Package 'scBubbletree'

April 24, 2025

Type Package

Title Quantitative visual exploration of scRNA-seq data

Version 1.10.0

Description scBubbletree is a quantitative method for the visual exploration of scRNA-seq data, preserving key biological properties such as local and global cell distances and cell density distributions across samples. It effectively resolves overplotting and enables the visualization of diverse cell attributes from multiomic single-cell experiments. Additionally, scBubbletree is user-friendly and integrates seamlessly with popular scRNA-seq analysis tools, facilitating comprehensive and intuitive data interpretation.

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Depends R (>= 4.2.0)

Imports reshape2, BiocParallel, ape, scales, Seurat, ggplot2, ggtree, patchwork, proxy, methods, stats, base, utils, dplyr

Suggests BiocStyle, knitr, testthat, cluster, SingleCellExperiment

Encoding UTF-8

NeedsCompilation no

biocViews Visualization, Clustering, SingleCell, Transcriptomics, RNASeq

BugReports https://github.com/snaketron/scBubbletree/issues

URL https://github.com/snaketron/scBubbletree

SystemRequirements Python (>= 3.6), leidenalg (>= 0.8.2)

RoxygenNote 6.1.1

VignetteBuilder knitr

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scBubbletree-package The R package scBubbletree

Description

Method for quantitative visualization of single cell RNA-seq data

Details

This package contains functions for clustering, hierarchical grouping of clusters and visualization of scRNA-seq data.

Author(s)

Authors and maintainers:

• Simo Kitanovski <simokitanovski@uni-due.de>(ORCID)

compare_bubbletrees

See Also

Useful links:

- https://github.com/snaketron/scBubbletree
- Report bugs at https://github.com/snaketron/scBubbletree/issues

compare_bubbletrees Comparison of two bubbletrees generated from the same scRNA-seq data

Description

compare_bubbletrees takes as its main input two bubbletrees generated from the **same input data** but potentially with different input parameters (e.g. clustering method or resolutions).

It then does the following two operations:

1. computes the Jaccard distance (JD) and the intersection between paris of clusters from the two bubbletrees. This is visualized as a heatmap.

2. it visualizes the two bubbletrees together with the heatmap.

Usage

Arguments

btd_1	bubbletree object
btd_2	bubbletree object
<pre>tile_text_size</pre>	integer, size of tile labels (default = 3)
tile_bw	logical, tile grayscale (tile_bw = TRUE) vs. color (tile_bw = FALSE, default)
ratio_heatmap	nummeric, probability (default = 0.5) that dictates the relative width and height of the heatmap and the bubbletrees

Details

compare_bubbletrees takes as its main input two bubbletrees generated from the **same input data** but potentially with different input parameters (e.g. clustering method or resolutions).

It then does the following two operations:

1. computes the Jaccard distance and the intersection between paris of clusters from the two bubbletrees. This is visualized as a heatmap.

2. it visualizes the two bubbletrees together with the heatmap.

Value

comparison	ggplot2 objects assembled by R-package patchwork
m	data.frame object with JD and intersection for each pair of clusters from the two bubbletrees

Author(s)

Simo Kitanovski <simo.kitanovski@uni-due.de>

See Also

get_k, get_bubbletree_dummy, get_bubbletree_graph, get_bubbletree_kmeans, get_gini, get_gini_k, d_500, get_num_tiles, get_num_violins, get_cat_tiles

Examples

```
# input data
data("d_500", package = "scBubbletree")
A <- d_500$A
btd_1 <- get_bubbletree_graph(x = A,</pre>
                               r = 1,
                               n_{start} = 20,
                               iter_max = 100,
                               algorithm = "original",
                               knn_k = 50,
                               hclust_method = "average",
                               hclust_distance = "euclidean",
                               cores = 1,
                               round_digits = 2,
                               show_simple_count = FALSE)
btd_2 <- get_bubbletree_kmeans(x = A,</pre>
                                k = 8,
                                cores = 1,
                                round_digits = 1,
                                show_simple_count = FALSE,
                                kmeans_algorithm = "MacQueen",
                                hclust_distance = "euclidean",
                                hclust_method = "average")
btd_comparison <- compare_bubbletrees(btd_1 = btd_1,</pre>
                                        btd_2 = btd_2,
                                        tile_bw = FALSE,
                                        tile_text_size = 3,
                                        ratio_heatmap = 0.5)
```

plot
btd_comparison\$tree_comparison

data.frame of heatmap data
btd_comparison\$m

d_500

Dataset: 500 PBMCs

Description

d_500 is a list with 3 elements:

1. A = numeric matrix $A^{5}00x15$ with n=500 rows for PBMCs and f=15 principal components.

2. f = character vector f of length 500. Each element in f represents the predicted cell type of a specific cell.

