# Package 'cellmarkeraccordion'

December 2, 2024

**Title** Annotation of normal and aberrant hematopoietic cell types in single-cell datasets **Version** 0.9.5

Description cellmarkeraccordion is an R package designed to obtain a robust identification of normal and aberrant hematopoietic cell types in single-cell datasets and easy interpretation of the results. The cellmarkeraccordion package allows to automatically annotate normal and disease ritical cell populations based on the built-in Accordion gene marker database. It requires in input only the counts matrix (raw or normalized) or a Seurat object. In addition, by exploiting the built-in cell cycle labeling you can easily assign the cell cycle phase to each cell. The users can also customize the annotation by simply providing their own genes list as input, which can be associated with cell types or even to specific pathways or signatures of interest. Importantly, the Accordion implements novel options to explore annotation results by inspecting for each group of cells the top marker genes which mostly impacted the annotation, together with the top cell types and their relationship based on the cell ontology tree.

```
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      plyr (>= 1.8.8),
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      ontoProc (>= 1.20.0),
      ontologyIndex (\geq 2.10),
      igraph (>= 1.4.2),
      ontologyPlot (>= 1.6),
      ggraph (>= 2.1.0),
      cowplot(>= 1.1.0),
      ggnewscale (>= 0.4.0),
      purrr (>= 1.0.0),
```

## R topics documented:

accordion																				2
accordion_	cell	_cy	cle						 				 							6

Index		22
	list_tissues	20
	list_disease_tissues	
	list_diseases	
	list_celltypes	19
	list_aberrant_celltypes	18
	accordion_plot	17
	accordion_disease	13
	accordion_custom	

accordion

Automatically annotating and interpreting single-cell populations with the built-in Cell Marker Accordion database

## **Description**

This function performs cell types annotation exploiting the built-in Accordion gene marker database. It takes in input either a Seurat object or a raw or normalized count matrix and return in output the cell types assignment and the detailed informations of the annotation results (added to the Seurat object or as a list).

## Usage

```
accordion(
 data,
  cluster_info = "seurat_clusters",
  assay = "RNA",
 CL_celltypes = NULL,
  species = "Human",
  tissue = NULL,
  include_descendants = FALSE,
 evidence_consistency_score_threshold = NULL,
  specificity_score_threshold = NULL,
  log2FC_threshold = NULL,
 min_n_marker = 5,
 max_n_marker = NULL,
  combined_score_quantile_threshold = NULL,
 annotation_resolution = "cluster",
 cluster_score_quantile_threshold = 0.75,
 allow_unknown = TRUE,
  annotation_name = "accordion",
  include_detailed_annotation_info = TRUE,
 condition_group_info = NULL,
 CL_celltype_group_info = NULL,
 group_markers_by = "celltype_cluster",
  top_cell_score_quantile_threshold = 0.9,
 n_{top_celltypes} = 5,
 n_{top_markers} = 5,
  top_marker_score_quantile_threshold = 0.75,
 plot = TRUE,
  color_by = "cell_type"
```

#### **Arguments**

data

Either a Seurat object (version 4 or 5) or a raw or normalized count matrix with genes on rows and cells on columns. If raw counts are provided, data are lognormalized exploiting the NormalizeData() function from the Seurat package.

cluster\_info

in case object is a Seurat object, cluster\_info should be need to be a character string specifying the name of the column in the metadata that contains cluster ids; if object is a count matrix, cluster\_info should be need to be a data frame or data table containing cluster identity for each cell. The data frame or data table should contain at least two columns, one named "cell", which specifies cell id's, and one named "cluster", which specifies the clustering id's for each cell. This parameter is necessary only when the input is a count matrix and only if the annotation\_resolution parameter is set to "cluster". Default is "seurat clusters".

Character string specifying the Assay of the Seurat object. This parameter is assay necessary only in case data is a Seurat object. Default is "RNA".

Character string or character string vector specifying the cell types to annotate. Run the function "list\_celltypes()" to obtain the available cell types. If this parameter is not specified, all cell types present in the Accordion database are used

for the annotation. Default is NULL.

Character string or character string vector specifying the species. Currently, species either "Human" and/or "Mouse" are supported. If multiple species are selected,

marker genes are merged together. Default is "Human".

