusPartition-method: get the membership of partitions generated from random samplings.
get_stats, ConsensusPartition-method: get statistics for the consensus partitioning.
get_classes, ConsensusPartition-method: get the consensus subgroup labels and other columns.
suggest_best_k, ConsensusPartition-method: guess the best number of subgroups.
test_to_known_factors, ConsensusPartition-method: test correlation between predicted subgroups and known factors, if available.
cola_report, ConsensusPartition-method: generate a HTML report for the whole analysis.
functional_enrichment, ConsensusPartition-method: perform functional enrichment analysis on significant genes if rows in the matrix can be corresponded to genes.

Author(s)
Zuguang Gu <z.gu@dkfz.de>

Examples
# There is no example
NULL

ConsensusPartitionList-class

The ConsensusPartitionList class

Description
The ConsensusPartitionList class

Details
The object contains results from all combinations of top-value methods and partitioning methods.

Methods
The ConsensusPartitionList-class provides following methods:
run_all_consensus_partition_methods: constructor method.
top_rows_overlap, ConsensusPartitionList-method: plot the overlaps of top rows under different top-value methods.
top_rows_heatmap, ConsensusPartitionList-method: plot the heatmap of top rows under different top-value methods.
get_classes, ConsensusPartitionList-method: get consensus subgroup labels merged from all methods.
generate_matrix, ConsensusPartition-method: get the original matrix.
**get_stats,ConsensusPartitionList-method:** get statistics for the partition for a specified k.

**get_membership,ConsensusPartitionList-method:** get consensus membership matrix summarized from all methods.

**suggest_best_k,ConsensusPartitionList-method:** guess the best number of subgroups for all methods.

**collect_plots,ConsensusPartitionList-method:** collect plots from all combinations of top-value methods and partitioning methods with choosing a plotting function.

**collect_classes,ConsensusPartitionList-method:** make a plot which contains predicted subgroups from all combinations of top-value methods and partitioning methods.

**test_to_known_factors,ConsensusPartitionList-method:** test correlation between predicted subgroups and known annotations, if provided.

**cola_report,ConsensusPartitionList-method:** generate a HTML report for the whole analysis.

**functional_enrichment,ConsensusPartitionList-method:** perform functional enrichment analysis on significant genes if rows in the matrix can be corresponded to genes.

**Author(s)**

Zuguang Gu <z.gu@dkfz.de>

**See Also**

The `ConsensusPartition-class`.

**Examples**

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

---

**consensus_heatmap-ConsensusPartition-method**

*Heatmap of the consensus matrix*

---

**Description**

Heatmap of the consensus matrix

**Usage**

```r
## S4 method for signature 'ConsensusPartition'
consensus_heatmap(object, k, internal = FALSE,
                   anno = object$anno, anno_col = get_anno_col(object),
                   show_row_names = FALSE, show_column_names = FALSE, row_names_gp = gpar(fontsize = 8),
                   simplify = FALSE, ...)```
Arguments

- **object**
  A ConsensusPartition-class object.

- **k**
  Number of subgroups.

- **internal**
  Used internally.

- **anno**
  A data frame of annotations for the original matrix columns. By default it uses the annotations specified in the consensus_partition or run_all_consensus_partition_methods.

- **anno_col**
  A list of colors (color is defined as a named vector) for the annotations. If anno is a data frame, anno_col should be a named list where names correspond to the column names in anno.

- **show_row_names**
  Whether plot row names on the consensus heatmap (which are the column names in the original matrix).

- **show_column_names**
  Whether show column names.

- **row_names_gp**
  Graphics parameters for row names.

- **simplify**
  Internally used.

- **...**
  Other arguments.

Details

For row i and column j in the consensus matrix, the value of corresponding x_ij is the probability of sample i and sample j being in a same group from all partitions.

There are following heatmaps from left to right:

- probability of the sample to stay in the corresponding group
- silhouette scores which measure the distance of an item to the second closest subgroups.
- predicted subgroups
- consensus matrix.
- more annotations if provided as anno

One thing that is very important to note is that since we already know the consensus subgroups from consensus partition, in the heatmap, only rows or columns within the group is clustered.

Value

No value is returned.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

See Also

membership_heatmap, ConsensusPartition-method
Examples

```r
data(golub_cola)
consensus_heatmap(golub_cola[, "ATC", "skmeans"], k = 3)
```

---

**Description**

Consensus partition

**Usage**

```r
consensus_partition(data,
top_value_method = "ATC",
top_n = NULL,
partition_method = "skmeans",
max_k = 6,
k = NULL,
sample_by = "row",
p_sampling = 0.8,
partition_repeat = 50,
partition_param = list(),
anno = NULL,
anno_col = NULL,
scale_rows = NULL,
verbose = TRUE,
mc.cores = 1, cores = mc.cores,
prefix = "",
.env = NULL,
help = cola_opt$help)
```

**Arguments**

- **data**: A numeric matrix where subgroups are found by columns.
- **top_value_method**: A single top-value method. Available methods are in `all_top_value_methods`. Use `register_top_value_methods` to add a new top-value method.
- **top_n**: Number of rows with top values. The value can be a vector with length > 1. When n > 5000, the function only randomly sample 5000 rows from top n rows. If `top_n` is a vector, partition will be applied to every values in `top_n` and consensus partition is summarized from all partitions.
- **partition_method**: A single partitioning method. Available methods are in `all_partition_methods`. Use `register_partition_methods` to add a new partition method.
max_k  Maximal number of subgroups to try. The function will try for 2:max_k subgroups.
k  Alternatively, you can specify a vector k.
sample_by  Should randomly sample the matrix by rows or by columns?
p_sampling  Proportion of the submatrix which contains the top n rows to sample.
partition_repeat  Number of repeats for the random sampling.
partition_param  Parameters for the partition method which are passed to ... in a registered partitioning method. See register_partition_methods for detail.
anno  A data frame with known annotation of samples. The annotations will be plotted in heatmaps and the correlation to predicted subgroups will be tested.
anno_col  A list of colors (color is defined as a named vector) for the annotations. If anno is a data frame, anno_col should be a named list where names correspond to the column names in anno.
scale_rows  Whether to scale rows. If it is TRUE, scaling method defined in register_partition_methods is used.
verbose  Whether print messages.
mc.cores  Multiple cores to use. This argument will be removed in future versions.
cores  Number of cores, or a cluster object returned by makeCluster.
prefix  Internally used.
.env  An environment, internally used.
help  Whether to print help messages.

Details

The function performs analysis in following steps:

• calculate scores for rows by top-value method,
• for each top_n value, take top n rows,
• randomly sample p_sampling rows from the top_n-row matrix and perform partitioning for partition_repeats times,
• collect partitions from all individual partitions and summarize a consensus partition.

Value

A ConsensusPartition-class object. Simply type object in the interactive R session to see which functions can be applied on it.

Author(s)

Zuguang Gu <z.gu@dkfz.de>
See Also

run_all_consensus_partition_methods runs consensus partitioning with multiple top-value methods and multiple partitioning methods.

Examples

```r
set.seed(123)
m = cbind(rbind(matrix(rnorm(20*20, mean = 1, sd = 0.5), nr = 20),
    matrix(rnorm(20*20, mean = 0, sd = 0.5), nr = 20),
    matrix(rnorm(20*20, mean = 0, sd = 0.5), nr = 20)),
    rbind(matrix(rnorm(20*20, mean = 0, sd = 0.5), nr = 20),
    matrix(rnorm(20*20, mean = 1, sd = 0.5), nr = 20),
    matrix(rnorm(20*20, mean = 0, sd = 0.5), nr = 20)),
    rbind(matrix(rnorm(20*20, mean = 0.5, sd = 0.5), nr = 20),
    matrix(rnorm(20*20, mean = 0.5, sd = 0.5), nr = 20),
    matrix(rnorm(20*20, mean = 1, sd = 0.5), nr = 20))
) + matrix(rnorm(60*60, sd = 0.5), nr = 60)
res = consensus_partition(m, partition_repeat = 10, top_n = c(10, 20, 50))
res
```

---

**consensus_partition_by_down_sampling**

Consensus partitioning only with a subset of columns

**Description**

Consensus partitioning only with a subset of columns

**Usage**

```r
consensus_partition_by_down_sampling(data,
    top_value_method = "ATC",
    top_n = NULL,
    partition_method = "skmeans",
    max_k = 6, k = NULL,
    subset = min(round(ncol(data)*0.2), 250), pre_select = TRUE,
    verbose = TRUE, prefix = "", anno = NULL, anno_col = NULL,
    predict_method = "centroid",
    dist_method = c("euclidean", "correlation", "cosine"),
    .env = NULL, .predict = TRUE, mc.cores = 1, cores = mc.cores, ...)
```

**Arguments**

- **data**
  A numeric matrix where subgroups are found by columns.

- **top_value_method**
  A single top-value method. Available methods are in all_top_value_methods. Use register_top_value_methods to add a new top-value method.
**top_n**  Number of rows with top values. The value can be a vector with length > 1. When n > 5000, the function only randomly sample 5000 rows from top n rows. If top_n is a vector, partition will be applied to every values in top_n and consensus partition is summarized from all partitions.

**partition_method**  
A single partitioning method. Available methods are in `all_partition_methods`. Use `register_partition_methods` to add a new partition method.

**max_k**  
Maximal number of subgroups to try. The function will try for 2:max_k subgroups

**k**  
Alternatively, you can specify a vector k.

**subset**  
Number of columns to randomly sample, or a vector of selected indices.

**pre_select**  
Whether to pre-select by k-means.

**verbose**  
Whether to print messages.

**prefix**  
Internally used.

**anno**  
Annotation data frame.

**anno_col**  
Annotation colors.

**predict_method**  
Method for predicting class labels. Possible values are "centroid", "svm" and "randomForest".

**dist_method**  
Method for predict the class for other columns.

**.env**  
An environment, internally used.

**.predict**  
Internally used.

**mc.cores**  
Number of cores. This argument will be removed in future versions.

**cores**  
Number of cores, or a cluster object returned by `makeCluster`.

**...**  
All pass to `consensus_partition`.

## Details

The function performs consensus partitioning only with a small subset of columns and the class of other columns are predicted by `predict_classes,ConsensusPartition-method`.

## Examples

```r
## Not run:
data(golub_cola)
m = get_matrix(golub_cola)
set.seed(123)
golub_cola_ds = consensus_partition_by_down_sampling(m, subset = 50,
anno = get_anno(golub_cola), anno_col = get_anno_col(golub_cola),
top_value_method = "SD", partition_method = "kmeans")
```

## End(Not run)
correspond_between_rankings

Correspond between a list of rankings

Description

Correspond between a list of rankings

Usage

correspond_between_rankings(lt, top_n = length(lt[[1]]),
  col = cola_opt$color_set_1[1:length(lt)], ...)

Arguments

lt              A list of scores under different metrics.
top_n           Top n elements to show the correspondance.
col             A vector of colors for lt.
...             Pass to correspond_between_two_rankings.

Details

It makes plots for every pairwise comparison in lt.

Value

No value is returned.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

require(matrixStats)
mat = matrix(runif(1000), ncol = 10)
x1 = rowSds(mat)
x2 = rowMads(mat)
x3 = rowSds(mat)/rowMeans(mat)
correspond_between_rankings(lt = list(SD = x1, MAD = x2, CV = x3),
  top_n = 20, col = c("red", "blue", "green"))
correspond_between_two_rankings

Correspond two rankings

**Description**

Correspond two rankings

**Usage**

```r
correspond_between_two_rankings(x1, x2, name1, name2,
    col1 = 2, col2 = 3, top_n = round(0.25*length(x1)), transparency = 0.9,
    pt_size = unit(1, "mm"), newpage = TRUE, ratio = c(1, 1, 1))
```

**Arguments**

- `x1` A vector of scores calculated by one metric.
- `x2` A vector of scores calculated by another metric.
- `name1` Name of the first metric.
- `name2` Name of the second metric.
- `col1` Color for the first metric.
- `col2` Color for the second metric.
- `top_n` Top n elements to show the correspondance.
- `transparency` Transparency of the connecting lines.
- `pt_size` Size of the points, must be a `unit` object.
- `newpage` Whether to plot in a new graphic page.
- `ratio` Ratio of width of the left barplot, connection lines and right barplot. The three values will be scaled to a sum of 1.

**Details**

In `x1` and `x2`, the i\(^{th}\) element in both vectors corresponds to the same object (e.g. same row if they are calculated from a matrix) but with different scores under different metrics.

`x1` and `x2` are sorted in the left panel and right panel respectively. The top n elements under corresponding metric are highlighted by vertical colored lines in both panels. The left and right panels also shown as barplots of the scores in the two metrics. Between the left and right panels, there are lines connecting the same element (e.g. i\(^{th}\) element in `x1` and `x2`) in the two ordered vectors so that you can see how a same element has two different ranks in the two metrics.

Under the plot is a simple Venn diagram showing the overlaps of the top n elements by the two metrics.

**Value**

No value is returned.
Author(s)
Zuguang Gu <z.gu@dkfz.de>

See Also
 correspond_between_rankings draws for more than 2 sets of rankings.

Examples
require(matrixStats)
mat = matrix(runif(1000), ncol = 10)
x1 = rowSds(mat)
x2 = rowMads(mat)
correspond_between_two_rankings(x1, x2, name1 = "SD", name2 = "MAD", top_n = 20)

david_enrichment

Perform DAVID enrichment analysis

Description
Perform DAVID enrichment analysis

Usage


david_enrichment(genes, email,
catalog = c("GOTERM_CC_FAT", "GOTERM_BP_FAT", "GOTERM_MF_FAT", "KEGG_PATHWAY"),
idtype = "ENSEMBL_GENE_ID", species = "Homo sapiens")

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>genes</td>
<td>A vector of gene identifiers.</td>
</tr>
<tr>
<td>email</td>
<td>The email that user registered on DAVID web service (<a href="https://david.ncifcrf.">https://david.ncifcrf.</a></td>
</tr>
<tr>
<td></td>
<td>gov/content.jsp?file=WS.html )</td>
</tr>
<tr>
<td>catalog</td>
<td>A vector of function catalogs. Valid values should be in cola:::DAVID_ALL_CATALOGS.</td>
</tr>
<tr>
<td>idtype</td>
<td>ID types for the input gene list. Valid values should be in cola:::DAVID_ALL_ID_TYPES.</td>
</tr>
<tr>
<td>species</td>
<td>Full species name if the ID type is not uniquely mapped to one single species.</td>
</tr>
</tbody>
</table>

Details
This function directly sends the HTTP request to DAVID web service (https://david.ncifcrf.gov/content.jsp?file=WS.html ) and parses the returned XML. The reason of writing this function is I have problems with other R packages doing DAVID analysis (e.g. RDAVIDWebService, https://bioconductor.org/packages/devel/bioc/html/RDAVIDWebService.html ) because the rJava package RDAVIDWebService depends on can not be installed on my machine.

Users are encouraged to use more advanced gene set enrichment tools such as clusterProfiler (http://www.bioconductor.org/packages/release/bioc/html/clusterProfiler.html ), or fgsea (http://www.bioconductor.org/packages/release/bioc/html/fgsea.html ).

If you want to run this function multiple times, please set time intervals between runs.
Value
   A data frame with functional enrichment results.

Author(s)
   Zuguang Gu <z.gu@dkfz.de>

See Also
   Now cola has a replacement function `functional_enrichment` to perform enrichment analysis.

Examples
   # There is no example
   NULL
### dim.ConsensusPartitionList

**Dimension of the matrix**

**Description**

Dimension of the matrix

**Usage**