3. fs = numeric matrix containing normalized gene expressions of 12 marker genes in 500 cells.

Usage

data("d_500", package = "scBubbletree")

Format

Format of d_500: list

Details

This data is a sample drawn from a larger dataset of 2,700 PBMCs. The original dataset was processed as described in vignette (accessed 23, Sep, 2022):

https://satijalab.org/seurat/articles/multimodal_reference_mapping.html

See R script inst/script/get_d_500.R to see how this dataset was created.

Source

https://satijalab.org/seurat/articles/multimodal_reference_mapping.html

Examples

```
data("d_500", package = "scBubbletree")
```

```
A <- d_500$A
base::dim(A)
f <- d_500$f
base::table(f)
fs <- d_500$fs
base::dim(fs)</pre>
```

Dataset: scRNA-seq data of 3,918 cells from 5 adenocarcinoma cell lines

Description

d_ccl is a list with 3 elements:

1. A = numeric matrix with n=3,918 rows for cells and f=15 principal components

2. m = data.frame meta data

3. e = numeric matrix containing normalized gene expressions of 5 marker genes

Usage

data("d_ccl", package = "scBubbletree")

Format

Format of d_ccl: list

Details

d_ccl is a scRNA-seq dataset containing a mixture of 3,918 cells from five human lung adenocarcinoma cell lines (HCC827, H1975, A549, H838 and H2228). The dataset is available here:

https://github.com/LuyiTian/sc_mixology/blob/master/data/ sincell_with_class_5cl.RData

The library has been prepared with 10x Chromium platform and sequenced with Illumina NextSeq 500 platform. Raw data has been processed with Cellranger. The tool demuxlet has been used to predict the identity of each cell based on known genetic differences between the different cell lines.

See R script inst/script/get_d_ccl.R to see how this dataset was created.

Source

https://github.com/LuyiTian/sc_mixology/blob/master/data/ sincell_with_class_5cl.RData

References

Tian, Luyi, et al. "Benchmarking single cell RNA-sequencing analysis pipelines using mixture control experiments." Nature methods 16.6 (2019): 479-487

Examples

```
data("d_ccl", package = "scBubbletree")
A <- d_ccl$A
base::dim(A)
m <- d_ccl$m
utils::head(m)</pre>
```

d_ccl

```
e <- d_ccl$e
base::dim(e)</pre>
```

get_bubbletree_dummy Build bubbletree from matrix A of low-dimensional projections and vector cs of externally generated cluster IDs

Description

get_bubbletree_dummy takes two main inputs:

1. numeric matrix $A^{n \times f}$, which represents a low-dimensional projection (obtained e.g. by PCA) of the original high-dimensional scRNA-seq data, with n rows as cells and f columns as low-dimension features.

2. vector cs of cluster IDs of each cell

The function get_bubbletree_dummy performs one main operation. It organizes the bubbles (defined by cs) in a hierarchical dendrogram (bubbletree) which represents the hierarchical relationships between the clusters (bubbles).

Usage

Arguments

х	numeric matrix $(A^{n \times f}$ with <i>n</i> cells, and <i>f</i> low-dimensional projections of the original single cell RNA-seq dataset)	
cs	vector, cluster IDs	
В	integer, number of bootstrap iterations to perform in order to generate bubbletree	
N_eff	integer, number of cells to draw randomly from each cluster when computing inter-cluster distances	
hclust_distance		
	distance measure to be used: euclidean (default) or manhattan, see documenta- tion of stats::dist	
hclust_method	agglomeration method to be used, default = average. See documentation of stats::hclust	

cores	integer, number of PC cores for parallel execution	
round_digits	integer, number of decimal places to keep when showing the relative frequency of cells in each bubble	
show_simple_count		
	logical, if show_simple_count=T, cell counts in each bubble will be divided by 1,000 to improve readability. This is only useful for samples that are composed of millions of cells.	
verbose	logical, progress messages	

Details

This function is similar to get_bubbletree_kmeans and get_bubbletree_graph but skips the clustering step. See the documentation of the respective functions.