Character string or character string vector specifying the tissue. Run the function tissue "list\_tissues()" to obtain the available tissues. If NULL, all tissues information

are aggregated together. Default is NULL.

evidence\_consistency\_score\_threshold

Integer value (currently in 1,7) specifying the minimum evidence consistency (EC) score for each marker. Only markers >= this threshold are kept. If NULL, no filter is applied. Default is NULL.

specificity\_score\_threshold

numeric value in (0,1] specifying the minimum specificity score for each marker. Only markers <= this threshold are kept. If NULL, no filter is applied. Default is NULL.

log2FC\_threshold

numeric value specifying the minimum log2FC threshold for each marker reporting this information. Only markers <= this threshold or without any log2FC are kept. If NULL, no filter is applied. Default is NULL.

Integer value specifying the minimum number of markers to keep for each cell min\_n\_marker type. Only cell types with a number of markers >= this threshold are kept.

Default is 5.

max\_n\_marker Integer value specifying the maximum number of markers to keep for each cell type. For the selection, markers are ranked according to their combined score,

obtained by multiplying evidence consistency score and specificity score. If

NULL, no filter is applied. Default is NULL.

combined\_score\_quantile\_threshold

numeric value in (0,1] specifying the combined score quantile threshold. For the selection, markers are ranked according to their combined score, obtained by multiplying evidence consistency score and specificity score. Only markers > the quantile\_threshold are kept. If NULL, no filter is applied. Default is NULL.

CL\_celltypes

#### annotation\_resolution

Character string or character string vector specifying the resolution of the annotation. Either "cluster" and/or "cell" are supported. Default is "cluster".

#### cluster\_score\_quantile\_threshold

numeric value in 0,1 specifying the cluster score quantile threshold. For each cell a score specific for each cell type is computed. To annotate a cluster cl, for each cell type the cluster\_score\_quantile\_threshold is computed across cells belonging to that cluster and the cell type with the maximum score is then assigned to the cluster cl. Default is 0.75.

allow\_unknown

Logical value indicating whether to allow cells or clusters to be labeled as "unknown". If it is set to TRUE, cells or clusters with negative scores are assigned to the "unknown" category. Default is TRUE.

#### annotation\_name

Character string specifying the name of the column in either the metadata of the input Seurat object or in the input cluster\_info where the annotation will be stored. Per cluster and per cell annotation results will be stored in the annotation\_name\_per\_cluster and annotation\_name\_per\_cell columns respectively. If include\_detailed\_annotation\_info parameter is set to TRUE, the detailed information the stored in a list named annotation\_name. Default is "accordion".

## include\_detailed\_annotation\_info

Logical value indicating whether to store information on the top cell types and markers in the output. If TRUE, a nested list named annotation\_name is created. If resolution\_annotation is set to "cluster" and/or "cell, sublists named "cluster\_resolution" and/or "cell\_resolution" are then added. Inside the sublist "detailed\_annotation\_info" the n\_top\_markers markers, group by group\_markers\_by and the n\_top\_celltypes cell types are then included. If a Seurat object is provided as input the list is stored in the misc slot of the object (object@misc@annotation\_name). If the input is a count matrix, the list is returned in the final output. Default is TRUE.

#### condition\_group\_info

in case object is a Seurat object, condition\_group\_info should be need to be a character string specifying the name of the column in the metadata that contains condition ids for each cell; if object is a count matrix, condition\_group\_info should be need to be a data frame or data table containing condition identity for each cell. The data frame or data table should contain at least two columns, one named "cell", which specifies cell id's, and one named "condition", which specifies the condition id's for each cell. Default is NULL.

## CL\_celltype\_group\_info

in case object is a Seurat object, CL\_celltype\_group\_info should be need to be a character string specifying the name of the column in the metadata that contains cell types ids for each cell; if object is a count matrix, CL\_celltype\_group\_info should be need to be a data frame or data table containing cell types identity for each cell. The data frame or data table should contain at least two columns, one named "cell", which specifies cell id's, and one named "CL\_celltype", which specifies the cell types for each cell. Default is NULL.

#### group\_markers\_by

Character string or character string vector specifying the classification of marker genes. It possible to retrieve n\_top\_markers marker genes for each cell type identified with cluster ("celltype\_cluster") or cell ("celltype\_cell") resolution; n\_top\_markers marker genes per cluster ("cluster") or per cell ("cell") can be also obtained. Additionally, by setting group\_markers\_by to "score\_cell", the

n\_top\_markers marker genes only for cells with a score greater than top\_cell\_score\_quantile\_that are retrieved. Either "celltype\_cluster", "celltype\_cell", "cluster", "cell" or "score\_cell".