```r
## S3 method for class 'ConsensusPartitionList'

dim(x)
```

**Arguments**

- `x` A `ConsensusPartitionList-class` object.

**Examples**

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

---

### dim.DownSamplingConsensusPartition

**Dimension of the matrix**

**Description**

Dimension of the matrix

**Usage**

```r
## S3 method for class 'DownSamplingConsensusPartition'

dim(x)
```

**Arguments**

- `x` A `DownSamplingConsensusPartition-class` object.

**Examples**

```r
# There is no example
NULL
```
**dim.HierarchicalPartition**

*Dimension of the matrix*

**Description**

Dimension of the matrix

**Usage**

```r
## S3 method for class 'HierarchicalPartition'
dim(x)
```

**Arguments**

- `x` A `HierarchicalPartition-class` object.

**Examples**

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

---

**dimension_reduction-ConsensusPartition-method**

*Visualize column after dimension reduction*

**Description**

Visualize samples (the matrix columns) after dimension reduction

**Usage**

```r
## S4 method for signature 'ConsensusPartition'
dimension_reduction(object, k, top_n = NULL,
                   method = c("PCA", "MDS", "t-SNE", "UMAP"),
                   control = list(), color_by = NULL,
                   internal = FALSE, nr = 5000,
                   silhouette_cutoff = 0.5, remove = FALSE,
                   scale_rows = object@scale_rows, verbose = TRUE, ...)
```

---
Arguments

- **object**
  - A `ConsensusPartition-class` object.

- **k**
  - Number of subgroups.

- **top_n**
  - Top n rows to use. By default it uses all rows in the original matrix.

- **method**
  - Which method to reduce the dimension of the data. MDS uses `cmdscale`, PCA uses `prcomp`. t-SNE uses `Rtsne`. UMAP uses `umap`.

- **color_by**
  - If annotation table is set, an annotation name can be set here.

- **control**
  - A list of parameters for `Rtsne` or `umap`.

- **internal**
  - Internally used.

- **nr**
  - If number of matrix rows is larger than this value, random nr rows are used.

- **silhouette_cutoff**
  - Cutoff of silhouette score. Data points with values less than it will be mapped with cross symbols.

- **remove**
  - Whether to remove columns which have less silhouette scores than the cutoff.

- **scale_rows**
  - Whether to perform scaling on matrix rows.

- **verbose**
  - Whether print messages.

- **...**
  - Pass to `dimension_reduction,matrix-method`.

Value

Locations of the points.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

data(golub_cola)
dimension_reduction(golub_cola["ATC", "skmeans"], k = 3)

Description

Method dispatch page for `dimension_reduction`.
Dispatch

dimension_reduction can be dispatched on following classes:

- dimension_reduction,ConsensusPartition-method, ConsensusPartition-class class method
- dimension_reduction,DownSamplingConsensusPartition-method, DownSamplingConsensusPartition-class class method
- dimension_reduction,HierarchicalPartition-method, HierarchicalPartition-class class method
- dimension_reduction,matrix-method, matrix-class class method

Examples

```r
# no example
NULL
```

dimension_reduction-DownSamplingConsensusPartition-method

*Visualize column after dimension reduction*

Description

Visualize samples (the matrix columns) after dimension reduction

Usage

```r
## S4 method for signature 'DownSamplingConsensusPartition'
dimension_reduction(object, k, top_n = NULL,
method = c("PCA", "MDS", "t-SNE", "UMAP"),
control = list(), color_by = NULL,
internal = FALSE, nr = 5000,
p_cutoff = 0.05, remove = FALSE,
scale_rows = TRUE, verbose = TRUE, ...)
```

Arguments

- `object`: A `DownSamplingConsensusPartition-class` object.
- `k`: Number of subgroups.
- `top_n`: Top n rows to use. By default it uses all rows in the original matrix.
- `method`: Which method to reduce the dimension of the data. MDS uses `cmdscale`, PCA uses `prcomp`. t-SNE uses `Rtsne`. UMAP uses `umap`.
- `color_by`: If annotation table is set, an annotation name can be set here.
- `control`: A list of parameters for `Rtsne` or `umap`. 

dimension_reduction-HierarchicalPartition-method

internal

\(nr\)

If number of matrix rows is larger than this value, random \(nr\) rows are used.

p_cutoff

Cutoff of p-value of class label prediction. Data points with values higher than it will be mapped with cross symbols.

remove

Whether to remove columns which have high p-values than the cutoff.

scale_rows

Whether to perform scaling on matrix rows.

verbose

Whether print messages.

Details

This function is basically very similar as \texttt{dimension_reduction,ConsensusPartition-method}.

Value

No value is returned.

Examples

data(golub_cola_ds)
dimension_reduction(golub_cola_ds, k = 2)
dimension_reduction(golub_cola_ds, k = 3)

---

dimension_reduction-HierarchicalPartition-method

\textit{Visualize columns after dimension reduction}

Description

Visualize columns after dimension reduction

Usage

```r
## S4 method for signature 'HierarchicalPartition'
dimension_reduction(object, merge_node = merge_node_param(),
                   parent_node, top_n = NULL, top_value_method = object@list[[1]]@top_value_method,
                   method = c("PCA", "MDS", "t-SNE", "UMAP"), color_by = NULL,
                   scale_rows = object@list[[1]]@scale_rows, verbose = TRUE, ...)
```

Arguments

- **object**
  A \texttt{HierarchicalPartition-class} object.
- **merge_node**
  Parameters to merge sub-dendrograms, see \texttt{merge_node_param}.
- **top_n**
  Top \(n\) rows to use. By default it uses all rows in the original matrix.
- **top_value_method**
  Which top-value method to use.
parent_node Parent node. If it is set, the function call is identical to `dimension_reduction(object[parent_node])`

method Which method to reduce the dimension of the data. MDS uses `cmdscale`, PCA uses `prcomp`, t-SNE uses `Rtsne`, UMAP uses `umap`.

color_by If annotation table is set, an annotation name can be set here.
scale_rows Whether to perform scaling on matrix rows.
verbose Whether print messages.
... Other arguments passed to `dimension_reduction,ConsensusPartition-method`.

Details
The class IDs are extract at depth.

Value
No value is returned.

Author(s)
Zuguang Gu <z.gu@dkfz.de>

Examples
```r
data(golub_cola_rh)
dimension_reduction(golub_cola_rh)
```

---

**dimension_reduction-matrix-method**

*Visualize columns after dimension reduction*

Description
Visualize columns after dimension reduction

Usage
```r
## S4 method for signature 'matrix'
dimension_reduction(object,
pch = 16, col = "black", cex = 1, main = NULL,
method = c("PCA", "MDS", "t-SNE", "UMAP"),
pc = NULL, control = list(),
scale_rows = FALSE, nr = 5000,
internal = FALSE, verbose = TRUE)
```
Arguments

- **object**: A numeric matrix.
- **method**: Which method to reduce the dimension of the data. MDS uses cmdscale, PCA uses prcomp, t-SNE uses Rtsne, UMAP uses umap.
- **pc**: Which two principle components to visualize
- **control**: A list of parameters for Rtsne or umap.
- **pch**: Shape of points.
- **col**: Color of points.
- **cex**: Size of points.
- **main**: Title of the plot.
- **scale_rows**: Whether perform scaling on matrix rows.
- **nr**: If number of matrix rows is larger than this value, random nr rows are used.
- **internal**: Internally used.
- **verbose**: Whether print messages.

Value

- Locations of the points.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

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

---

DownSamplingConsensusPartition-class

The DownSamplingConsensusPartition class

Description

The DownSamplingConsensusPartition class

Details

The DownSamplingConsensusPartition performs consensus partitioning only with a small subset of columns and the class of other columns are predicted by predict_classes, ConsensusPartition-class.

The DownSamplingConsensusPartition-class is a child class of ConsensusPartition-class. It inherits all methods of ConsensusPartition-class.
FCC

Flatness of the CDF curve

Description
Flatness of the CDF curve

Usage
FCC(consensus_mat, diff = 0.1)

Arguments
- consensus_mat: A consensus matrix.
- diff: Difference of F(b) - F(a).

Details
For a in [0, 0.5] and for b in [0.5, 1], the flatness measures the flatness of the CDF curve of the consensus matrix. It is calculated as the maximum width that fits F(b) - F(a) <= diff.

Value
A numeric value.

Examples
```r
data(golub_cola)
FCC(get_consensus(golub_cola[1, 1], k = 2))
FCC(get_consensus(golub_cola[1, 1], k = 3))
FCC(get_consensus(golub_cola[1, 1], k = 4))
FCC(get_consensus(golub_cola[1, 1], k = 5))
FCC(get_consensus(golub_cola[1, 1], k = 6))
```
find_best_km

**Description**

Find a best k for the k-means clustering

**Usage**

```r
find_best_km(mat, max_km = 15)
```

**Arguments**

- `mat`: A matrix where k-means clustering is executed by rows.
- `max_km`: Maximal k to try.

**Details**

The best k is determined by looking for the knee/elbow of the WSS curve (within-cluster sum of square).

Note this function is only for a rough and quick estimation of the best k.

**Examples**

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

---

functional_enrichment-ANY-method

**Description**

Perform functional enrichment on signature genes

**Usage**

```r
# S4 method for signature 'ANY'
functional_enrichment(object, 
  id_mapping = guess_id_mapping(object, org_db, verbose), 
  org_db = "org.Hs.eg.db", ontology = "BP", 
  min_set_size = 10, max_set_size = 1000, 
  verbose = TRUE, prefix = "", ...) 
```
functional_enrichment-ConsensusPartition-method

Arguments

object A vector of gene IDs.

id_mapping If the gene IDs are not Entrez IDs, a named vector should be provided where the names are the gene IDs and values are the corresponding Entrez IDs. The value can also be a function that converts gene IDs.

org_db Annotation database.

ontology Following ontologies are allowed: BP, CC, MF, KEGG, Reactome. MsigDb with the gmt file set by gmt_file argument, or gmt for general gmt gene sets.

min_set_size The minimal size of the gene sets.

max_set_size The maximal size of the gene sets.

verbose Whether to print messages.

prefix Used internally.

... Pass to enrichGO, enrichKEGG, enricher, enrichDO or enrichPathway.

Details

The function enrichment is applied by clusterProfiler, DOSE or ReactomePA packages.

Value

A data frame.

See Also


Examples

# There is no example
NULL

Perform functional enrichment on signature genes
Usage

```r
## S4 method for signature 'ConsensusPartition'
functional_enrichment(object, gene_fdr_cutoff = cola_opt$fdr_cutoff, k = suggest_best_k(object, help = FALSE),
  row_km = NULL, id_mapping = guess_id_mapping(rownames(object), org_db, verbose),
  org_db = "org.Hs.eg.db", ontology = "BP",
  min_set_size = 10, max_set_size = 1000,
  verbose = TRUE, prefix = "", ...)
```

Arguments

- `object`: a `ConsensusPartition-class` object from `run_all_consensus_partition_methods`.
- `gene_fdr_cutoff`: Cutoff of FDR to define significant signature genes.
- `k`: Number of subgroups.
- `row_km`: Number of row clusterings by k-means to separate the matrix that only contains signatures.
- `id_mapping`: If the gene IDs which are row names of the original matrix are not Entrez IDs, a named vector should be provided where the names are the gene IDs in the matrix and values are corresponding Entrez IDs. The value can also be a function that converts gene IDs.
- `org_db`: Annotation database.
- `ontology`: See corresponding argument in `functional_enrichment,ANY-method`.
- `min_set_size`: The minimal size of the gene sets.
- `max_set_size`: The maximal size of the gene sets.
- `verbose`: Whether to print messages.
- `prefix`: Used internally.
- `...`: Pass to `functional_enrichment,ANY-method`.

Details

For how to control the parameters of functional enrichment, see help page of `functional_enrichment,ANY-method`.

Value

A list of data frames which correspond to results for the functional ontologies:

See Also


Examples

```r
# There is no example
NULL
```
Perform functional enrichment on signature genes

Usage

```r
## S4 method for signature 'ConsensusPartitionList'
functional_enrichment(object, gene_fdr_cutoff = cola_opt$fdr_cutoff,
    id_mapping = guess_id_mapping(rownames(object), org_db, FALSE),
    org_db = "org.Hs.eg.db", ontology = "BP",
    min_set_size = 10, max_set_size = 1000, ...)
```

Arguments

- `object`: A `ConsensusPartitionList-class` object from `run_all_consensus_partition_methods`.
- `gene_fdr_cutoff`: Cutoff of FDR to define significant signature genes.
- `id_mapping`: If the gene IDs which are row names of the original matrix are not Entrez IDs, a named vector should be provided where the names are the gene IDs in the matrix and values are corresponding Entrez IDs. The value can also be a function that converts gene IDs.
- `org_db`: Annotation database.
- `ontology`: See corresponding argument in `functional_enrichment,ANY-method`.
- `min_set_size`: The minimal size of the gene sets.
- `max_set_size`: The maximal size of the gene sets.
- `...`: Pass to `functional_enrichment,ANY-method`.

Details

For each method, the signature genes are extracted based on the best k.
It calls `functional_enrichment,ConsensusPartition-method` on the consensus partitioning results for each method.
For how to control the parameters of functional enrichment, see help page of `functional_enrichment,ANY-method`.

Value

A list where each element in the list corresponds to enrichment results from a single method.

See Also

Examples

# There is no example
NULL

---

**functional_enrichment-dispatch**

*Method dispatch page for functional_enrichment*

---

**Description**

Method dispatch page for functional_enrichment.

**Dispatch**

functional_enrichment can be dispatched on following classes:

- `functional_enrichment,HierarchicalPartition-method,HierarchicalPartition-class` class method
- `functional_enrichment,ANY-method,ANY-class` class method
- `functional_enrichment,ConsensusPartition-method,ConsensusPartition-class` class method
- `functional_enrichment,ConsensusPartitionList-method,ConsensusPartitionList-class` class method

**Examples**

# no example
NULL

---

**functional_enrichment-HierarchicalPartition-method**

*Perform functional enrichment on signature genes*

---

**Description**

Perform functional enrichment on signature genes
Usage