Value

A	input x matrix
k	number of clusters
km	NULL
ph	boot_ph: bootstrap dendrograms H_b ; main_ph: bubbletree H
ph_data	two phlogenies: $ph_c = phylogenity$ constructed from bubble centroids (computed from $A^{n \times f}$); $ph_p = main_ph = phylogeny$ constructed from intercell distances
pair_dist	inter-cluster distances used to generate the dendrograms
cluster	cluster assignments of each cell
input_par	list of all input parameters
tree	ggtree bubbletree object
tree_simple	simplified ggtree bubbletree object
tree_meta	meta-data associated with the bubbletree

Author(s)

Simo Kitanovski <simo.kitanovski@uni-due.de>

See Also

get_k, get_r, get_bubbletree_kmeans, get_bubbletree_graph, get_bubbletree_comparison, get_gini, get_gini_k, get_num_tiles, get_num_violins, get_cat_tiles, d_500

Examples

```
# input data
data("d_500", package = "scBubbletree")
A <- d_500$A
cs <- base::sample(x = LETTERS[1:5], size = nrow(A), replace = TRUE)</pre>
```

get_bubbletree_graph Louvain clustering and hierarchical grouping of k' clusters (bubbles)

Description

get_bubbletree_graph has two main inputs:

1. numeric matrix $A^{n \times f}$, which represents a low-dimensional projection (obtained e.g. by PCA) of the original high-dimensional scRNA-seq data, with n rows as cells and f columns as low-dimension features.

2. clustering resolution r

The function get_bubbletree_graph performs two main operations. First, it performs Louvain clustering to identify groups (bubbles) of transcriptionally similar cells; second, it organizes the bubbles in a hierarchical dendrogram (bubbletree) which adequatly represents inter-cluster relationships.

Usage

```
get_bubbletree_graph(x,
```

```
r,
B = 200,
N_eff = 100,
n_start = 20,
iter_max = 100,
algorithm = "original",
knn_k = 20,
hclust_method = "average",
hclust_distance = "euclidean",
cores = 1,
round_digits = 2,
show_simple_count = FALSE,
verbose = TRUE)
```

Arguments

Х	numeric matrix $(A^{n \times f}$ with <i>n</i> cells, and <i>f</i> low-dimensional projections of the original single cell RNA-seq dataset)
r	number, clustering resolution

В	integer, number of bootstrap iterations to perform in order to generate bubble- tree. If $B = 200$ (default), cluster centroids are used to compute inter-cluster distances and N_{eff} is ignored, i.e. all cells are used to compute centroids.	
N_eff	integer, number of cells to draw randomly from each cluster when computing inter-cluster distances.	
n_start,iter_m	ax	
	parameters for Louvain clustering, see documentation of function FindClusters, R-package Seurat	
algorithm	character, four clustering algorithms: 'original', 'LMR', 'SLM' and 'Leiden', see documentation of function FindClusters, R-package Seurat	
knn_k	integer, defines k for the k-nearest neighbor algorithm, see documentation of function FindClusters, R-package Seurat	
hclust_method	the agglomeration method to be used (default = average). See documentation of stats::hclust	
hclust_distance		
	distance measure to be used: euclidean (default) or manhattan, see documenta- tion of stats::dist	
cores	integer, number of PC cores for parallel execution	
round_digits	integer, number of decimal places to keep when showing the relative frequency of cells in each bubble	
show_simple_count		
	logical, if show_simple_count=T, cell counts in each bubble will be divided by 1,000 to improve readability. This is only useful for samples that are composed of millions of cells.	
verbose	logical, progress messages	

Details

For Louvain clustering get_bubbletree_graph uses the function FindClusters implemented in R-package Seurat. For additional information on the clustering procedure see the documentation of FindClusters. To organize the resulting clusters in a hierarchical dendrogram, then the following steps are performed:

- 1. In bootrap iteration b from 1: B
- 2. draw up to N_{eff} number of cells at random from each cluster without replacement
- 3. compute distances (in space $A^{n \times f}$) between all pairs of cells in cluster *i* and cluster *j*
- 4. compute mean distance between cluster i and j and populate inter-cluster distance matrix $D_{h}^{k \times k}$

5. perform hierarchical clustering with user-specified agglomeration method based on $D_b^{k \times k}$ to generate dendrogram H_b

6. quantify branch robustness in H by counting how many times each branch is found among bootrap dendrograms H_b

Value

А

input x matrix

k	number of clusters
r	clustering resolution
ph	boot_ph: bootstrap dendrograms H_b ; main_ph: bubbletree H
ph_data	two phlogenies: $ph_c = phylogenity$ constructed from bubble centroids (computed from $A^{n \times f}$); $ph_p = main_ph = phylogeny$ constructed from intercell distances
pair_dist	inter-cluster distances used to generate the dendrograms
cluster	cluster assignments of each cell
input_par	list of all input parameters
tree	ggtree bubbletree object
tree_simple	simplified ggtree bubbletree object
tree_meta	meta-data associated with the bubbletree