Default is "celltype\_cluster".

top\_cell\_score\_quantile\_threshold

numeric value in (0,1] specifying the cell score quantile threshold. For each cell type a score specific for each cell is computed. The top\_cell\_score\_quantile\_threshold is computed across cells belonging to the same cell type, and only cells with a score greater than the top\_cell\_score\_quantile\_threshold are kept. This parameter is necessary only when group\_markers\_by is set to "score\_cell". Default is 0.90.

n\_top\_celltypes

Integer value specifying the number of the top cell types to be included in the output for each cluster and cell depending on the selected annotation\_resolution parameter Default is 5.

n\_top\_markers Integer value specifying the number of the top markers to be included in the output for each cell type, cluster or cell depending on the selected annotation\_resolution and group\_markers\_by parameters. Default is 5.

top\_marker\_score\_quantile\_threshold

numeric value in (0,1] specifying the marker score quantile threshold. For each marker a score specific for each cell is computed. To identify the n\_top\_markers for a cluster cl or a cell type ct, the top\_marker\_score\_quantile\_threshold is computed across cells belonging to that cluster or labeled as ct, and the n\_top\_markers with the maximum score are reported. Default is 0.75.

Logical value indicating whether to store plots displaying detailed annotation information. This parameter can be set to TRUE only when include\_detailed\_annotation\_info is set to TRUE. If TRUE, lollipop plots displaying the top n\_top\_markers group by group\_markers\_by and top n\_top\_celltypes for each annotation\_resolution together with the cell types hierarchies based on the cell ontology structure are

stored in the annotation\_name list. Default is TRUE.

Character string specifying if the plot reporting the top cell types for each cluster/cell is colored based on the assigned cell type ("CL\_celltype") or on cluster id ("cluster"). Default is "CL\_celltype.

## **Details**

If a Seurat object was provided in input, the function returns the Seurat object with markers-based scaled data in the scale.data slot and cell types annotation results in the metadata. If include\_detailed\_annotation\_in and plot were set to TRUE, a list containing cell types and markers information, together with ggplot objects, is stored in the "misc@annotation\_name" slot. If a count matrix was provided in input, the function returns a list containing the following elements:

"scaled\_matrix":normalized and scaled expression matrix;

If annotation\_resolution is set to "cell":

• "cell\_annotation":data table containing cell types annotation results for each cell;

If annotation\_resolution is set to "cluster":

• "cluster\_annotation":data table containing cell types annotation results for each cell;

If include\_detailed\_annotation\_info is set to TRUE:

• "annotation\_name":list containing detailed information of cell types annotation.

plot

color\_by

6 accordion\_cell\_cycle

#### Value

A Seurat object or a list

accordion\_cell\_cycle Automatically identify and interpreting cell cycle state of single-cell populations

## Description

This function identifies cell cycle states exploiting the collection of marker genes associated to each phase, including G0. It takes in input either a Seurat object or a raw or normalized count matrix and return in output the cell cycle assignment and the detailed informations of the annotation results (added to the Seurat object or as a list).

#### Usage

```
accordion_cell_cycle(
  data,
  cluster_info = "seurat_clusters",
  assay = "RNA",
  species = "Human",
  annotation_resolution = "cell",
  annotation_name = "accordion_cell_cycle",
  cluster_score_quantile_threshold = 0.75,
  allow_unknown = FALSE,
  include_detailed_annotation_info = FALSE,
  condition_group_info = NULL,
  cell_type_group_info = NULL,
  group_markers_by = "celltype_cell",
 n_top_celltypes = 5,
 n_{top_markers} = 5,
  top_marker_score_quantile_threshold = 0.75,
  plot = FALSE
)
```

## **Arguments**

data

Either a Seurat object (version 4 or 5) or a raw or normalized count matrix with genes on rows and cells on columns. If raw counts are provided, data are lognormalized exploiting the NormalizeData() function from the Seurat package.