```r
## S4 method for signature 'HierarchicalPartition'
functional_enrichment(object, merge_node = merge_node_param(),
gene_fdr_cutoff = cola_opt$fdr_cutoff,
row_km = NULL, id_mapping = guess_id_mapping(rownames(object), org_db, verbose),
org_db = "org.Hs.eg.db", ontology = "BP",
min_set_size = 10, max_set_size = 1000,
verbose = TRUE, ...)
```

Arguments

- **object**: a HierarchicalPartition-class object from hierarchical_partition.
- **merge_node**: Parameters to merge sub-dendrograms, see merge_node_param.
- **gene_fdr_cutoff**: Cutoff of FDR to define significant signature genes.
- **row_km**: Number of row clusterings by k-means to separate the matrix that only contains signatures.
- **id_mapping**: If the gene IDs which are row names of the original matrix are not Entrez IDs, a named vector should be provided where the names are the gene IDs in the matrix and values are corresponding Entrez IDs. The value can also be a function that converts gene IDs.
- **org_db**: Annotation database.
- **ontology**: See corresponding argument in functional_enrichment,ANY-method.
- **min_set_size**: The minimal size of the gene sets.
- **max_set_size**: The maximal size of the gene sets.
- **verbose**: Whether to print messages.
- **...**: Pass to functional_enrichment,ANY-method.

Details

For how to control the parameters of functional enrichment, see help page of functional_enrichment,ANY-method.

Value

A list of data frames which correspond to results for the functional ontologies:

Examples

```r
# There is no example
NULL
```
Description
Get annotations

Usage
## S4 method for signature 'ConsensusPartition'
get_anno(object)

Arguments

object       A ConsensusPartition-class object.

Value
A data frame if anno was specified in run_all_consensus_partition_methods or consensus_partition, or else NULL.

Author(s)
Zuguang Gu <z.gu@dkfz.de>

Examples

# There is no example
NULL

Description
Get annotations

Usage
## S4 method for signature 'ConsensusPartitionList'
get_anno(object)
Arguments

object A ConsensusPartitionList-class object.

Value

A data frame if anno was specified in run_all_consensus_partition_methods, or else NULL.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

# There is no example
NULL
get_anno-DownSamplingConsensusPartition-method

Description
Get annotations

Usage
## S4 method for signature 'DownSamplingConsensusPartition'
get_anno(object, reduce = FALSE)

Arguments
object A DownSamplingConsensusPartition-class object.
reduce Used internally.

Value
A data frame if anno was specified in consensus_partition_by_down_sampling, or else NULL.

Author(s)
Zuguang Gu <z.gu@dkfz.de>

Examples
data(golub_cola_ds)
get_anno(golub_cola_ds)

get_anno-HierarchicalPartition-method

Description
Get annotations

Usage
## S4 method for signature 'HierarchicalPartition'
get_anno(object)

Arguments
object A HierarchicalPartition-class object.
get_anno_col-ConsensusPartition-method

Value

A data frame if anno was specified in hierarchical_partition, or NULL.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

# There is no example
NULL
**Description**

Get annotation colors

**Usage**

```r
## S4 method for signature 'ConsensusPartitionList'
get_anno_col(object)
```

**Arguments**

- `object` A `ConsensusPartitionList-class` object.

**Value**

A list of color vectors or else NULL.

**Author(s)**

Zuguang Gu <z.gu@dkfz.de>

**Examples**

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

---

**get_anno_col-dispatch**  
*Method dispatch page for get_anno_col*

**Description**

Method dispatch page for `get_anno_col`.

**Dispatch**

`get_anno_col` can be dispatched on following classes:

- `get_anno_col,HierarchicalPartition-method,HierarchicalPartition-class` class method
- `get_anno_col,ConsensusPartitionList-method,ConsensusPartitionList-class` class method
- `get_anno_col,ConsensusPartition-method,ConsensusPartition-class` class method
get_anno_col-HierarchicalPartition-method

Get annotation colors

Description

Get annotation colors

Usage

## S4 method for signature 'HierarchicalPartition'
get_anno_col(object)

Arguments

object A `HierarchicalPartition-class` object.

Value

A list of color vectors or NULL.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

# There is no example
NULL
### get_children_nodes-HierarchicalPartition-method

Test whether a node is a leaf node

---

**Description**

Test whether a node is a leaf node

**Usage**

```r
## S4 method for signature 'HierarchicalPartition'
geet_children_nodes(object, node, merge_node = merge_node_param())
```

**Arguments**

- `node`: A vector of node IDs.
- `merge_node`: Parameters to merge sub-dendrograms, see `merge_node_param`.

**Value**

A vector of children nodes.

**Examples**

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

---

### get_classes-ConsensusPartition-method

Get subgroup labels

---

**Description**

Get subgroup labels

**Usage**

```r
## S4 method for signature 'ConsensusPartition'
get_classes(object, k = object@k)
```

**Arguments**

- `k`: Number of subgroups.
Value

A data frame with subgroup labels and other columns which are entropy of the percent membership matrix and the silhouette scores which measure the stability of a sample to stay in its group.

If k is not specified, it returns a data frame with subgroup labels from all k.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

data(golub_cola)
obj = golub_cola[“ATC”, “skmeans”]
get_classes(obj, k = 2)
get_classes(obj)

Description

Get subgroup labels

Usage

## S4 method for signature 'ConsensusPartitionList'
get_classes(object, k)

Arguments

object A ConsensusPartitionList-class object.
k Number of subgroups.

Details

The subgroup labels are inferred by merging partitions from all methods by weighting the mean silhouette scores in each method.

Value

A data frame with subgroup labels and other columns which are entropy of the percent membership matrix and the silhouette scores which measure the stability of a sample to stay in its group.

Author(s)

Zuguang Gu <z.gu@dkfz.de>
Examples

data(golub_cola)
get_classes(golub_cola, k = 2)

get_classes-dispatch  Method dispatch page for get_classes

Description

Method dispatch page for get_classes.

Dispatch

get_classes can be dispatched on following classes:

- `get_classes,HierarchicalPartition-method`, `HierarchicalPartition-class` class method
- `get_classes,ConsensusPartitionList-method`, `ConsensusPartitionList-class` class method
- `get_classes,ConsensusPartition-method`, `ConsensusPartition-class` class method
- `get_classes,DownSamplingConsensusPartition-method`, `DownSamplingConsensusPartition-class` class method

Examples

# no example
NULL

get_classes-DownSamplingConsensusPartition-method

Getting subgroup labels

Description

Get subgroup labels

Usage

## S4 method for signature 'DownSamplingConsensusPartition'
get_classes(object, k = object@k, p_cutoff = 0.05, reduce = FALSE)
get_classes-HierarchicalPartition-method

Arguments

- object: A `DownSamplingConsensusPartition-class` object.
- k: Number of subgroups.
- p_cutoff: Cutoff of p-values of class label prediction. It is only used when k is a vector.
- reduce: Used internally.

Value

If k is a scalar, it returns a data frame with two columns:
- the class labels
- the p-value for the prediction of class labels.

If k is a vector, it returns a data frame of class labels for each k. The class label with prediction p-value > p_cutoff is set to NA.

Examples

```r
data(golub_cola_ds)
get_classes(golub_cola_ds, k = 3)
get_classes(golub_cola_ds)
```

Description

Get class IDs from the HierarchicalPartition object

Usage

```r
## S4 method for signature 'HierarchicalPartition'
get_classes(object, merge_node = merge_node_param())
```

Arguments

- merge_node: Parameters to merge sub-dendrograms, see `merge_node_param`.

Value

A data frame of classes IDs. The class IDs are the node IDs where the subgroup sits in the hierarchy.

Author(s)

Zuguang Gu <z.gu@dkfz.de>
**Examples**

```r
data(golub_cola)
get_classes(golub_cola)
```

---

**Description**

Get consensus matrix

**Usage**

```r
## S4 method for signature 'ConsensusPartition'
get_consensus(object, k)
```

**Arguments**

- `object` A `ConsensusPartition-class` object.
- `k` Number of subgroups.

**Details**

For row i and column j in the consensus matrix, the value of corresponding x_ij is the probability of sample i and sample j being in the same group from all partitions.

**Value**

A consensus matrix corresponding to the current k.

**Author(s)**

Zuguang Gu <z.gu@dkfz.de>

**Examples**

```r
data(golub_cola)
obj = golub_cola["ATC", "skmeans"]
get_consensus(obj, k = 2)
```
get_matrix-ConsensusPartition-method

Description

Get the original matrix

Usage

## S4 method for signature 'ConsensusPartition'
get_matrix(object, full = FALSE, include_all_rows = FALSE)

Arguments

- `full`: Whether to extract the complete original matrix.
- `include_all_rows`: Internally used.

Value

A numeric matrix.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

data(golub_cola)
obj = golub_cola["ATC", "skmeans"]
get_matrix(obj)

get_matrix-ConsensusPartitionList-method

Description

Get the original matrix

Usage

## S4 method for signature 'ConsensusPartitionList'
get_matrix(object)

Description

Get the original matrix

Usage

## S4 method for signature 'ConsensusPartitionList'
get_matrix(object)
get_matrix-dispatch

Arguments

object  A `ConsensusPartitionList-class` object.

Value

A numeric matrix.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

```r
data(golub_cola)
get_matrix(golub_cola)
```

Description

Method dispatch page for `get_matrix`.

Dispatch

`get_matrix` can be dispatched on following classes:

- `get_matrix,ConsensusPartition-method,ConsensusPartition-class` class method
- `get_matrix,ConsensusPartitionList-method,ConsensusPartitionList-class` class method
- `get_matrix,DownSamplingConsensusPartition-method,DownSamplingConsensusPartition-class` class method
- `get_matrix,HierarchicalPartition-method,HierarchicalPartition-class` class method

Examples

```r
# no example
NULL
```
Description
Get the original matrix

Usage
```r
## S4 method for signature 'DownSamplingConsensusPartition'
get_matrix(object, reduce = FALSE)
```

Arguments
- `object`: A `DownSamplingConsensusPartition-class` object.
- `reduce`: Whether to return the reduced matrix where columns are randomly sampled.

Value
A numeric matrix

Examples
```r
# There is no example
NULL
```

Description
Get the original matrix

Usage
```r
## S4 method for signature 'HierarchicalPartition'
get_matrix(object)
```

Arguments
get_membership-ConsensusPartition-method

Value

A numeric matrix.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

# There is no example
NULL

Description

Get membership matrix

Usage

## S4 method for signature 'ConsensusPartition'
get_membership(object, k, each = FALSE)

Arguments

object A ConsensusPartition-class object.
k Number of subgroups.
each Whether to return the percentage membership matrix which is summarized from all partitions or the individual membership in every single partition run.

Details

If each == FALSE, the value in the membership matrix is the probability to be in one subgroup, while if each == TRUE, the membership matrix contains the subgroup labels for every single partitions which are from randomly sampling from the original matrix.

The percent membership matrix is calculated by cl_consensus.

Value

- If each == FALSE, it returns a membership matrix where rows correspond to the columns from the subgroups.
- If each == TRUE, it returns a membership matrix where rows correspond to the columns from the original matrix.
Author(s)
Zuguang Gu <z.gu@dkfz.de>

See Also

get_membership, ConsensusPartitionList-method summarizes membership from partitions from all combinations of top-value methods and partitioning methods.

Examples

data(golub_cola)
obj = golub_cola["ATC", "skmeans"]
get_membership(obj, k = 2)
get_membership(obj, k = 2, each = TRUE)

Description

Get membership matrix

Usage

## S4 method for signature 'ConsensusPartitionList'
get_membership(object, k)

Arguments

  object   A ConsensusPartitionList-class object.
  k        Number of subgroups.

Details

The membership matrix (the probability of each sample to be in one subgroup, if assuming columns represent samples) is inferred from the consensus partition of every combination of methods, weighted by the mean silhouette score of the partition for each method. So methods which give unstable partitions have lower weights when summarizing membership matrix from all methods.

Value

A membership matrix where rows correspond to the columns in the original matrix.

Author(s)

Zuguang Gu <z.gu@dkfz.de>
See Also

`get_membership,ConsensusPartition-method` returns membership matrix for a single top-value method and partitioning method.

Examples

```r
data(golub_cola)
get_membership(golub_cola, k = 2)
```

Description

Method dispatch page for `get_membership`.

Dispatch

`get_membership` can be dispatched on following classes:

- `get_membership,ConsensusPartition-method,ConsensusPartition-class` class method
- `get_membership,ConsensusPartitionList-method,ConsensusPartitionList-class` class method

Examples

```r
# no example
NULL
```

Description

Get parameters

Usage

```r
## S4 method for signature 'ConsensusPartition'
get_param(object, k = object@k, unique = TRUE)
```
get_signatures-ConsensusPartition-method

Arguments

object A ConsensusPartition-class object.
k Number of subgroups.
unique Whether to apply unique to rows of the returned data frame.

Details

It is mainly used internally.

Value

A data frame of parameters corresponding to the current k. In the data frame, each row corresponds to a partition run.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

```r
data(golub_cola)
obj = golub_cola["ATC", "skmeans"]
get_param(obj)
get_param(obj, k = 2)
get_param(obj, unique = FALSE)
```

get_signatures-ConsensusPartition-method

Get signature rows

Description

Get signature rows

Usage