Author(s)

Simo Kitanovski <simo.kitanovski@uni-due.de>

See Also

get_k, get_bubbletree_dummy, get_bubbletree_kmeans, get_bubbletree_comparison, get_gini, get_gini_k, d_500, get_num_tiles, get_num_violins, get_cat_tiles

Examples

```
# input data
data("d_500", package = "scBubbletree")
A <- d_500$A
b <- get_bubbletree_graph(x = A,</pre>
                           r = 1,
                           B = 200,
                           N_{eff} = 100,
                           n_{start} = 20,
                           iter_max = 100,
                           algorithm = "original",
                           knn_k = 20,
                           hclust_method = "average",
                           hclust_distance = "euclidean",
                           cores = 1,
                           round_digits = 2,
                           show_simple_count = FALSE)
```

b\$tree

get_bubbletree_kmeans k-means clustering and hierarchical grouping of k clusters (bubbles)

Description

get_bubble_kmeans takes two main inputs:

1. numeric matrix $A^{n \times f}$, which represents a low-dimensional projection (obtained e.g. by PCA) of the original high-dimensional scRNA-seq data, with n rows as cells and f columns as low-dimension features.

2. number k of clusters

The function get_bubble_kmeans performs two main operations. First, it performs k-means clustering to identify groups (bubbles) of transcriptionally similar cells. Second, it organizes the bubbles in a hierarchical dendrogram (bubbletree) which adequatly represents inter-cluster relationships.

Usage

```
get_bubbletree_kmeans(x,
```

```
k,
B = 200,
N_eff = 100,
n_start = 1000,
iter_max = 300,
kmeans_algorithm = "MacQueen",
hclust_distance = "euclidean",
hclust_method = "average",
cores = 1,
round_digits = 2,
show_simple_count = FALSE,
verbose = TRUE)
```

Arguments

x	numeric matrix $(A^{n \times f}$ with <i>n</i> cells, and <i>f</i> low-dimensional projections of the original single cell RNA-seq dataset)	
k	integer, number of clusters	
В	integer, number of bootstrap iterations to perform in order to generate bubbletree	
N_eff	integer, number of cells to draw randomly from each cluster when computing inter-cluster distances	
n_start,iter_max,kmeans_algorithm		
	parameters for k-means clustering, see documentation of function k-means, R-package stats	
hclust_distance		
	distance measure to be used: euclidean (default) or manhattan, see documenta- tion of stats::dist	

hclust_method	the agglomeration method to be used, default = average. See documentation of stats::hclust	
cores	integer, number of PC cores for parallel execution	
round_digits	integer, number of decimal places to keep when showing the relative frequency of cells in each bubble	
show_simple_count		
	logical, if show_simple_count=T, cell counts in each bubble will be divided by 1,000 to improve readability. This is only useful for samples that are composed of millions of cells.	
verbose	logical, progress messages	

Details

For k-means clustering get_bubble_kmeans uses the function kmeans implemented in R-package stats (version 4.2.0). For additional information on the clustering procedure see the documentation of kmeans. To organize the resulting clusters in a hierarchical dendrogram these steps are performed:

- 1. In bootrap iteration b from 1: B
- 2. draw up to N_{eff} number of cells at random from each cluster without replacement
- 3. compute distances (in space $A^{n \times f}$) between pairs of cells in cluster *i* and cluster *j*
- 4. compute mean distance between cluster i and j and populate inter-cluster distance matrix $D_b^{k \times k}$

5. perform hierarchical clustering with user-specified agglomeration method based on $D_b^{k \times k}$ to generate dendrogram H_b

6. quantify branch robustness in H by counting how many times each branch is found among the bootrap dendrograms H_b

Value

А	input matrix x
k	number of clusters
km	k-means clustering results identical to those generated by function k-means from R-package stats
ph	boot_ph: bootstrap dendrograms H_b ; main_ph: bubbletree H
ph_data	two phlogenies: $ph_c = phylogenity$ constructed from bubble centroids (computed from $A^{n \times f}$); $ph_p = main_ph = phylogeny$ constructed from intercell distances
pair_dist	inter-cluster distances used to generate the dendrograms
cluster	cluster assignments of each cell
input_par	list of all input parameters
tree	ggtree bubbletree object
<pre>tree_simple</pre>	simplified ggtree bubbletree object
tree_meta	meta-data associated with the bubbletree