cluster\_info

in case object is a Seurat object, cluster\_info should be need to be a character string specifying the name of the column in the metadata that contains cluster ids; if object is a count matrix, cluster\_info should be need to be a data frame or data table containing cluster identity for each cell. The data frame or data table should contain at least two columns, one named "cell", which specifies cell id's, and one named "cluster", which specifies the clustering id's for each cell. This parameter is necessary only when the input is a count matrix and only if the annotation\_resolution parameter is set to "cluster". Default is "seurat\_clusters".

accordion\_cell\_cycle 7

assay Character string specifying the Assay of the Seurat object. This parameter is

necessary only in case data is a Seurat object. Default is "RNA".

species Character string or character string vector specifying the species. Currently,

either "Human" and/or "Mouse" are supported. If multiple species are selected,

marker genes are merged together. Default is "Human".

annotation\_resolution

Character string or character string vector specifying the resolution of the annotation. Either "cluster" and/or "cell" are supported. Default is "cell".

annotation\_name

Character string specifying the name of the column in either the metadata of the input Seurat object or in the input cluster\_info where the annotation will be stored. Per cluster and per cell annotation results will be stored in the annotation\_name\_per\_cluster and annotation\_name\_per\_cell columns respectively. If include\_detailed\_annotation\_info parameter is set to TRUE, the detailed information the stored in a list named annotation\_name. Default is "accordion\_cell\_cycle".

cluster\_score\_quantile\_threshold

numeric value in 0,1 specifying the cluster score quantile threshold. For each cell a score specific for each cell type is computed. To annotate a cluster cl, for each cell type the cluster\_score\_quantile\_threshold is computed across cells belonging to that cluster and the cell type with the maximum score is then assigned to the cluster cl. Default is 0.75.

allow\_unknown

Logical value indicating whether to allow cells or clusters to be labeled as "unknown". If it is set to TRUE, cells or clusters with negative scores are assigned to the "unknown" category. Default is TRUE.

include\_detailed\_annotation\_info

Logical value indicating whether to store information on the top cell types and markers in the output. If TRUE, a nested list named annotation\_name is created. If resolution\_annotation is set to "cluster" and/or "cell, sublists named "cluster\_resolution" and/or "cell\_resolution" are then added. Inside the sublist "detailed\_annotation\_info" the n\_top\_markers markers, group by group\_markers\_by and the n\_top\_celltypes cell types are then included. If a Seurat object is provided as input the list is stored in the misc slot of the object (object@misc@annotation\_name). If the input is a count matrix, the list is returned in the final output. Default is FALSE. @param condition group info in case object is a Seurat object, condition\_group\_info should be need to be a character string specifying the name of the column in the metadata that contains condition ids for each cell; if object is a count matrix, condition\_group\_info should be need to be a data frame or data table containing condition identity for each cell. The data frame or data table should contain at least two columns, one named "cell", which specifies cell id's, and one named "condition", which specifies the condition id's for each cell. Default is NULL. @param cell\_type\_group\_info in case object is a Seurat object, cell\_type\_group\_info should be need to be a character string specifying the name of the column in the metadata that contains cell types ids for each cell; if object is a count matrix, cell\_type\_group\_info should be need to be a data frame or data table containing cell types identity for each cell. The data frame or data table should contain at least two columns, one named "cell", which specifies cell id's, and one named "cell\_type", which specifies the cell types for each cell. Default is NULL.

group\_markers\_by

Character string or character string vector specifying the classification of marker genes. It possible to retrieve n\_top\_markers marker genes for each cell type

8 accordion\_cell\_cycle

identified with cluster ("celltype\_cluster") or cell ("celltype\_cell") resolution; n\_top\_markers marker genes per cluster ("cluster") or per cell ("cell") can be also obtained. Either "celltype\_cluster", "celltype\_cell", "cluster" and/or "cell". Default is "celltype\_cell".

n\_top\_celltypes

Integer value specifying the number of the top cell types to be included in the output for each cluster and cell depending on the selected annotation\_resolution parameter Default is 5.

n\_top\_markers

Integer value specifying the number of the top markers to be included in the output for each cell type, cluster or cell depending on the selected annotation\_resolution and group\_markers\_by parameters. Default is 5.

top\_marker\_score\_quantile\_threshold

numeric value in (0,1] specifying the marker score quantile threshold. For each marker a score specific for each cell is computed. To identify the n\_top\_markers for a cluster cl or a cell type ct, the top\_marker\_score\_quantile\_threshold is computed across cells belonging to that cluster or labeled as ct, and the n\_top\_markers with the maximum score are reported. Default is 0.75.

plot

Logical value indicating whether to store plots displaying detailed annotation information. This parameter can be set to TRUE only when include\_detailed\_annotation\_info is set to TRUE. If TRUE, lollipop plots displaying the top n\_top\_markers group by group\_markers\_by and n\_top\_celltypes for each annotation\_resolution together with the cell types hierarchies based on the cell ontology structure are stored in the annotation\_name list. Default is FALSE.