```r
## S4 method for signature 'ConsensusPartition'
get_signatures(object, k,
    col = if(scale_rows) c("green", "white", "red") else c("blue", "white", "red"),
    silhouette_cutoff = 0.5,
    fdr_cutoff = cola_opt$fdr_cutoff,
    top_signatures = NULL,
    group_diff = cola_opt$group_diff,
    scale_rows = object@scale_rows, .scale_mean = NULL, .scale_sd = NULL,
    row_km = NULL,
    diff_method = c("Ftest", "ttest", "samr", "pamr", "one_vs_others", "uniquely_high_in_one_group"),
    anno = get_anno(object),
```

get_signatures-ConsensusPartition-method

anno_col = get_anno_col(object),
internal = FALSE,
show_row_dend = FALSE,
show_column_names = FALSE,
column_names_gp = gpar(fontsize = 8),
use_raster = TRUE,
plot = TRUE, verbose = TRUE, seed = 888,
left_annotation = NULL, right_annotation = NULL,
simplify = FALSE, prefix = "", enforce = FALSE, hash = NULL, from_hc = FALSE,
...)

Arguments

object A ConsensusPartition-class object.
k Number of subgroups.
col Colors for the main heatmap.
silhouette_cutoff Cutoff for silhouette scores. Samples with values less than it are not used for finding signature rows. For selecting a proper silhouette cutoff, please refer to https://www.stat.berkeley.edu/~s133/Cluster2a.html#tth_tAb1.
fdr_cutoff Cutoff for FDR of the difference test between subgroups.
top_signatures Top signatures with most significant fdr. Note since fdr might be same for multiple rows, the final number of signatures might not be exactly the same as the one that has been set.
grup_diff Cutoff for the maximal difference between group means.
scale_rows Whether apply row scaling when making the heatmap.
.scale_mean Internally used.
.scale_sd Internally used.
row_km Number of groups for performing k-means clustering on rows. By default it is automatically selected.
diff_method Methods to get rows which are significantly different between subgroups, see 'Details' section.
anno A data frame of annotations for the original matrix columns. By default it uses the annotations specified in consensus_partition or run_all_consensus_partition_methods.
anno_col A list of colors (color is defined as a named vector) for the annotations. If anno is a data frame, anno_col should be a named list where names correspond to the column names in anno.
internal Used internally.
show_row_dend Whether show row dendrogram.
show_column_names Whether show column names in the heatmap.
column_names_gp Graphics parameters for column names.
use_raster Internally used.
plot  Whether to make the plot.
verbose Whether to print messages.
seed  Random seed.
left_annotation  Annotation put on the left of the heatmap. It should be a HeatmapAnnotation-class object. The number of items should be the same as the number of the original matrix rows. The subsetting to the significant rows are automatically performed on the annotation object.
right_annotation  Annotation put on the right of the heatmap. Same format as left_annotation.
simplify Only used internally.
prefix Only used internally.
enforce The analysis is cached by default, so that the analysis with the same input will be automatically extracted without rerunning them. Set enforce to TRUE to enforce the function to re-perform the analysis.
hash Use internally.
from_hc Is the ConsensusPartition-class object a node of a HierarchicalPartition object?
... Other arguments.

Details

Basically the function applies statistical test for the difference in subgroups for every row. There are following methods which test significance of the difference:

- **ttest**  First it looks for the subgroup with highest mean value, compare to each of the other subgroups with t-test and take the maximum p-value. Second it looks for the subgroup with lowest mean value, compare to each of the other subgroups again with t-test and take the maximum p-values. Later for these two list of p-values take the minimal p-value as the final p-value.
- **samr/pamr**  use SAM (from samr package)/PAM (from pamr package) method to find significantly different rows between subgroups.
- **Ftest**  use F-test to find significantly different rows between subgroups.
- **one_vs_others**  For each subgroup i in each row, it uses t-test to compare samples in current subgroup to all other samples, denoted as p_i. The p-value for current row is selected as min(p_i).
- **uniquely_high_in_one_group**  The signatures are defined as, if they are uniquely up-regulated in subgroup A, then it must fit following criterions: 1. in a two-group t-test of A ~ other_merged_groups, the statistic must be > 0 (high in group A) and p-value must be significant, and 2. for other groups (excluding A), t-test in every pair of groups should not be significant.

diff_method can also be a self-defined function. The function needs two arguments which are the matrix for the analysis and the predicted classes. The function should returns a vector of FDR from the difference test.
Value

A data frame with more than two columns:

which_row: row index corresponding to the original matrix.
fdr: the FDR.
km: the k-means groups if row_km is set.
other_columns: the mean value (depending rows are scaled or not) in each subgroup.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

data(golub_cola)
res = golub_cola["ATC", "skmeans"]
tb = get_signatures(res, k = 3)
head(tb)
get_signatures(res, k = 3, top_signatures = 100)

Description

Method dispatch page for get_signatures.

Dispatch

get_signatures can be dispatched on following classes:

- get_signatures,ConsensusPartition-method,ConsensusPartition-class class method
- get_signatures,DownSamplingConsensusPartition-method,DownSamplingConsensusPartition-class class method
- get_signatures,HierarchicalPartition-method,HierarchicalPartition-class class method

Examples

# no example
NULL
Description

Get signature rows

Usage

```r
## S4 method for signature 'DownSamplingConsensusPartition'
get_signatures(object, k,
               p_cutoff = 1, ...)
```

Arguments

- `object`: A `DownSamplingConsensusPartition-class` object.
- `k`: Number of subgroups.
- `p_cutoff`: Cutoff for p-values of class label prediction. Samples with values higher than it are not used for finding signature rows.
- `...`: Other arguments passed to `get_signatures,ConsensusPartition-method`.

Details

This function is very similar as `get_signatures,ConsensusPartition-method`.

Examples

```r
data(golub_cola_ds)
get_signatures(golub_cola_ds, k = 2)
gt_signatures(golub_cola_ds, k = 3)
```
get_signatures-HierarchicalPartition-method

Usage

```r
## S4 method for signature 'HierarchicalPartition'
get_signatures(object, merge_node = merge_node_param(),
               group_diff = object@param$group_diff,
               row_km = NULL, diff_method = "Ftest", fdr_cutoff = object@param$fdr_cutoff,
               scale_rows = object[[1]]@scale_rows,
               anno = get_anno(object),
               anno_col = get_anno_col(object),
               show_column_names = FALSE, column_names_gp = gpar(fontsize = 8),
               verbose = TRUE, plot = TRUE, seed = 888,
               ...)```

Arguments

- **object**: a `HierarchicalPartition-class` object.
- **merge_node**: Parameters to merge sub-dendrograms, see `merge_node_param`.
- **group_diff**: Cutoff for the maximal difference between group means.
- **row_km**: Number of groups for performing k-means clustering on rows. By default it is automatically selected.
- **diff_method**: Methods to get rows which are significantly different between subgroups.
- **fdr_cutoff**: Cutoff for FDR of the difference test between subgroups.
- **scale_rows**: whether apply row scaling when making the heatmap.
- **anno**: a data frame of annotations for the original matrix columns. By default it uses the annotations specified in `hierarchical_partition`.
- **anno_col**: a list of colors (color is defined as a named vector) for the annotations. If `anno` is a data frame, `anno_col` should be a named list where names correspond to the column names in `anno`.
- **show_column_names**: whether show column names in the heatmap.
- **column_names_gp**: Graphic parameters for column names.
- **verbose**: whether to print messages.
- **plot**: whether to make the plot.
- **seed**: Random seed.
- **...**: other arguments pass to `get_signatures,ConsensusPartition-method`.

Details

The function calls `get_signatures,ConsensusPartition-method` to find signatures at each node of the partition hierarchy.
Value

A data frame with more than two columns:

- **which_row**: row index corresponding to the original matrix.
- **km**: the k-means groups if `row_km` is set.
- **other_columns**: the mean value (depending rows are scaled or not) in each subgroup.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

```r
data(golub_cola_rh)
tb = get_signatures(golub_cola_rh)
head(tb)
```

Description

Get statistics

Usage

```r
## S4 method for signature 'ConsensusPartition'
get_stats(object, k = object@k, all_stats = FALSE)
```

Arguments

- **object**: A `ConsensusPartition-class` object.
- **k**: Number of subgroups. The value can be a vector.
- **all_stats**: Whether to show all statistics that were calculated. Used internally.

Details

The statistics are:

- **1-PAC**: 1 - proportion of ambiguous clustering, calculated by `PAC`.
- **concordance**: The mean probability that each partition fits the consensus partition, calculated by `concordance`. 
**area_increased**  The increased area under eCDF (the empirical cumulative distribution function) curve to the previous $k$.

**Rand**  This is the percent of pairs of samples that are both in a same cluster or both are not in a same cluster in the partition of $k$ and $k-1$. See [https://en.wikipedia.org/wiki/Rand_index](https://en.wikipedia.org/wiki/Rand_index).

**Jaccard**  The ratio of pairs of samples are both in a same cluster in the partition of $k$ and $k-1$ and the pairs of samples are both in a same cluster in the partition $k$ or $k-1$.

Value

A matrix of partition statistics.

Author(s)

Zuguang Gu &lt;z.gu@dkfz.de&gt;

Examples

```r
data(golub_cola)
obj = golub_cola["ATC", "skmeans"]
get_stats(obj)
get_stats(obj, k = 2)
```

Description

Get statistics

Usage

```r
## S4 method for signature 'ConsensusPartitionList'
get_stats(object, k, all_stats = FALSE)
```

Arguments

- **object**  A `ConsensusPartitionList-class` object.
- **k**  Number of subgroups. The value can only be a single value.
- **all_stats**  Whether to show all statistics that were calculated. Used internally.

Value

A matrix of partition statistics for a selected $k$. Rows in the matrix correspond to combinations of top-value methods and partitioning methods.
Author(s)
Zuguang Gu <z.gu@dkfz.de>

Examples

```r
data(golub_cola)
get_stats(golub_cola, k = 2)
```

---

get_stats-dispatch  

**Method dispatch page for get_stats**

Description

Method dispatch page for get_stats.

Dispatch

get_stats can be dispatched on following classes:

- `get_stats,ConsensusPartitionList-method,ConsensusPartitionList-class` class method
- `get_stats,ConsensusPartition-method,ConsensusPartition-class` class method

Examples

```r
# no example
NULL
```

---

golub_cola  

**Example ConsensusPartitionList object from Golub dataset**

Description

Example ConsensusPartitionList object from Golub dataset

Usage

```r
data(golub_cola)
```
Details

Following code was used to generate golub_cola:

```r
library(cola)

library(golubEsets)  # from bioc
data(Golub_Merge)
m = exprs(Golub_Merge)
colnames(m) = paste0("sample_", colnames(m))
anno = pData(Golub_Merge)

m[m <= 1] = NA
m = log10(m)
m = adjust_matrix(m)

library(preprocessCore)  # from bioc
cn = colnames(m)
rn = rownames(m)
m = normalize.quantiles(m)
colnames(m) = cn
rownames(m) = rn

set.seed(123)
golub_cola = run_all_consensus_partition_methods(
m, cores = 6,
anno = anno[, c("ALL.AML"), drop = FALSE],
anno_col = c("ALL" = "red", "AML" = "blue")
)
```

Author(s)

Zuguang Gu <z.gu@dkfz.de>

See Also

https://jokergoo.github.io/cola_examples/Golub_leukemia/

Examples

```r
data(golub_cola)
golub_cola
```
Example DownSamplingConsensusPartition object from Golub dataset

Usage

data(golub_cola_ds)

Details

Following code was used to generate golub_cola_ds:

```r
library(cola)
data(golub_cola)
m = get_matrix(golub_cola)
set.seed(123)
golub_cola_ds = consensus_partition_by_down_sampling(
  m, subset = 50, cores = 6,
  anno = get_anno(golub_cola),
  anno_col = get_anno_col(golub_cola),
)
```

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

data(golub_cola_ds)
golub_cola_ds

golub_cola_rh Example HierarchicalPartition object from Golub dataset

Description

Example HierarchicalPartition object from Golub dataset

Usage

data(golub_cola_rh)
Details

Following code was used to generate golub_cola_rh:

```r
library(cola)
data(golub_cola)
m = get_matrix(golub_cola)
set.seed(123)
golub_cola_rh = hierarchical_partition(
  m, cores = 6,
  anno = get_anno(golub_cola),
  anno_col = get_anno_col(golub_cola)
)
```

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

```r
data(golub_cola_rh)
golub_cola_rh
```

HierarchicalPartition-class

The HierarchicalPartition class

Description

The HierarchicalPartition class

Methods

The HierarchicalPartition-class has following methods:

- `hierarchical_partition`: constructor method.
- `collect_classes,HierarchicalPartition-method`: plot the hierarchy of subgroups predicted.
- `get_classes,HierarchicalPartition-method`: get the class IDs of subgroups.
- `suggest_best_k,HierarchicalPartition-method`: guess the best number of partitions for each node.
- `get_matrix,HierarchicalPartition-method`: get the original matrix.
- `get_signatures,HierarchicalPartition-method`: get the signatures for each subgroup.
- `compare_signatures,HierarchicalPartition-method`: compare signatures from different nodes.
- `test_to_known_factors,HierarchicalPartition-method`: test correlation between predicted subgrouping and known annotations, if available.
Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

# There is no example
NULL

hierarchical_partition

Hierarchical partition

Description

Hierarchical partition

Usage

hierarchical_partition(data, 
  top_n = NULL, 
  top_value_method = "ATC", 
  partition_method = "skmeans", 
  combination_method = expand.grid(top_value_method, partition_method), 
  anno = NULL, anno_col = NULL, 
  mean_silhouette_cutoff = 0.9, min_samples = max(6, round(ncol(data)*0.01)), 
  subset = Inf, predict_method = "centroid", 
  group_diff = ifelse(scale_rows, 0.5, 0), 
  fdr_cutoff = cola_opt$fdr_cutoff, 
  min_n_signatures = NULL, 
  filter_fun = function(mat) { 
    s = rowSds(mat) 
    s > quantile(unique(s[s > 1e-10]), 0.05, na.rm = TRUE) 
  }, 
  max_k = 4, scale_rows = TRUE, verbose = TRUE, mc.cores = 1, cores = mc.cores, help = TRUE, ...)

Arguments

data         a numeric matrix where subgroups are found by columns.
top_n        Number of rows with top values.
top_value_method a single or a vector of top-value methods. Available methods are in all_top_value_methods.
partition_method a single or a vector of partition methods. Available methods are in all_partition_methods.
**combination_method**

A list of combinations of top-value methods and partitioning methods. The value can be a two-column data frame where the first column is the top-value methods and the second column is the partitioning methods. Or it can be a vector of combination names in a form of "top_value_method:partitioning_method".

**anno**

A data frame with known annotation of samples. The annotations will be plotted in heatmaps and the correlation to predicted subgroups will be tested.

**anno_col**

A list of colors (color is defined as a named vector) for the annotations. If anno is a data frame, anno_col should be a named list where names correspond to the column names in anno.

**mean_silhouette_cutoff**

The cutoff to test whether partition in current node is stable.

**min_samples**

The cutoff of number of samples to determine whether to continue looking for subgroups.

**group_diff**

Pass to `get_signatures,ConsensusPartition-method`.

**fdr_cutoff**

Pass to `get_signatures,ConsensusPartition-method`.

**subset**

Number of columns to randomly sample.

**predict_method**

Method for predicting class labels. Possible values are "centroid", "svm" and "randomForest".

**min_n_signatures**

Minimal number of signatures under the best classification.

**filter_fun**

A self-defined function which filters the original matrix and returns a submatrix for partitioning.

**max_k**

Maximal number of partitions to try. The function will try 2:max_k partitions. Note this is the number of partitions that will be tried out on each node of the hierarchical partition. Since more subgroups will be found in the whole partition hierarchy, on each node, max_k should not be set to a large value.

**scale_rows**

Whether rows are scaled?

**verbose**

Whether print message.

**mc.cores**

Multiple cores to use. This argument will be removed in future versions.

**cores**

Number of cores, or a cluster object returned by `makeCluster`.

**help**

Whether to show the help message.

**...**

Pass to `consensus_partition`.

**Details**

The function looks for subgroups in a hierarchical way.

There is a special way to encode the node in the hierarchy. The length of the node name is the depth of the node in the hierarchy and the substring excluding the last digit is the name node of the parent node. E.g. for the node 0011, the depth is 4 and the parent node is 001.

**Value**

A `HierarchicalPartition-class` object. Simply type object in the interactive R session to see which functions can be applied on it.
is_best_k-ConsensusPartition-method

Test whether the current k is the best/optional k

Description

Test whether the current k is the best/optional k

Usage

## S4 method for signature 'ConsensusPartition'
is_best_k(object, k, ...)

Arguments

object A ConsensusPartition-class object.
k Number of subgroups.
... Pass to suggest_best_k,ConsensusPartition-method.

Details

Optional best k is also assigned as TRUE.

Value

Logical scalar.