Author(s)

Simo Kitanovski <simo.kitanovski@uni-due.de>

See Also

get_k, get_bubbletree_dummy, get_bubbletree_graph, get_gini, get_gini_k, d_500, get_num_tiles, get_num_violins, get_cat_tiles, get_bubbletree_comparison

Examples

b\$tree

get_cat_tiles Visualization of categorical cell features using tile plots

Description

get_cat_tiles creates tile plot to visualize the relative frequency of categorical cell features between and within the bubbles of a bubbletree

Usage

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get_cat_tiles

Arguments

btd	bubbletree object	
f	character vector, categorical cell features	
integrate_verti	cal	
	logical, if integrate_vertical=TRUE: relative frequency of the features is shown in each bubble, if integrate_vertical=FALSE: relative frequencies of the features is shown within each bubble	
round_digits	integer, number of decimal places to keep when showing the relative frequency of cells in each bubble	
<pre>tile_text_size</pre>	integer, size of tile labels	
x_axis_name	character, x-axis title	
rotate_x_axis_labels		
	logical, should the x-axis labels be shown horizontally (rotate_x_axis_labels = FALSE) or vertically (rotate_x_axis_labels = TRUE)	
tile_bw	logical, tile grayscale (tile_bw = TRUE) vs. color (tile_bw = FALSE, default)	

Details

get_cat_tiles uses two main inputs:

1. bubbletree object

2. character vector of categorical cell features.

The order of the cells used to generat the bubbletree (input 1.) should correspond to the order of cells in the vector of categorical cell features (input 2.)

This function computes:

1. with integrate_vertical=T: relative frequencies of each feature across the different bubbles

2. with integrate_vertical\=F: within-bubble relative frequencies (composition) of different features

Value

plot	ggplot2, tile plot
table	data.frame, raw data used to generate the plot

Author(s)

Simo Kitanovski <simo.kitanovski@uni-due.de>

See Also

get_k, get_r get_bubbletree_dummy, get_bubbletree_kmeans, get_bubbletree_graph, get_gini, get_gini_k, get_num_tile, get_num_violins, d_500

Examples

```
# input data
data("d_500", package = "scBubbletree")
A <- d_500$A
f <- d_500$f
b <- get_bubbletree_graph(x = A,</pre>
                            r = 0.8,
                           N_{eff} = 100)
g_v <- get_cat_tiles(btd = b,</pre>
                      f = f,
                      integrate_vertical = TRUE,
                      round_digits = 2,
                      tile_text_size = 3,
                      x_axis_name = "Feature",
                      rotate_x_axis_labels = TRUE)
g_h <- get_cat_tiles(btd = b,</pre>
                     f = f,
                     integrate_vertical = FALSE,
                     round_digits = 2,
                     tile_text_size = 3,
                     x_axis_name = "Feature",
                     rotate_x_axis_labels = TRUE)
```

b\$tree|g_v\$plot|g_h\$plot

get	ŀα	in	١i
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Gini impurity index computed for a clustering solution and a vector of categorical cell feature labels

Description

How well is a set of categorical feature labels (e.g. cell type predictions) partitioned accross the different clusters of a clustering solution? We can assess this using the Gini impurity index (see details below).

Inputs are two equal-sized vectors:

1) clusters IDs

2) labels

Output:

1) cluster-specific purity -> Gini impurity (GI) index

2) clustering solution impurity -> Weighted Gini impurity (WGI) index

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get_gini

Usage

get_gini(labels, clusters)

Arguments

labels	character or numeric vector of labels
clusters	character or numeric vector of cluster IDs

Details

To quantify the purity of a cluster (or bubble) i with n_i number of cells, each of which carries one of L possible labels (e.g. cell type), we computed the Gini impurity index:

$$GI_i = \sum_{j=1}^{L} \pi_{ij} (1 - \pi_{ij}),$$

with π_{ij} as the relative frequency of label j in cluster i. In homogeneous ('pure') clusters most cells carry a distinct label. Hence, the π 's are close to either 1 or 0, and GI takes on a small value close to zero. In 'impure' clusters cells carry a mixture of different labels. In this case most π are far from either 1 or 0, and GI diverges from 0 and approaches 1. If the relative frequencies of the different labels in cluster i are equal to the (background) relative frequencies of the labels in the sample, then cluster i is completely 'impure'.