#### **Details**

If a Seurat object was provided in input, the function returns the Seurat object with markers-based scaled data in the scale.data slot and cell types annotation results in the metadata. If include\_detailed\_annotation\_in and plot were set to TRUE, a list named annotation\_name containing cell types and markers information, together with ggplot objects, is stored in the "misc" slot. If a count matrix was provided in input, the function returns a list containing the following elements:

"scaled\_matrix":normalized and scaled expression matrix;

If annotation\_resolution is set to "cell":

• "cell\_annotation":data table containing cell types annotation results for each cell;

If annotation\_resolution is set to "cluster":

• "cluster\_annotation":data table containing cell types annotation results for each cell;

If include\_detailed\_annotation\_info is set to TRUE:

• "annotation\_name":list containing detailed information of cell types annotation.

## Value

A Seurat object or a list

accordion\_custom

Automatically annotating and interpreting single-cell populations with custom marker genes sets

## **Description**

This function performs cell types or signatures/pathways annotation based on cusom marker genes set. It takes in input either a Seurat object or a raw or normalized count matrix and a table of marker genes associated to cell types or even to pathways and return in output the cell types/pathways assignment and the detailed informations of the annotation results (added to the Seurat object or as a list).

## Usage

```
accordion_custom(
  data,
 marker_table,
  category_column = "cell_type",
 marker_column = "marker",
 marker_type_column = "marker_type",
 weight_column = "weight",
  cluster_info = "seurat_clusters",
 assay = "RNA".
 min_n_marker = 5,
 max_n_marker = NULL,
  combined_score_quantile_threshold = NULL,
  annotation_resolution = "cluster",
  cluster_score_quantile_threshold = 0.75,
  allow_unknown = TRUE,
  annotation_name = "accordion_custom",
  include_detailed_annotation_info = TRUE,
  condition_group_info = NULL,
  cell_type_group_info = NULL,
  group_markers_by = "celltype_cluster",
  top_cell_score_quantile_threshold = 0.9,
 n_{top_celltypes} = 5,
 n_{top_markers} = 5,
  top_marker_score_quantile_threshold = 0.75,
 plot = TRUE
)
```

#### Arguments

data

Either a Seurat object (version 4 or 5) or a raw or normalized count matrix with genes on rows and cells on columns. If raw counts are provided, data are lognormalized exploiting the NormalizeData() function from the Seurat package.

marker\_table

Data table or data frame containing cell type markers. The table needs to have at least two columns, the category\_column, which specifies cell types or categories, and the marker\_column, which specifies the corresponding markers on each row. Columns indicating the marker type (either positive or negative), and the marker weight can be optionally included.

category\_column

String characters specifying the name of the marker\_table column containing cell types or categories. Default is "cell\_type".

marker\_column

String characters specifying the name of the marker\_table column containing markers. Default is "marker".

marker\_type\_column

Optional string characters specifying the name of the marker\_table column containing string characters indicating the type of markers, either "positive" or "negative". If no marker\_type\_column is found in the marker\_table all markers are considered "positive". Default is "marker\_type".

weight\_column

Optional string characters specifying the name of the marker\_table column containing numeric value indicating the weight for each marker. If no weight\_column is found in the marker\_table all markers are equally weighted as 1. Default is "weight".

cluster\_info

in case object is a Seurat object, cluster\_info should be need to be a character string specifying the name of the column in the metadata that contains cluster ids; if object is a count matrix, cluster\_info should be need to be a data frame or data table containing cluster identity for each cell. The data frame or data table should contain at least two columns, one named "cell", which specifies cell id's, and one named "cluster", which specifies the clustering id's for each cell. This parameter is necessary only when the input is a count matrix and only if the annotation\_resolution parameter is set to "cluster". Default is "seurat clusters".

assay

Character string specifying the Assay of the Seurat object. This parameter is necessary only in case data is a Seurat object. Default is "RNA".

min\_n\_marker

Integer value specifying the minimum number of markers to keep for each cell type. Only cell types with a number of markers >= this threshold are kept. Default is 5.

max\_n\_marker

Integer value specifying the maximum number of markers to keep for each cell type. For the selection, markers are ranked according to their combined score, obtained by multiplying evidence consistency score and specificity score. If NULL, no filter is applied. Default is NULL.