Examples

```r
## Not run:
set.seed(123)
m = cbind(rbind(matrix(rnorm(20*20, mean = 2, sd = 0.3), nr = 20),
               matrix(rnorm(20*20, mean = 0, sd = 0.3), nr = 20),
               matrix(rnorm(20*20, mean = 0, sd = 0.3), nr = 20)),
          rbind(matrix(rnorm(20*20, mean = 0, sd = 0.3), nr = 20),
                 matrix(rnorm(20*20, mean = 1, sd = 0.3), nr = 20),
                 matrix(rnorm(20*20, mean = 0, sd = 0.3), nr = 20)),
          rbind(matrix(rnorm(20*20, mean = 0, sd = 0.3), nr = 20),
                 matrix(rnorm(20*20, mean = 0, sd = 0.3), nr = 20),
                 matrix(rnorm(20*20, mean = 1, sd = 0.3), nr = 20))
) + matrix(rnorm(60*60, sd = 0.5), nr = 60)
rh = hierarchical_partition(m, top_value_method = "SD", partition_method = "kmeans")
## End(Not run)
```
is_best_k-ConsensusPartitionList-method

Examples

```r
data(golub_cola)
obj = golub_cola["ATC", "skmeans"]
is_best_k(obj, k = 2)
is_best_k(obj, k = 3)
```

is_best_k-ConsensusPartitionList-method
Test whether the current k is the best/optional k

Description

Test whether the current k is the best/optional k

Usage

```r
## S4 method for signature 'ConsensusPartitionList'
is_best_k(object, k, ...)
```

Arguments

- `object`: A `ConsensusPartitionList-class` object.
- `k`: Number of subgroups.
- `...`: Pass to `suggest_best_k`, `ConsensusPartitionList-method`.

Details

It tests on the partitions for every method.

Value

Logical vector.

Examples

```r
data(golub_cola)
is_best_k(golub_cola, k = 3)
```
is_best_k-method

**Description**
Method dispatch page for is_best_k.

**Dispatch**
is_best_k can be dispatched on following classes:

- `is_best_k,ConsensusPartition-method`, `ConsensusPartition-class` class method
- `is_best_k,ConsensusPartitionList-method`, `ConsensusPartitionList-class` class method

**Examples**

```r
# no example
NULL
```

is_leaf_node-HierarchicalPartition-method

**Description**
Test whether a node is a leaf node

**Usage**

```r
## S4 method for signature 'HierarchicalPartition'
is_leaf_node(object, node, merge_node = merge_node_param())
```

**Arguments**

- `object` A `HierarchicalPartition-class` object.
- `node` A vector of node IDs.
- `merge_node` Parameters to merge sub-dendrograms, see `merge_node_param`.

**Examples**

```r
data(golub_cola_rh)
is_leaf_node(golub_cola_rh, all_leaves(golub_cola_rh))
```
is_stable_k-ConsensusPartition-method

Test whether the current k corresponds to a stable partition

Description

Test whether the current k corresponds to a stable partition

Usage

## S4 method for signature 'ConsensusPartition'

is_stable_k(object, k, stable_PAC = 0.1, ...)

Arguments

object

A ConsensusPartition-class object.

k

Number of subgroups.

stable_PAC

Cutoff for stable PAC.

...

Pass to suggest_best_k,ConsensusPartition-method.

Details

if 1-PAC for the k is larger than 0.9 (10% ambiguity for the partition), cola marks it as a stable partition.

Value

Logical scalar.

Examples

data(golub_cola)
obj = golub_cola["ATC", "skmeans"]
is_stable_k(obj, k = 2)
is_stable_k(obj, k = 3)
is_stable_k-ConsensusPartitionList-method

Test whether the current k corresponds to a stable partition

Description
Test whether the current k corresponds to a stable partition

Usage
## S4 method for signature 'ConsensusPartitionList'
is_stable_k(object, k, ...)

Arguments
object A ConsensusPartitionList-class object.
k Number of subgroups.
... Pass to suggest_best_k,ConsensusPartitionList-method.

Details
It tests on the partitions for every method.

Value
Logical vector

Examples
data(golub_cola)
is_stable_k(golub_cola, k = 3)

is_stable_k-dispatch Method dispatch page for is_stable_k

Description
Method dispatch page for is_stable_k.

Dispatch
is_stable_k can be dispatched on following classes:

• is_stable_k,ConsensusPartitionList-method, ConsensusPartitionList-class class method
• is_stable_k,ConsensusPartition-method, ConsensusPartition-class class method
knee_finder2

Examples

# no example
NULL

knee_finder2  Find the knee/elbow of a list of sorted points

Description

Find the knee/elbow of a list of sorted points

Usage

knee_finder2(x, plot = FALSE)

Arguments

x  A numeric vector.
plot  Whether to make the plot.

Value

A vector of two numeric values. One for the left knee and the second for the right knee.

Examples

x = rnorm(1000)
knee_finder2(x, plot = TRUE)

knitr_add_tab_item  Add JavaScript tab in the report

Description

Add JavaScript tab in the report

Usage

knitr_add_tab_item(code, header, prefix, desc = "", opt = NULL, message = NULL, hide_and_show = FALSE)
Arguments

- code: R code to execute.
- header: Header or the title for the tab.
- prefix: Prefix of the chunk label.
- desc: Description in the tab.
- opt: Options for the knitr chunk.
- message: Message to print.
- hide_and_show: Whether to hide the code output.

Details

Each tab contains the R source code and results generated from it (figure, tables, text, ...).
This function is only for internal use.

Value

No value is returned.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

See Also

knitr_insert_tabs produces a complete HTML fragment.

Examples

# There is no example
NULL

---

**knitr_insert_tabs**  
*Generate the HTML fragment for the JavaScript tabs*

Description

Generate the HTML fragment for the JavaScript tabs

Usage

knitr_insert_tabs(uid)

Arguments

- uid: A unique identifier for the div.
map_to_entrez_id

Details

The jQuery UI is used to generate html tabs (https://jqueryui.com/tabs/).

knitr_insert_tabs should be used after several calls of knitr_add_tab_item to generate a complete HTML fragment for all tabs with all necessary Javascript and css code.

This function is only for internal use.

Value

No value is returned.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

# There is no example
NULL

map_to_entrez_id Map to Entrez IDs

Description

Map to Entrez IDs

Usage

map_to_entrez_id(from, org_db = "org.Hs.eg.db")

Arguments

from The input gene ID type. Valid values should be in, e.g. columns(org.Hs.eg.db::org.Hs.eg.db).
org_db The annotation database.

Details

If there are multiple mappings from the input ID type to an unique Entrez ID, it randomly picks one.

Value

A named vectors where names are IDs with input ID type and values are the Entrez IDs.

The returned object normally is used in functional_enrichment.
Examples

```r
map = map_to_entrez_id("ENSEMBL")
head(map)
```

Description

Max depth of the hierarchy

Usage

```r
## S4 method for signature 'HierarchicalPartition'
max_depth(object)
```

Arguments

- **object**: A `HierarchicalPartition-class` object.

Value

A numeric value.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

```r
data(golub_cola_rh)
max_depth(golub_cola_rh)
```
Heatmap of membership in each partition

Description
Heatmap of membership in each partition

Usage
```r
## S4 method for signature 'ConsensusPartition'
membership_heatmap(object, k, internal = FALSE,
                   anno = object@anno, anno_col = get_anno_col(object),
                   show_column_names = FALSE, column_names_gp = gpar(fontsize = 8), ...)
```

Arguments
- `k`: Number of subgroups.
- `internal`: Used internally.
- `anno`: A data frame of annotations for the original matrix columns. By default it uses the annotations specified in `consensus_partition` or `run_all_consensus_partition_methods`.
- `anno_col`: A list of colors (color is defined as a named vector) for the annotations. If `anno` is a data frame, `anno_col` should be a named list where names correspond to the column names in `anno`.
- `show_column_names`: Whether show column names in the heatmap (which is the column name in the original matrix).
- `column_names_gp`: Graphics parameters for column names.
- `...`: Other arguments.

Details
Each row in the heatmap is the membership in one single partition.
Heatmap is split on rows by `top_n`.

Value
No value is returned.

Author(s)
Zuguang Gu <z.gu@dkfz.de>
Examples

```r
data(golub_cola)
membership_heatmap(golub_cola['ATC', 'skmeans'], k = 3)
```

merge_node

**Description**

Merge node

**Usage**

```r
## S4 method for signature 'HierarchicalPartition'
merge_node(object, node_id)
```

**Arguments**

- `node_id`: A vector of node IDs where each node is merged as a leaf node.

**Value**

A `HierarchicalPartition-class` object.

**Examples**

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

merge_node_param

**Description**

Parameters to merge branches in subgroup dendrogram.

**Usage**

```r
merge_node_param(depth = Inf, min_n_signatures = -Inf, min_p_signatures = -Inf)
```
ncol-ConsensusPartition-method

Arguments

- **depth**  
  Depth of the dendrogram.
- **min_n_signatures**  
  Minimal number of signatures for the partitioning on each node.
- **min_p_signatures**  
  Minimal fraction of signatures compared to the total number of rows on each node.

Examples

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

---

ncol-ConsensusPartition-method

*Number of columns in the matrix*

Description

Number of columns in the matrix

Usage

```r
## S4 method for signature 'ConsensusPartition'
ncol(x)
```

Arguments

- **x**  
  A ConsensusPartition-class object.

Examples

```r
# There is no example
NULL
```
Description

Number of columns in the matrix

Usage

## S4 method for signature 'ConsensusPartitionList'
ncol(x)

Arguments

x
A ConsensusPartitionList-class object.

Examples

# There is no example
NULL

Dispatch

ncol can be dispatched on following classes:

- ncol,ConsensusPartitionList-method, ConsensusPartitionList-class class method
- ncol,ConsensusPartition-method, ConsensusPartition-class class method
- ncol,DownSamplingConsensusPartition-method, DownSamplingConsensusPartition-class class method
- ncol,HierarchicalPartition-method, HierarchicalPartition-class class method

Examples

# no example
NULL
Description

Number of columns in the matrix

Usage

## S4 method for signature 'DownSamplingConsensusPartition'
ncol(x)

Arguments

  x

A \code{DownSamplingConsensusPartition-class} object.

Examples

  # There is no example
  NULL

---

Description

Number of columns in the matrix

Usage

## S4 method for signature 'HierarchicalPartition'
ncol(x)

Arguments

  x

A \code{HierarchicalPartition-class} object.

Examples

  # There is no example
  NULL
Description
Information on the nodes

Usage
## S4 method for signature 'HierarchicalPartition'
node_info(object)

Arguments

object A HierarchicalPartition-class object.

Details
It returns the following node-level information:

id  Node id.
n_columns  Number of columns.
n_signatures  Number of signatures.
p_signatures  Percent of signatures.
is_leaf  Whether the node is a leaf

Examples

# There is no example
NULL

Description
Information on the nodes

Usage
## S4 method for signature 'HierarchicalPartition'
node_level(object)
nrow-ConsensusPartition-method

Arguments

object A HierarchicalPartition-class object.

Details

It is the same as node_info, HierarchicalPartition-method.

Examples

# There is no example
NULL

Description

Number of rows in the matrix

Usage

## S4 method for signature 'ConsensusPartition'
nrow(x)

Arguments

x A ConsensusPartition-class object.

Examples

# There is no example
NULL
nrow-ConsensusPartitionList-method

Number of rows in the matrix

Description

Number of rows in the matrix

Usage

## S4 method for signature 'ConsensusPartitionList'
nrow(x)

Arguments

x

A ConsensusPartitionList-class object.

Examples

# There is no example
NULL

nrow-dispatch

Method dispatch page for nrow

Description

Method dispatch page for nrow.

Dispatch

nrow can be dispatched on following classes:

- nrow,HierarchicalPartition-method, HierarchicalPartition-class class method
- nrow,ConsensusPartitionList-method, ConsensusPartitionList-class class method
- nrow,ConsensusPartition-method, ConsensusPartition-class class method

Examples

# no example
NULL
nrow-HierarchicalPartition-method

Number of rows in the matrix

Description

Number of rows in the matrix

Usage

```r
## S4 method for signature 'HierarchicalPartition'
nrow(x)
```

Arguments

- `x`: A `HierarchicalPartition-class` object.

Examples

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

PAC

The proportion of ambiguous clustering (PAC score)

Description

The proportion of ambiguous clustering (PAC score)

Usage

```r
PAC(consensus_mat, x1 = 0.1, x2 = 0.9, class = NULL)
```

Arguments

- `consensus_mat`: A consensus matrix.
- `x1`: Lower bound to define "ambiguous clustering".
- `x2`: Upper bound to define "ambiguous clustering".
- `class`: Subgroup labels. If it is provided, samples with silhouette score less than the 5-th percential are removed from PAC calculation.

Details

The PAC score is defined as $F(x2) - F(x1)$ where $F(x)$ is the CDF of the consensus matrix.
plot_ecdf-ConsensusPartition-method

Value

A single numeric value.

See

See https://www.nature.com/articles/srep06207 for explanation of PAC score.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

data(golub_cola)
PAC(get_consensus(golub_cola[1, 1], k = 2))
PAC(get_consensus(golub_cola[1, 1], k = 3))
PAC(get_consensus(golub_cola[1, 1], k = 4))
PAC(get_consensus(golub_cola[1, 1], k = 5))
PAC(get_consensus(golub_cola[1, 1], k = 6))

# with specifying `class`
PAC(get_consensus(golub_cola[1, 1], k = 2),
   class = get_classes(golub_cola[1, 1], k = 2)[, 1])

plot_ecdf-ConsensusPartition-method

Plot the empirical cumulative distribution (eCDF) curve of the consensus matrix

Description

Plot the empirical cumulative distribution (eCDF) curve of the consensus matrix

Usage

## S4 method for signature 'ConsensusPartition'
plot_ecdf(object, ...)

Arguments

object A ConsensusPartition-class object.
...
Other arguments.

Details

It plots eCDF curve for each k.
This function is mainly used in collect_plots and select_partition_number functions.
Value

No value is returned.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

See Also

See `ecdf` for a detailed explanation of the empirical cumulative distribution function.

Examples

```r
data(golub_cola)
plot_ecdf(golub_cola[, "ATC", "skmeans"])
```

Description

Predict classes for new samples based on cola classification.