To compute the overall Gini impurity of a bubbletree, which represents a clustering solution with k bubbles, we estimated the weighted Gini impurity (WGI) by computing the weighted (by the cluster size) average of the GIs:

 $WGI = \sum_{i=1}^{k} GI_i n_i / n,$

with n_i as the number of cells in cluster *i* and $n = \sum_i n_i$.

Value

gi	Gini impurity of each bubble
wgi	Weighted Gini impurity index of the bubbletree

Author(s)

Simo Kitanovski <simo.kitanovski@uni-due.de>

See Also

get_k, get_r, get_bubbletree_kmeans, get_bubbletree_dummy, get_bubbletree_graph, get_gini_k, d_500

Examples

get_gini_k

Gini impurity index computed for a list of clustering solutions obtained by functions get_k or get_r and a vector of categorical cell feature labels

Description

Given The Gini impurity (GI) index allows us to quantitatively evaluate how well a set of labels (categorical features) are split across a set of bubbles. We have a completely perfect split (GI = 0) when each bubble is 'pure', i.e. each bubble contains labels coming from distinct a class. In contrast to this, we have completely imperfect split (GI = 1) when the relative frequency distribution of the labels in each bubble is identical to the background relative frequency distribution of the labels.

Cell type predictions are a type of categorical features that are often used to evaluate the goodness of the clustering. get_gini_k takes as input: 1) a vector of labels for each cell (e.g. cell types) and 2) object returned by function get_k or get_r . Then it computes for each k or r the cluster purity and weightred gini impurity of each clustering solution mean GI, which is another way of finding an optimal clustering resolution.

Usage

get_gini_k(labels, obj)

Arguments

labels	character/factor vector of labels
obj	object returned by functions get_k or get_r

Details

To quantify the purity of a cluster (or bubble) i with n_i number of cells, each of which carries one of L possible labels (e.g. cell type), we computed the Gini impurity index:

 $GI_i = \sum_{j=1}^{L} \pi_{ij} (1 - \pi_{ij}),$

with π_{ij} as the relative frequency of label j in cluster i. In homogeneous ('pure') clusters most cells carry a distinct label. Hence, the π 's are close to either 1 or 0, and GI takes on a small value close to zero. In 'impure' clusters cells carry a mixture of different labels. In this case most π are far from either 1 or 0, and GI diverges from 0 and approaches 1. If the relative frequencies of the different labels in cluster i are equal to the (background) relative frequencies of the labels in the sample, then cluster i is completely 'impure'.

To compute the overall Gini impurity of a bubbletree, which represents a clustering solution with k bubbles, we estimated the weighted Gini impurity (WGI) by computing the weighted (by the cluster size) average of the GIs:

$$WGI = \sum_{i=1}^{\kappa} GI_i n_i / n_i$$

with n_i as the number of cells in cluster *i* and $n = \sum_i n_i$.

get_k

Value

gi_summary	GI for each bubble of a clustering solution with clustering resolution k or r
wgi_summary	WGI for each clustering solution with clustering resolution k or r

Author(s)

Simo Kitanovski <simo.kitanovski@uni-due.de>

See Also

get_k, get_r, get_gini, get_bubbletree_kmeans, get_bubbletree_graph, get_bubbletree_dummy, d_500, get_num_tiles, get_num_violins, get_cat_tiles

Examples

```
# input data
data("d_500", package = "scBubbletree")
A <- d_500$A
f <- d_500$f
b_k \leq get_k(x = A)
           ks = 1:5,
           B_gap = 5,
           n_start = 100,
           iter_max = 200,
           kmeans_algorithm = "MacQueen",
           cores = 1)
b_r <- get_r(x = A)
            rs = c(0.1, 0.5, 1),
            B_gap = 5,
            n_{start} = 20,
            iter_max = 100,
            algorithm = "original",
            cores = 1)
get_gini_k(labels = f, obj = b_k)
get_gini_k(labels = f, obj = b_r)
```

```
get_k
```

Finding optimal number k of clusters

Description

To perform k-means clustering we must specify a number k of clusters. Data-driven metrics, such as the Gap statistic or the within-cluster sum of squares (WCSS), can be used to infer appropriate k from the data. get_k computes the Gap statistic and WCSS for a number of clusters ks.

Usage

```
get_k(x,
    ks,
    B_gap = 20,
    n_start = 1000,
    iter_max = 300,
    kmeans_algorithm = "MacQueen",
    cores = 1,
    verbose = TRUE)
```

Arguments

х	numeric matrix A^{nxf} with n cells, and f low-dimensional projections	
ks	integer vector, k values to consider	
B_gap	integer, number of Monte Carlo ("bootstrap") samples taken when computing the Gap statistic (see documentation of function clusGap, R-package cluster)	
n_start,iter_max,kmeans_algorithm		
	parameters for k-means clustering, see documentation of function kmeans, $R\mbox{-}\xspackage \mbox{stats}$	
cores	integer, number of PC cores for parallel execution	
verbose	logical, progress messages	

Details

To compute the Gap statistic get_k adapts the algorithm of function clustGap from R-package cluster (version 2.1.3). For k-means clustering get_k uses the function kmeans implemented in R-package stats (version 4.2.0). For additional information see the respective documentations.

Value

boot_obj	The results: k-means clustering solutions, the Gap statistic and WCSS	
gap_stats_sum	nmary, wcss_stats_summary	
	main results; Gap statistic and WCSS estimates. Means, standard errors and 95% confidence intervals are provided for each k	
gap_stats, wcs	ss_stats	
	intermediate results; Gap statistic and WCSS estimates for each k and bootstrap iteration b	

Author(s)

Simo Kitanovski <simo.kitanovski@uni-due.de>

See Also

get_r, get_bubbletree_dummy, get_bubbletree_graph, get_bubbletree_kmeans, get_gini, get_gini_k, d_500, get_num_tiles, get_num_violins, get_cat_tiles

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get_num_cell_tiles

Examples

b\$gap_stats_summary

get_num_cell_tiles Visualization of numeric features of individual cells using tile plots

Description

get_num_cell_tiles creates one heatmap from the cells in each bubble. The heatmap visualizes a gradient of the sorted (from high to low) values of a numeric feature (e.g. expression of a certain gene) among the cells of that bubble.

Usage

Arguments

btd	bubbletree object	
f	numeric vector, numeric cell feature	
x_axis_name	character, x-axis title	
feature_name	character, color legend title	
rotate_x_axis_labels		
	<pre>logical, should the x-axis labels be shown horizontally (rotate_x_axis_labels = FALSE) or vertically (rotate_x_axis_labels = TRUE)</pre>	
tile_bw	logical, tile grayscale (tile_bw = TRUE) vs. color (tile_bw = FALSE, default)	

get_num_cell_tiles uses two main inputs:

1. bubbletree object

2. numeric vector of a numeric cell feature.

The order of the cells used to generate the bubbletree (input 1.) should correspond to the order of cell features in input vector f (input 2.)

This function does the following procedure for each bubble: 1. sort and rank the cells in each bubble: rank = 1 for the cell with the highest f value, rank = $n\$ for the bubble with the lowest f value 2. draw a heatmap with x=rank, y=bubble, tile-color=f

Value

plot	ggplot2, tile plot
table	data.frame, raw data used to generate the plot

Author(s)

Simo Kitanovski <simo.kitanovski@uni-due.de>

See Also

get_k, get_r get_bubbletree_dummy, get_bubbletree_kmeans, get_bubbletree_graph, get_gini, get_gini_k, get_cat_tile, get_num_tiles, get_num_violins, d_500, d_ccl

Examples

```
# input data
data("d_500", package = "scBubbletree")
A <- d_500$A
f <- as.vector(d_500$fs[,1])
b <- get_bubbletree_kmeans(x = A, k = 8)
g <- get_num_cell_tiles(btd = b, f = f)
b$tree|g$plot</pre>
```

get_num_tiles Visualization of numeric cell features using tile plots

Description

get_num_tiles creates tile plot to visualize a summary (e.g. mean, median or sum) of a numeric cell feature (e.g. gene expression of a specific gene) in each bubble of a bubbletree

get_num_tiles

Usage

Arguments

btd	bubbletree object	
fs	numeric vector or matrix, numeric cell features	
summary_function	on	
	character, "mean", "median" or "sum", "pct nonzero", "pct zero", summaries are allowed	
round_digits	integer, number of decimal places to keep when showing the relative frequency of cells in each bubble	
<pre>tile_text_size</pre>	integer, size of tile labels	
x_axis_name	character, x-axis title	
rotate_x_axis_labels		
	<pre>logical, should the x-axis labels be shown horizontally (rotate_x_axis_labels = FALSE) or vertically (rotate_x_axis_labels = TRUE)</pre>	
tile_bw	logical, tile grayscale (tile_bw = TRUE) vs. color (tile_bw = FALSE, default)	

Details

get_num_tiles uses two main inputs:

1. bubbletree object

2. numeric vector or matrix of numeric cell features.

The order of the cells used to generat the bubbletree (input 1.) should correspond to the order of cells in the vector/matrix of numeric cell features (input 2.)