Usage

```r
## S4 method for signature 'ConsensusPartition'
predict_classes(object, k, mat,
    silhouette_cutoff = 0.5,
    fdr_cutoff = cola_opt$fdr_cutoff,
    group_diff = cola_opt$group_diff,
    scale_rows = object@scale_rows,
    diff_method = "Ftest",
    method = "centroid",
    dist_method = c("euclidean", "correlation", "cosine"), nperm = 1000,
    p_cutoff = 0.05, plot = TRUE, col_fun = NULL,
    split_by_sigatures = FALSE, force = FALSE,
    verbose = TRUE, help = TRUE, prefix = "",
    mc.cores = 1, cores = mc.cores)
```

Arguments

- `k`: Number of subgroups to get the classifications.
- `mat`: The new matrix where the sample classes are going to be predicted. The number of rows should be the same as the original matrix for cola analysis (also make sure the row orders are the same). Be careful that the scaling of `mat` should be the same as that applied in cola analysis.
predict_classes-ConsensusPartition-method

- **silhouette_cutoff**: Send to `get_signatures,ConsensusPartition-method` for determining signatures.
- **fdr_cutoff**: Send to `get_signatures,ConsensusPartition-method` for determining signatures.
- **group_diff**: Send to `get_signatures,ConsensusPartition-method` for determining signatures.
- **scale_rows**: Send to `get_signatures,ConsensusPartition-method` for determining signatures.
- **diff_method**: Send to `get_signatures,ConsensusPartition-method` for determining signatures.
- **method**: Method for predicting class labels. Possible values are "centroid", "svm" and "randomForest".
- **dist_method**: Distance method. Value should be "euclidean", "correlation" or "cosine". Send to `predict_classes,matrix-method`.
- **nperm**: Number of permutations. It is used when `dist_method` is set to "euclidean" or "cosine". Send to `predict_classes,matrix-method`.
- **p_cutoff**: Cutoff for the p-values for determining class assignment. Send to `predict_classes,matrix-method`.
- **plot**: Whether to draw the plot that visualizes the process of prediction. Send to `predict_classes,matrix-method`.
- **col_fun**: A color mapping function generated from `colorRamp2`. It is set to both heatmaps.
- **split_by_signatures**: Should the heatmaps be split based on k-means on the main heatmap, or on the patterns of the signature heatmap.
- **force**: If the value is `TRUE` and when `get_signatures,ConsensusPartition-method` internally failed, top 1000 rows with the highest between-group mean difference are used for constructing the signature centroid matrix. It is basically used internally.
- **verbose**: Whether to print messages. Send to `predict_classes,matrix-method`.
- **help**: Whether to print help messages.
- **prefix**: Used internally.
- **mc.cores**: Number of cores. This argument will be removed in future versions.
- **cores**: Number of cores, or a cluster object returned by `makeCluster`.

**Details**

The prediction is based on the signature centroid matrix from cola classification. The processes are as follows:

1. For the provided `ConsensusPartition-class` object and a selected k, the signatures that discriminate classes are extracted by `get_signatures,ConsensusPartition-method`. If number of signatures is more than 2000, only 2000 signatures are randomly sampled.
2. The signature centroid matrix is a k-column matrix where each column is the centroid of samples in the corresponding class, i.e. the mean across samples. If rows were scaled in cola analysis, the signature centroid matrix is the mean of scaled values and vise versa. Please note the samples with silhouette score less
than silhouette_cutoff are removed for calculating the centroids. 3. With the signature centroid matrix and the new matrix, it calls `predict_classes,matrix-method` to perform the prediction. Please see more details of the prediction on that help page. Please note, the scales of the new matrix should be the same as the matrix used for cola analysis.

**Value**

A data frame with two columns: the class labels (in numeric) and the corresponding p-values.

**See Also**

`predict_classes,matrix-method` that predicts the classes for new samples.

**Examples**

```r
data(golub_cola)
res = golub_cola["ATC:skmeans"]
mat = get_matrix(res)
# note scaling should be applied here because the matrix was scaled in the cola analysis
mat2 = t(scale(t(mat)))
c1 = predict_classes(res, k = 3, mat2)
# compare the real classification and the predicted classification
data.frame(cola_class = get_classes(res, k = 3)[, "class"],
           predicted = c1[, "class"])
# change to correlation method
c1 = predict_classes(res, k = 3, mat2, dist_method = "correlation")
# compare the real classification and the predicted classification
data.frame(cola_class = get_classes(res, k = 3)[, "class"],
           predicted = c1[, "class"])
```

**Description**

Method dispatch page for `predict_classes`.

**Dispatch**

`predict_classes` can be dispatched on following classes:

- `predict_classes,matrix-method, matrix-class` class method
- `predict_classes,ConsensusPartition-method, ConsensusPartition-class` class method
predict_classes-matrix-method

Examples

```r
# no example
NULL
```

predict_classes-matrix-method

*Predict classes for new samples based on signature centroid matrix*

Description

Predict classes for new samples based on signature centroid matrix

Usage

```r
## S4 method for signature 'matrix'
predict_classes(object, mat, dist_method = c("euclidean", "correlation", "cosine"),
               nperm = 1000, p_cutoff = 0.05, plot = TRUE, col_fun = NULL, split_by_signatures = FALSE,
               verbose = TRUE, prefix = ",", mc.cores = 1, cores = mc.cores, width1 = NULL, width2 = NULL)
```

Arguments

- **object**: The signature centroid matrix. See the Details section.
- **mat**: The new matrix where the classes are going to be predicted. The number of rows should be the same as the signature centroid matrix (also make sure the row orders are the same). Be careful that `mat` should be in the same scale as the centroid matrix.
- **dist_method**: Distance method. Value should be "euclidean", "correlation" or "cosine".
- **nperm**: Number of permutations. It is used when `dist_method` is set to "euclidean" or "cosine".
- **p_cutoff**: Cutoff for the p-values for determining class assignment.
- **plot**: Whether to draw the plot that visualizes the process of prediction.
- **col_fun**: A color mapping function generated from `colorRamp2`. It is set to both heatmaps.
- **verbose**: Whether to print messages.
- **split_by_signatures**: Should the heatmaps be split based on k-means on the main heatmap, or on the patterns of the signature heatmap.
- **prefix**: Used internally.
- **mc.cores**: Number of cores. This argument will be removed in future versions.
- **cores**: Number of cores, or a `cluster` object returned by `makeCluster`.
- **width1**: Width of the first heatmap.
- **width2**: Width of the second heatmap.
predict_classes-matrix-method

Details

The signature centroid matrix is a k-column matrix where each column is the centroid of samples in the corresponding class (k-group classification).

For each sample in the new matrix, the task is basically to test which signature centroid the current sample is the closest to. There are two methods: the Euclidean distance and the correlation (Spearman) distance.

For the Euclidean/cosine distance method, for the vector denoted as x which corresponds to sample i in the new matrix, to test which class should be assigned to sample i, the distance between sample i and all k signature centroids are calculated and denoted as d_1, d_2, ..., d_k. The class with the smallest distance is assigned to sample i. The distances for k centroids are sorted increasingly, and we design a statistic named "difference ratio", denoted as r and calculated as: (d_(1) - d_(2))/mean(d), which is the difference between the smallest distance and the second smallest distance, normalized by the mean distance. To test the statistical significance of r, we randomly permute rows of the signature centroid matrix and calculate r_rand. The random permutation is performed n_perm times and the p-value is calculated as the proportion of r_rand being larger than r.

For the correlation method, the distance is calculated as the Spearman correlation between sample i and signature centroid k. The label for the class with the maximal correlation value is assigned to sample i. The p-value is simply calculated by cor.test between sample i and centroid k.

If a sample is tested with a p-value higher than p_cutoff, the corresponding class label is set to NA.

Value

A data frame with two columns: the class labels (the column names of the signature centroid matrix are treated as class labels) and the corresponding p-values.

Examples

data(golub_cola)
res = golub_cola["ATC:skmeans"]
mat = get_matrix(res)
# note scaling should be applied here because the matrix was scaled in the cola analysis
mat2 = t(scale(t(mat)))

# calculate signature centroids
tb = get_signatures(res, k = 3, plot = FALSE)
sig_mat = tb[, grepl("scaled_mean", colnames(tb))]
sig_mat = as.matrix(sig_mat)
colnames(sig_mat) = paste0("class", seq_len(ncol(sig_mat)))
# this is how the signature centroid matrix looks like:
head(sig_mat)

mat2 = mat2[cbind(which_row, , drop = FALSE)]

# now we predict the class for mat2 based on sig_mat
predict_classes(sig_mat, mat2)
print.hc_table_suggest_best_k

Print the \textit{hc\_table\_suggest\_best\_k} object

\section*{Description}  
Print the \textit{hc\_table\_suggest\_best\_k} object

\section*{Usage}  
\begin{verbatim}
## S3 method for class 'hc_table_suggest_best_k'
print(x, ...)
\end{verbatim}

\section*{Arguments}  
\begin{itemize}
\item \textbf{x} \hspace{1cm} A \textit{hc\_table\_suggest\_best\_k} object from \texttt{suggest\_best\_k,HierarchicalPartition-method}.
\item \textbf{...} \hspace{1cm} Other arguments.
\end{itemize}

\section*{Examples}  
\begin{verbatim}
# There is no example
NULL
\end{verbatim}

\section*{recalc_stats}  
\textit{Recalculate statistics in the ConsensusPartitionList object}

\section*{Description}  
Recalculate statistics in the \texttt{ConsensusPartitionList} object

\section*{Usage}  
\begin{verbatim}
recalc_stats(rl)
\end{verbatim}

\section*{Arguments}  
\begin{itemize}
\item \textbf{rl} \hspace{1cm} A \texttt{ConsensusPartitionList-class} object.
\end{itemize}

\section*{Details}  
It updates the \texttt{stat} slot in the \texttt{ConsensusPartitionList} object, used internally.

\section*{Examples}  
\begin{verbatim}
# There is no example
NULL
\end{verbatim}
### register_NMF

*Register NMF partitioning method*

#### Description

Register NMF partitioning method

#### Usage

```r
register_NMF()
```

#### Details

NMF analysis is performed by `nmf`.

#### Examples

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

### register_partition_methods

*Register user-defined partitioning methods*

#### Description

Register user-defined partitioning methods

#### Usage

```r
register_partition_methods(..., scale_method = c("z-score", "min-max", "none"))
```

#### Arguments

- `...` A named list of functions.
- `scale_method` Normally, data matrix is scaled by rows before sent to the partition function. The default scaling is applied by `scale`. However, some partition functions may not accept negative values which are produced by `scale`. Here `scale_method` can be set to `min-max` which scales rows by `(x - min)/(max - min)`. Note here `scale_method` only means the method to scale rows. When `scale_rows` is set to `FALSE` in `consensus_partition` or `run_all_consensus_partition_methods`, there will be no row scaling when doing partitioning. The value for `scale_method` can be a vector if user specifies more than one partition function.
Details

The user-defined function should accept at least two arguments. The first two arguments are the data matrix and the number of subgroups. The third optional argument should always be ... so that parameters for the partition function can be passed by partition_param from consensus_partition. If users forget to add ..., it is added internally.

The function should return a vector of partitions (or class labels) or an object which can be recognized by cl_membership.

The partition function should be applied on columns (Users should be careful with this because some R functions apply on rows and some R functions apply on columns). E.g. following is how we register kmeans partition method:

```r
register_partition_methods(
  kmeans = function(mat, k, ...) {
    # mat is transposed because kmeans() applies on rows
    kmeans(t(mat), centers = k, ...)$centers
  }
)
```

The registered partitioning methods will be used as defaults in run_all_consensus_partition_methods.

To remove a partitioning method, use remove_partition_methods.

There are following default partitioning methods:

- "hclust" hierarchclal clustering with Euclidean distance, later columns are partitioned by cutree.
  If users want to use another distance metric or clustering method, consider to register a new partitioning method. E.g. register_partition_methods(hclust_cor = function(mat, k) cutree(hclust(as.dist(cor(mat)))).
- "kmeans" by kmeans.
- "skmeans" by skmeans.
- "pam" by pam.
- "mclust" by Mclust. mclust is applied to the first three principle components from rows.

Users can register two other pre-defined partitioning methods by register_NMF and register_SOM.

Value

No value is returned.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

See Also

all_partition_methods lists all registered partitioning methods.
Examples

```r
all_partition_methods()
register_partition_methods(
  random = function(mat, k) sample(k, ncol(mat), replace = TRUE)
)
all_partition_methods()
remove_partition_methods("random")
```

Description

Register SOM partitioning method

Usage

```r
register_SOM()
```

Details

The SOM analysis is performed by `som`.

Examples

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

Description

Register user-defined top-value methods

Usage

```r
register_top_value_methods(..., validate = TRUE)
```

Arguments

```r
...
validate
```

A named list of functions.

Whether validate the functions.
**register_top_value_methods**

Details

The user-defined function should accept one argument which is the data matrix where the scores are calculated by rows. Rows with top scores are treated as "top rows" in cola analysis. Following is how we register "SD" (standard deviation) top-value method:

\[
\text{register\_top\_value\_methods(} SD = \text{function(mat) apply(mat, 1, sd))}
\]

Of course, you can use `rowSds` to give a faster calculation of row SD:

\[
\text{register\_top\_value\_methods(} SD = \text{rowSds})
\]

The registered top-value method will be used as defaults in `run_all_consensus_partition_methods`. To remove a top-value method, use `remove_top_value_methods`.

There are four default top-value methods:

- "SD" standard deviation, by `rowSds`.
- "CV" coefficient variance, calculated as \( \frac{sd}{(mean+s)} \) where \( s \) is the 10\(^{th} \) percentile of all row means.
- "MAD" median absolute deviation, by `rowMads`.
- "ATC" the ATC method.

Value

No value is returned.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

See Also

`all_top_value_methods` lists all registered top-value methods.

Examples

```r
all_top_value_methods()
register_top_value_methods(
  ATC_spearman = function(mat) ATC(mat, method = "spearman")
)
all_top_value_methods()
remove_top_value_methods("ATC_spearman")
```
relabel_class

Relabel class labels according to the reference labels

Description

Relabel class labels according to the reference labels

Usage

relabel_class(class, ref, full_set = union(class, ref), return_map = TRUE)

Arguments

class A vector of class labels.
ref A vector of reference labels.
full_set The full set of labels.
return_map Whether to return the mapping of the adjusted labels.

Details

In partitions, the exact value of the class label is not of importance. E.g. for two partitions a, a, a, b, b, b and b, b, a, a, a, a, they are the same partitions although the labels of a and b are switched in the two partitions. Even the partition c, c, d, d, d is the same as the previous two although it uses a different set of labels. Here relabel_class function relabels class vector according to the labels in ref vector by looking for a mapping m() to maximize sum(m(class) == ref).

Mathematically, this is called linear sum assignment problem and it is solved by solve_LSAP.

Value

A named vector where names correspond to the labels in class and values correspond to ref, which means map = relabel_class(class, ref); map[class] returns the relabelled labels.

The returned object attaches a data frame with three columns:

- original labels. in class
- adjusted labels. according to ref
- reference labels. in ref

If return_map in the relabel_class is set to FALSE, the function simply returns a vector of adjusted class labels.

If the function returns the mapping vector (when return_map = TRUE), the mapping variable is always character, which means, if your class and ref are numeric, you need to convert them back to numeric explicitly. If return_map = FALSE, the returned relabelled vector has the same mode as class.
Examples

class = c(rep("a", 10), rep("b", 3))
ref = c(rep("b", 4), rep("a", 9))
relabel_class(class, ref)
relabel_class(class, ref, return_map = FALSE)
# if class and ref are from completely different sets
class = c(rep("A", 10), rep("B", 3))
relabel_class(class, ref)

# class labels are numeric
class = c(rep(1, 10), rep(2, 3))
ref = c(rep(2, 4), rep(1, 9))
relabel_class(class, ref)
relabel_class(class, ref, return_map = FALSE)

remove_partition_methods

| Remove partitioning methods |

Description

Remove partitioning methods

Usage

remove_partition_methods(method)

Arguments

method Name of the partitioning methods to be removed.