This function computes summaries of numeric cell feature in each bubble: 1. mean = mean of feature 2. median = median of feature 3. sum = sum of feature 4. pct nonzero = sum of cells with feature > 0 5. pct zero = sum of cells with feature = 0

Important note: NA and NULL values are omitted.

Value

plot	ggplot2, tile plot
table	data.frame, raw data used to generate the plot

Author(s)

Simo Kitanovski <simo.kitanovski@uni-due.de>

See Also

get_k, get_r get_bubbletree_dummy, get_bubbletree_kmeans, get_bubbletree_graph, get_gini, get_gini_k, get_cat_tile, get_num_violins, d_500, d_ccl

Examples

b\$tree|g\$plot

get_num_violins Visualization of numeric cell features using violin plots

Description

get_num_violins creates violin plot to visualize the distribution of of numeric cell features (e.g. gene expressions) in each bubble of a bubbletree

Usage

get_num_violins(btd,

fs, x_axis_name = "Feature distribution", rotate_x_axis_labels = TRUE)

Arguments

btd	bubbletree object	
fs	numeric vector or matrix, numeric cell features	
x_axis_name	character, x-axis title	
rotate_x_axis_labels		
	logical, should the x-axis labels be shown horizontally (rotate_x_axis_labels	
	= FALSE) or vertically (rotate_x_axis_labels = TRUE)	

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get_r

Details

get_num_violins uses two main inputs:

1. bubbletree object

2. numeric vector or matrix of numeric cell features.

The order of the cells used to generat the bubbletree (input 1.) should correspond to the order of cells in the vector/matrix of numeric cell features (input 2.)

This function visualizes densities of numeric cell feature in the different bubble.

Value

plot	ggplot2, violin plot
table	data.frame, raw data used to generate the plot

Author(s)

Simo Kitanovski <simo.kitanovski@uni-due.de>

See Also

get_k, get_r get_bubbletree_dummy, get_bubbletree_kmeans, get_bubbletree_graph, get_gini, get_gini_k, get_cat_tile, get_num_tiles, d_500

Examples

b\$tree|g\$plot

get_r

Finding optimal clustering resulution r and number of communities k'

Description

To perform Louvain clustering we must specify a clustering resulution r. Data-driven metrics, such as the Gap statistic or the within-cluster sum of squares (WCSS) can be used to infer appropriate r from the data. get_r computes the Gap statistic and WCSS for a vector of clustering resolutions rs.

Usage

```
get_r(x,
    rs,
    B_gap = 20,
    n_start = 20,
    iter_max = 100,
    algorithm = "original",
    knn_k = 20,
    cores = 1,
    verbose = TRUE)
```

Arguments

x	numeric matrix A^{nxf} with n cells, and f low-dimensional projections	
rs	number vector, r values to consider	
B_gap	integer, number of Monte Carlo ("bootstrap") samples taken when computing the Gap statistic (see documentation of function clusGap, R-package cluster)	
n_start,iter_max		
	parameters for Louvain clustering, see documentation of function FindClusters, $R\mbox{-}package$ Seurat	
algorithm	character, four clustering algorithms: 'original', 'LMR', 'SLM' and 'Leiden', see documentation of function FindClusters, R-package Seurat	
knn_k	integer, defines k for the k-nearest neighbor algorithm, see documentation of function FindClusters, R-package Seurat	
cores	integer, number of PC cores for parallel execution	
verbose	logical, progress messages	

Details

To compute the Gap statistic get_r adapts the algorithm of function clustGap from R-package cluster (version 2.1.3). For Louvain clustering get_r uses the function FindClusters implemented in the R-package Seurat. For additional information see the respective documentations.

Value

boot_obj	The results: k-means clustering solutions, the Gap statistic and WCSS	
gap_stats_sum	nary, wcss_stats_summary	
	main results; Gap statistic and WCSS estimates. Means, standard errors and 95% confidence intervals are provided for each r and k'	
gap_stats, wcss_stats		
	intermediate results; Gap statistic and WCSS estimates for each r and k' and	
	bootstrap iteration b	

Author(s)

Simo Kitanovski <simo.kitanovski@uni-due.de>

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get_r

See Also

get_k, get_bubbletree_dummy, get_bubbletree_graph, get_bubbletree_kmeans, get_gini, get_gini_k, d_500, get_num_tiles, get_num_violins, get_cat_tiles, d_ccl

Examples

```
iter_max = 100,
algorithm = "original",
cores = 1,
verbose = TRUE)
```

b\$gap_stats_summary

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