Value

No value is returned.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

# There is no example
NULL
remove_top_value_methods

Description
Remove top-value methods

Usage
remove_top_value_methods(method)

Arguments
method Name of the top-value methods to be removed.

Value
No value is returned.

Author(s)
Zuguang Gu <z.gu@dkfz.de>

Examples
# There is no example
NULL

rownames-ConsensusPartition-method

Description
Row names of the matrix

Usage
## S4 method for signature 'ConsensusPartition'
rownames(x)

Arguments
x A ConsensusPartition-class object.
Examples

# There is no example
NULL

rownames-ConsensusPartitionList-method
Row names of the matrix

Description
Row names of the matrix

Usage

## S4 method for signature 'ConsensusPartitionList'
rownames(x)

Arguments

x A ConsensusPartitionList-class object.

Examples

# There is no example
NULL

rownames-dispatch
Method dispatch page for rownames

Description
Method dispatch page for rownames.

Dispatch

rownames can be dispatched on following classes:

- rownames,HierarchicalPartition-method,HierarchicalPartition-class class method
- rownames,ConsensusPartition-method,ConsensusPartition-class class method
- rownames,ConsensusPartitionList-method,ConsensusPartitionList-class class method

Examples

# no example
NULL
### rownames-HierarchicalPartition-method

**Row names of the matrix**

#### Description

Row names of the matrix

#### Usage

```r
## S4 method for signature 'HierarchicalPartition'
rownames(x)
```

#### Arguments

- `x`: A `HierarchicalPartition-class` object.

#### Examples

```r
# There is no example
```

---

### run_all_consensus_partition_methods

**Consensus partitioning for all combinations of methods**

#### Description

Consensus partitioning for all combinations of methods

#### Usage

```r
run_all_consensus_partition_methods(data, 
  top_value_method = all_top_value_methods(), 
  partition_method = all_partition_methods(), 
  max_k = 6, k = NULL, 
  top_n = NULL, 
  mc.cores = 1, cores = mc.cores, anno = NULL, anno_col = NULL, 
  sample_by = "row", p_sampling = 0.8, partition_repeat = 50, 
  scale_rows = NULL, verbose = TRUE, help = cola_opt$help)
```
Arguments

- **data**
  A numeric matrix where subgroups are found by columns.

- **top_value_method**
  Method which are used to extract top n rows. Allowed methods are in `all_top_value_methods` and can be self-added by `register_top_value_methods`.

- **partition_method**
  Method which are used to partition samples. Allowed methods are in `all_partition_methods` and can be self-added by `register_partition_methods`.

- **max_k**
  Maximal number of subgroups to try. The function will try 2:max_k subgroups.

- **k**
  Alternatively, you can specify a vector k.

- **top_n**
  Number of rows with top values. The value can be a vector with length > 1. When n > 5000, the function only randomly sample 5000 rows from top n rows. If top_n is a vector, partition will be applied to every values in top_n and consensus partition is summarized from all partitions.

- **mc.cores**
  Number of cores to use. This argument will be removed in future versions.

- **cores**
  Number of cores, or a cluster object returned by `makeCluster`.

- **anno**
  A data frame with known annotation of columns.

- **anno_col**
  A list of colors (color is defined as a named vector) for the annotations. If anno is a data frame, anno_col should be a named list where names correspond to the column names in anno.

- **sample_by**
  Should randomly sample the matrix by rows or by columns?

- **p_sampling**
  Proportion of the top n rows to sample.

- **partition_repeat**
  Number of repeats for the random sampling.

- **scale_rows**
  Whether to scale rows. If it is TRUE, scaling method defined in `register_partition_methods` is used.

- **verbose**
  Whether to print messages.

- **help**
  Whether to print help messages.

Details

The function performs consensus partitioning by `consensus_partition` for all combinations of top-value methods and partitioning methods.

It also adjusts the subgroup labels for all methods and for all k to make them as consistent as possible.

Value

A ConsensusPartitionList-class object. Simply type object in the interactive R session to see which functions can be applied on it.

Author(s)

Zuguang Gu <z.gu@dkfz.de>
Examples

```r
## Not run:
set.seed(123)

m = cbind(rbind(matrix(rnorm(20*20, mean = 1), nr = 20),
               matrix(rnorm(20*20, mean = -1), nr = 20)),
           rbind(matrix(rnorm(20*20, mean = -1), nr = 20),
                  matrix(rnorm(20*20, mean = 1), nr = 20))

          ) + matrix(rnorm(40*40), nr = 40)

rl = run_all_consensus_partition_methods(data = m, top_n = c(20, 30, 40))

## End(Not run)
```

Description

Several plots for determining the optimized number of subgroups

Usage

```r
## S4 method for signature 'ConsensusPartition'
select_partition_number(object, mark_best = TRUE, all_stats = FALSE)
```

Arguments

- `mark_best`: Whether mark the best k in the plot.
- `all_stats`: Whether to show all statistics that were calculated. Used internally.

Details

There are following plots made:

- eCDF of the consensus matrix under each k, made by `plot_ecdf`, `ConsensusPartition-method`,
- PAC score,
- mean silhouette score,
- the `concordance` for each partition to the consensus partition,
- area increase of the area under the ECDF of consensus matrix with increasing k,
- Rand index for current k compared to k - 1,
- Jaccard coefficient for current k compared to k - 1,

Value

No value is returned.
Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

data(golub_cola)
select_partition_number(golub_cola["ATC", "skmeans"])

Description

Print the ConsensusPartition object

Usage

## S4 method for signature 'ConsensusPartition'
show(object)

Arguments

object A ConsensusPartition-class object.

Value

No value is returned.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

# There is no example
NULL
Print the ConsensusPartitionList object

Description
Print the ConsensusPartitionList object

Usage
## S4 method for signature 'ConsensusPartitionList'
show(object)

Arguments
object A ConsensusPartitionList-class object.

Value
No value is returned.

Author(s)
Zuguang Gu <z.gu@dkfz.de>

Examples
# There is no example
NULL

Method dispatch page for show

Description
Method dispatch page for show.

Dispatch
show can be dispatched on following classes:
- show, HierarchicalPartition-method, HierarchicalPartition-class class method
- show, ConsensusPartition-method, ConsensusPartition-class class method
- show, ConsensusPartitionList-method, ConsensusPartitionList-class class method
- show, DownSamplingConsensusPartition-method, DownSamplingConsensusPartition-class class method
show-DownSamplingConsensusPartition-method

Print the DownSamplingConsensusPartition object

Description

Print the DownSamplingConsensusPartition object

Usage

## S4 method for signature 'DownSamplingConsensusPartition'
show(object)

Arguments

object A DownSamplingConsensusPartition-class object.

Value

No value is returned.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

data(golub_cola_ds)
golub_cola_ds
show-HierarchicalPartition-method

Print the HierarchicalPartition object

Description

Print the HierarchicalPartition object

Usage

## S4 method for signature 'HierarchicalPartition'
show(object)

Arguments

object a HierarchicalPartition-class object

Value

No value is returned.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

data(golub_cola_rh)
golub_cola_rh

split_node-HierarchicalPartition-method

Split node

Description

Split node

Usage

## S4 method for signature 'HierarchicalPartition'
split_node(object, node_id,
subset = object@param$subset,
min_samples = object@param$min_samples, max_k = object@param$max_k, cores = object@param$cores, verbose = TRUE,
top_n = object@param$top_n, min_n_signatures = object@param$min_n_signatures,
group_diff = object@param$group_diff, fdr_cutoff = object@param$fdr_cutoff)
suggest_best_k-ConsensusPartition-method

Arguments

- **object**: A `HierarchicalPartition-class` object.
- **node_id**: A single ID of a node that is going to be split.
- **subset**: The same as in `hierarchical_partition`.
- **min_samples**: The same as in `hierarchical_partition`.
- **max_k**: The same as in `hierarchical_partition`.
- **cores**: Number of cores.
- **verbose**: Whether to print messages.
- **top_n**: The same as in `hierarchical_partition`.
- **min_n_signatures**: The same as in `hierarchical_partition`.
- **group_diff**: The same as in `hierarchical_partition`.
- **fdr_cutoff**: The same as in `hierarchical_partition`.

Details

It applies hierarchical consensus partitioning on the specified node.

Value

A `HierarchicalPartition-class` object.

Examples

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

Description

Suggest the best number of subgroups

Usage

```r
## S4 method for signature 'ConsensusPartition'
suggest_best_k(object,
    jaccard_index_cutoff = select_jaccard_cutoff(ncol(object)),
    mean_silhouette_cutoff = NULL,
    stable_PAC = 0.1, help = cola_opt$help)
```
suggest_best_k-ConsensusPartition-method

Arguments

object       A ConsensusPartition-class object.

jaccard_index_cutoff
The cutoff for Jaccard index for comparing to previous k.

mean_silhouette_cutoff
Cutoff for mean silhouette scores.

stable_PAC   Cutoff for stable PAC. This argument only take effect when mean_silhouette_cutoff
             is set to NULL.

help         Whether to print help message.

Details

The best k is selected according to following rules:

- All k with Jaccard index larger than jaccard_index_cutoff are removed because increasing
  k does not provide enough extra information. If all k are removed, it is marked as no subgroup
  is detected.
- If all k with Jaccard index larger than 0.75, k with the highest mean silhouette score is taken
  as the best k.
- For all k with mean silhouette score larger than mean_silhouette_cutoff, the maximal k is
  taken as the best k, and other k are marked as optional best k.
- If argument mean_silhouette_cutoff is set to NULL, which means we do not filter by mean
  silhouette scores while by 1-PAC scores. Similarly, k with the highest 1-PAC is taken the best
  k and other k are marked as optional best k.
- If it does not fit the second rule. The k with the maximal vote of the highest 1-PAC score,
  highest mean silhouette, and highest concordance is taken as the best k.

It should be noted that it is difficult to find the best k deterministically, we encourage users to
compare results for all k and determine a proper one which best explain their studies.

Value

The best k.

See

The selection of the best k can be visualized by select_partition_number.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

data(golub_cola)
obj = golub_cola["ATC", "skmeans"]
suggest_best_k(obj)
suggest_best_k-ConsensusPartitionList-method

Suggest the best number of subgroups

Description

Suggest the best number of subgroups

Usage

## S4 method for signature 'ConsensusPartitionList'
suggest_best_k(object, jaccard_index_cutoff = select_jaccard_cutoff(ncol(object)))

Arguments

- object: A ConsensusPartitionList-class object.
- jaccard_index_cutoff: The cutoff for Jaccard index for comparing to previous k.

Details

It basically gives the best k for each combination of top-value method and partitioning method by calling suggest_best_k,ConsensusPartition-method.

1-PAC score higher than 0.95 is treated as very stable partition (marked by **) and higher than 0.9 is treated as stable partition (marked by *).

Value

A data frame with the best k and other statistics for each combination of methods.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

data(golub_cola)
suggest_best_k(golub_cola)
suggest_best_k-dispatch

Method dispatch page for suggest_best_k

Description
Method dispatch page for suggest_best_k.

Dispatch
suggest_best_k can be dispatched on following classes:

- `suggest_best_k,HierarchicalPartition-method,HierarchicalPartition-class` class method
- `suggest_best_k,ConsensusPartitionList-method,ConsensusPartitionList-class` class method
- `suggest_best_k,ConsensusPartition-method,ConsensusPartition-class` class method

Examples

```r
# no example
NULL
```

suggest_best_k-HierarchicalPartition-method

Guess the best number of partitions

Description
Guess the best number of partitions

Usage

```r
## S4 method for signature 'HierarchicalPartition'
suggest_best_k(object)
```

Arguments

- `object` A `HierarchicalPartition-class` object.

Details
It basically gives the best k at each node.
Value
A data frame with the best k and other statistics for each node.

Author(s)
Zuguang Gu <z.gu@dkfz.de>

Examples
```r
data(golub_cola_rh)
suggest_best_k(golub_cola_rh)
```

---

test_between_factors  Test whether a list of factors are correlated

Description
Test whether a list of factors are correlated

Usage
test_between_factors(x, y = NULL, all_factors = FALSE, verbose = FALSE)

Arguments
- `x`: A data frame or a vector which contains discrete or continuous variables. If `y` is omit, pairwise testing for all columns in `x` is performed.
- `y`: A data frame or a vector which contains discrete or continuous variables.
- `all_factors`: Are all columns in `x` and `y` enforced to be factors?
- `verbose`: Whether to print messages.

Details
Pairwise test is applied to every two columns in the data frames. Methods are:

- two numeric variables: correlation test by `cor.test` is applied (Spearman method);
- two character or factor variables: `chisq.test` is applied;
- one numeric variable and one character/factor variable: oneway ANOVA test by `oneway.test` is applied.

This function can be used to test the correlation between the predicted classes and other known factors.

Value
A matrix of p-values. If there are NA values, basically it means there are no efficient data points to perform the test.
Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

def = data.frame(
    v1 = rnorm(100),
    v2 = sample(letters[1:3], 100, replace = TRUE),
    v3 = sample(LETTERS[5:6], 100, replace = TRUE)
)
test_between_factors(df)
x = runif(100)
test_between_factors(x, df)

Description

Test correspondance between predicted subgroups and known factors

Usage

## S4 method for signature 'ConsensusPartition'
test_to_known_factors(object, k, known = get_anno(object),
    silhouette_cutoff = 0.5, verbose = FALSE)

Arguments

object A ConsensusPartition-class object.
k Number of subgroups. It uses all k if it is not specified.
known A vector or a data frame with known factors. By default it is the annotation table set in consensus_partition or run_all_consensus_partition_methods.
silhouette_cutoff Cutoff for silhouette scores. Samples with value less than it are omit.
verbose Whether to print messages.

Details

The test is performed by test_between_factors between the predicted classes and user’s annotation table.
Value

A data frame with the following columns:

- number of samples used to test after filtered by silhouette_cutoff,
- p-values from the tests,
- number of subgroups.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

data(golub_cola)
res = golub_cola["ATC:skmeans"]
anno = get_anno(res)
anno
test_to_known_factors(res, k = 3)
# or explicitly specify known argument
test_to_known_factors(res, k = 3, known = anno)

Description

Test correspondance between predicted classes and known factors

Usage

```
## S4 method for signature 'ConsensusPartitionList'
test_to_known_factors(object, k, known = get_anno(object),
silhouette_cutoff = 0.5, verbose = FALSE)
```

Arguments

- `object`: A `ConsensusPartitionList-class` object.
- `k`: Number of subgroups. It uses all k if it is not set.
- `known`: A vector or a data frame with known factors. By default it is the annotation table set in `consensus_partition` or `run_all_consensus_partition_methods`.
- `silhouette_cutoff`: Cutoff for silhouette scores. Samples with value less than this are omit.
- `verbose`: Whether to print messages.
Details

The function basically sends each ConsensusPartition-class object to test_to_known_factors,ConsensusPartition-method and merges results afterwards.

Value

A data frame with the following columns:

- number of samples used to test after filtered by silhouette_cutoff,
- p-values from the tests,
- number of subgroups.

If there are NA values, basically it means there are no efficient data points to perform the test.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

See Also

test_between_factors, test_to_known_factors, ConsensusPartition-method

Examples

data(golub_cola)
test_to_known_factors(golub_cola)
Examples

```r
# no example
NULL
```

description

test correspondance between predicted subgroups and known factors

Usage

```r
## S4 method for signature 'DownSamplingConsensusPartition'

test_to_known_factors(object, k, known = get_anno(object),
                      p_cutoff = 0.05, verbose = FALSE)
```

Arguments

- `object`: A `DownSamplingConsensusPartition-class` object.
- `k`: Number of subgroups. It uses all k if it is not specified.
- `known`: A vector or a data frame with known factors. By default it is the annotation table set in `consensus_partition_by_down_sampling`.
- `p_cutoff`: Cutoff for p-values for the class prediction. Samples with p-value higher than it are omit.
- `verbose`: Whether to print messages.

Details

The test is performed by `test_between_factors` between the predicted classes and user's annotation table.

Value

A data frame with the following columns:

- number of samples used to test after filtered by `p_cutoff`,
- p-values from the tests,
- number of subgroups.

Author(s)

Zuguang Gu <z.gu@dkfz.de>
Examples

```r
data(golub_cola_ds)
test_to_known_factors(golub_cola_ds, k = 3)
test_to_known_factors(golub_cola_ds)
```

---

**Description**

Test correspondence between predicted classes and known factors

**Usage**

```r
## S4 method for signature 'HierarchicalPartition'
test_to_known_factors(object, known = get_anno(object[1]),
merge_node = merge_node_param(), verbose = FALSE)
```

**Arguments**

- `merge_node`: Parameters to merge sub-dendrograms, see `merge_node_param`.
- `known`: A vector or a data frame with known factors. By default it is the annotation table set in `hierarchical_partition`.
- `verbose`: Whether to print messages.

**Value**

A data frame with columns:

- number of samples
- p-values from the tests
- number of classes

The classifications are extracted for each depth.

**Author(s)**

Zuguang Gu <z.gu@dkfz.de>

**Examples**

```r
data(golub_cola_rh)
# golub_cola_rh already has known annotations, so test_to_known_factors()
# can be directly applied
test_to_known_factors(golub_cola_rh)
```
top_elements_overlap  
Overlap of top elements from different metrics

Description
Overlap of top elements from different metrics

Usage
```
top_elements_overlap(object, top_n = round(0.25*length(object[[1]])),
                     method = c("euler", "upset", "venn", "correspondance"),
                     fill = NULL, ...)
```

Arguments
- `object`: A list which contains values from different metrics.
- `top_n`: Number of top rows.
- `method`: `euler`: plot Euler diagram by `euler`; `upset`: draw the Upset plot by `UpSet`;
  `venn`: plot Venn diagram by `venn`; `correspondance`: use `correspond_between_rankings`.
- `fill`: Filled color for the Euler diagram. The value should be a color vector. Transparency of 0.5 are added internally.
- `...`: Additional arguments passed to `plot.euler`, `UpSet` or `correspond_between_rankings`.

Details
The i'th value in every vectors in `object` should correspond to the same element from the original data.

Value
No value is returned.

Author(s)
Zuguang Gu <z.gu@dkfz.de>

Examples
```
require(matrixStats)
set.seed(123)
mat = matrix(rnorm(1000), nrow = 100)
lt = list(sd = rowSds(mat), mad = rowMads(mat))
top_elements_overlap(lt, top_n = 20, method = "euler")
top_elements_overlap(lt, top_n = 20, method = "upset")
top_elements_overlap(lt, top_n = 20, method = "venn")
top_elements_overlap(lt, top_n = 20, method = "correspondance")
```
### Description

Heatmap of top rows

### Usage

```r
## S4 method for signature 'ConsensusPartition'

top_rows_heatmap(object, top_n = min(object@top_n), k = NULL,
                  anno = get_anno(object), anno_col = get_anno_col(object),
                  scale_rows = object@scale_rows, ...)
```

### Arguments

- `object` A `ConsensusPartition-class` object.
- `top_n` Number of top rows.
- `k` Number of subgroups. If it is not specified, it uses the "best k".
- `anno` A data frame of annotations.
- `anno_col` A list of colors (color is defined as a named vector) for the annotations. If `anno` is a data frame, `anno_col` should be a named list where names correspond to the column names in `anno`.
- `scale_rows` Whether to scale rows.
- `...` Pass to `top_rows_heatmap,matrix-method`.

### Value

No value is returned.

### Author(s)

Zuguang Gu <z.gu@dkfz.de>

### See Also

`top_rows_heatmap,matrix-method`

### Examples

```r
data(golub_cola)
top_rows_heatmap(golub_cola["ATC:skmeans"])```
### Description

Heatmap of top rows from different top-value methods

### Usage

```r
## S4 method for signature 'ConsensusPartitionList'
top_rows_heatmap(object, top_n = min(object@list[[1]]@top_n),
    anno = get_anno(object), anno_col = get_anno_col(object),
    scale_rows = object@list[[1]]@scale_rows, ...)
```

### Arguments

- **object**: A `ConsensusPartitionList-class` object.
- **top_n**: Number of top rows.
- **anno**: A data frame of annotations for the original matrix columns. By default it uses the annotations specified in `run_all_consensus_partition_methods`.
- **anno_col**: A list of colors (color is defined as a named vector) for the annotations. If `anno` is a data frame, `anno_col` should be a named list where names correspond to the column names in `anno`.
- **scale_rows**: Whether to scale rows.
- **...**: Pass to `top_rows_heatmap,matrix-method`.

### Value

No value is returned.

### Author(s)

Zuguang Gu <z.gu@dkfz.de>

### See Also

- `top_rows_heatmap,matrix-method`

### Examples

```r
data(golub_cola)
top_rows_heatmap(golub_cola)
```
Method dispatch page for top_rows_heatmap

Description

Method dispatch page for top_rows_heatmap.

Dispatch

top_rows_heatmap can be dispatched on following classes:

- top_rows_heatmap,ConsensusPartition-method,ConsensusPartition-class class method
- top_rows_heatmap,ConsensusPartitionList-method,ConsensusPartitionList-class class method
- top_rows_heatmap,HierarchicalPartition-method,HierarchicalPartition-class class method
- top_rows_heatmap,matrix-method,matrix-class class method

Examples

# no example
NULL

Heatmap of top rows from different top-value methods

Usage

## S4 method for signature 'HierarchicalPartition'
top_rows_heatmap(object, top_n = min(object@list[[1]]@top_n),
         anno = get_anno(object), anno_col = get_anno_col(object),
         scale_rows = object@list[[1]]@scale_rows, ...)

Heatmap of top rows from different top-value methods
# Arguments

- **object**: A `HierarchicalPartition-class` object.
- **top_n**: Number of top rows.
- **anno**: A data frame of annotations for the original matrix columns. By default it uses the annotations specified in `hierarchical_partition`.
- **anno_col**: A list of colors (color is defined as a named vector) for the annotations. If `anno` is a data frame, `anno_col` should be a named list where names correspond to the column names in `anno`.
- **scale_rows**: Whether to scale rows.
- **...**: Pass to `top_rows_heatmap`, `matrix-method`

# Value

No value is returned.

# Author(s)

Zuguang Gu <z.gu@dkfz.de>

# See Also

`top_rows_heatmap`, `matrix-method`

# Examples

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

---

# Description

Heatmap of top rows from different top-value methods

# Usage

```r
## S4 method for signature 'matrix'
top_rows_heatmap(object, all_top_value_list = NULL,
                 top_value_method = all_top_value_methods(),
                 bottom_annotation = NULL,
                 top_n = round(0.25*nrow(object)), scale_rows = TRUE, ...)
```
Arguments

- **object**: A numeric matrix.
- **all_top_value_list**: Top-values that have already been calculated from the matrix. If it is NULL the values are calculated by methods in **top_value_method** argument.
- **top_value_method**: Methods defined in `all_top_value_methods`.
- **bottom_annotation**: A `HeatmapAnnotation-class` object.
- **top_n**: Number of top rows to show in the heatmap.
- **scale_rows**: Whether to scale rows.
- **...**: Pass to `Heatmap`.

Details

The function makes heatmaps where the rows are scaled (or not scaled) for the top n rows from different top-value methods.

The top n rows are used for subgroup classification in cola analysis, so the heatmaps show which top-value method gives better candidate rows for the classification.

Value

No value is returned.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

```r
set.seed(123)
mat = matrix(rnorm(1000), nrow = 100)
top_rows_heatmap(mat, top_n = 25)
```

Description

Overlap of top rows from different top-value methods

Usage

```r
## S4 method for signature 'ConsensusPartitionList'
top_rows_overlap(object, top_n = min(object@list[[1]]@top_n),
    method = c("euler", "upset", "venn", "correspondance"), fill = NULL, ...)
```
Arguments

object A ConsensusPartitionList-class object.

top_n Number of top rows.

method euler: plot Euler diagram by euler; upset: draw the Upset plot by UpSet; venn: plot Venn diagram by venn; correspondance: use correspond_between_rankings.

fill Filled color for the Euler diagram. The value should be a color vector. Transparency of 0.5 are added internally.

... Additional arguments passed to plot.euler, UpSet or correspond_between_rankings.

Value

No value is returned.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

See Also

top_elements_overlap

Examples

data(golub_cola)
top_rows_overlap(golub_cola, method = "euler")
top_rows_overlap(golub_cola, method = "upset")
top_rows_overlap(golub_cola, method = "venn")
top_rows_overlap(golub_cola, method = "correspondance")

description

Method dispatch page for top_rows_overlap.

Dispatch

top_rows_overlap can be dispatched on following classes:

• top_rows_overlap,HierarchicalPartition-method,HierarchicalPartition-class class method
• top_rows_overlap,matrix-method,matrix-class class method
• top_rows_overlap,ConsensusPartitionList-method,ConsensusPartitionList-class class method
Examples

# no example
NULL

Description

Overlap of top rows on different nodes

Usage

## S4 method for signature 'HierarchicalPartition'

top_rows_overlap(object, method = c("euler", "upset", "venn"), fill = NULL, ...)

Arguments

- method: euler: plot Euler diagram by euler; upset: draw the Upset plot by UpSet; venn: plot Venn diagram by venn; correspondance: use correspond_between_rankings.
- fill: Filled color for the Euler diagram. The value should be a color vector. Transparency of 0.5 are added internally.
- ...: Additional arguments passed to plot.euler, UpSet or correspond_between_rankings.

Value

No value is returned.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

See Also

top_elements_overlap

Examples

data(golub_cola_rh)
top_rows_overlap(golub_cola_rh, method = "euler")
top_rows_overlap(golub_cola_rh, method = "upset")
top_rows_overlap(golub_cola_rh, method = "venn")
## S4 method for signature 'matrix'

`top_rows_overlap(object, top_value_method = all_top_value_methods(),
    top_n = round(0.25*nrow(object)),
    method = c("euler", "upset", "venn", "correspondance"),
    fill = NULL, ...)`

### Arguments

- **object**: A numeric matrix.
- **top_value_method**: Methods defined in `all_top_value_methods`.
- **top_n**: Number of top rows.
- **method**: `euler`: plot Euler diagram by `euler`; `upset`: draw the Upset plot by `UpSet`; `venn`: plot Venn diagram by `venn`; `correspondance`: use `correspond_between_rankings`.
- **fill**: Filled color for the Euler diagram. The value should be a color vector. Transparency of 0.5 are added internally.
- **...**: Additional arguments passed to `plot.euler` or `correspond_between_rankings`.

### Details

It first calculates scores for every top-value method and make plot by `top_elements_overlap`.

### Value

No value is returned.

### Author(s)

Zuguang Gu <z.gu@dkfz.de>

### See Also

- `top_elements_overlap`
Examples

```r
set.seed(123)
mat = matrix(rnorm(1000), nrow = 100)
top_rows_overlap(mat, top_n = 25)
```

## [.ConsensusPartitionList

Subset a ConsensusPartitionList object

### Description

Subset a ConsensusPartitionList object

### Usage

```r
## S3 method for class 'ConsensusPartitionList'
x[i, j, drop = TRUE]
```

### Arguments

- `x`: A `ConsensusPartitionList-class` object.
- `i`: Index for top-value methods, character or numeric.
- `j`: Index for partitioning methods, character or numeric.
- `drop`: Whether drop class

### Details

For a specific combination of top-value method and partitioning method, you can also subset by e.g. `x['SD:hclust']`.

### Value

A `ConsensusPartitionList-class` object or a `ConsensusPartition-class` object.

### Author(s)

Zuguang Gu <z.gu@dkfz.de>

### Examples

```r
data(golub_cola)
golub_cola[c("SD", "MAD"), c("hclust", "kmeans")]
golub_cola["SD", "kmeans"] # a ConsensusPartition object
golub_cola["SD:kmeans"] # a ConsensusPartition object
golub_cola["SD:kmeans", drop = FALSE] # still a ConsensusPartitionList object
golub_cola["SD:kmeans", drop = FALSE] # still a ConsensusPartitionList object
```
HierarchicalPartition

Subset the HierarchicalPartition object

Description

Subset the HierarchicalPartition object

Usage

## S3 method for class 'HierarchicalPartition'

x[i]

Arguments

x A HierarchicalPartition-class object.
i Index. The value should be numeric or a node ID.

Details

On each node, there is a ConsensusPartition-class object.

Note you cannot get a sub-hierarchy of the partition.

Value

A ConsensusPartition-class object.

Examples

data(golub_cola_rh)
golub_cola_rh["01"]
[[.ConsensusPartitionList

*Subset a ConsensusPartitionList object*

**Description**

Subset a ConsensusPartitionList object

**Usage**

```r
## S3 method for class 'ConsensusPartitionList'
x[[i]]
```

**Arguments**

- `x` A `ConsensusPartitionList-class` object.
- `i` Character index for combination of top-value methods and partitioning method in a form of e.g. SD:kmeans.

**Value**

A `ConsensusPartition-class` object.

**Author(s)**

Zuguang Gu <z.gu@dkfz.de>

**Examples**

```r
data(golub_cola)
golub_cola[["SD:kmeans"]]
```

[[.HierarchicalPartition

*Subset the HierarchicalPartition object*

**Description**

Subset the HierarchicalPartition object

**Usage**

```r
## S3 method for class 'HierarchicalPartition'
x[[i]]
```
Arguments

x  A **HierarchicalPartition-class** object
i  Index. The value should be numeric or a node ID.

Details

On each node, there is a **ConsensusPartition-class** object.
Note you cannot get a sub-hierarchy of the partition.

Value

A **ConsensusPartition-class** object.

Examples

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