 ${\tt combined\_score\_quantile\_threshold}$ 

numeric value in (0,1] specifying the combined score quantile threshold. For the selection, markers are ranked according to their combined score, obtained by multiplying weight and specificity score. Only markers > the quantile\_threshold are kept. If NULL, no filter is applied. Default is NULL.

annotation\_resolution

Character string or character string vector specifying the resolution of the annotation. Either "cluster" and/or "cell" are supported. Default is "cluster".

cluster\_score\_quantile\_threshold

numeric value in 0,1 specifying the cluster score quantile threshold. For each cell a score specific for each cell type is computed. To annotate a cluster cl, for each cell type the cluster\_score\_quantile\_threshold is computed across cells belonging to that cluster and the cell type with the maximum score is then assigned to the cluster cl. Default is 0.75.

allow\_unknown

Logical value indicating whether to allow cells or clusters to be labeled as "unknown". If it is set to TRUE, cells or clusters with negative scores are assigned to the "unknown" category. Default is TRUE.

#### annotation\_name

Character string specifying the name of the column in either the metadata of the input Seurat object or in the input cluster\_info where the annotation will be stored. Per cluster and per cell annotation results will be stored in the annotation\_name\_per\_cluster and annotation\_name\_per\_cell columns respectively. If include\_detailed\_annotation\_info parameter is set to TRUE, the detailed information the stored in a list named annotation\_name. Default is "accordion custom".

## $include\_detailed\_annotation\_info$

Logical value indicating whether to store information on the top cell types and markers in the output. If TRUE, a list named annotation\_name is created. If resolution\_annotation is set to "cluster" and/or "cell, sublists named "cluster\_resolution" and/or "cell\_resolution" are then added. Inside the sublist "detailed\_annotation\_info" the n\_top\_markers markers, group by group\_markers\_by and the n\_top\_celltypes cell types are then included. If a Seurat object is provided as input the list is stored in the misc slot of the object (object@misc@annotation\_name). If the input is a count matrix, the list is returned in the final output. Default is TRUE.

## condition\_group\_info

in case object is a Seurat object, condition\_group\_info should be need to be a character string specifying the name of the column in the metadata that contains condition ids for each cell; if object is a count matrix, condition\_group\_info should be need to be a data frame or data table containing condition identity for each cell. The data frame or data table should contain at least two columns, one named "cell", which specifies cell id's, and one named "condition", which specifies the condition id's for each cell. Default is NULL.

## cell\_type\_group\_info

in case object is a Seurat object, cell\_type\_group\_info should be need to be a character string specifying the name of the column in the metadata that contains cell types ids for each cell; if object is a count matrix, cell\_type\_group\_info should be need to be a data frame or data table containing cell types identity for each cell. The data frame or data table should contain at least two columns, one named "cell", which specifies cell id's, and one named "cell\_type", which specifies the cell types for each cell. Default is NULL.

## group\_markers\_by

Character string or character string vector specifying the classification of marker genes. It possible to retrieve n\_top\_markers marker genes for each cell type identified with cluster ("celltype\_cluster") or cell ("celltype\_cell") resolution; n\_top\_markers marker genes per cluster ("cluster") or per cell ("cell") can be also obtained. Additionally, by setting group\_markers\_by to "score\_cell", the n\_top\_markers marker genes only for cells with a score greater than top\_cell\_score\_quantile\_there retrieved. Either "celltype\_cluster", "celltype\_cell", "cluster", "cell" or "score\_cell". Default is "celltype\_cluster".

#### top\_cell\_score\_quantile\_threshold

numeric value in (0,1] specifying the cell score quantile threshold. For each cell type/signature a score specific for each cell is computed. The top\_cell\_score\_quantile\_threshol is computed across cells belonging to the same cell type/signature, and only cells with a score greater than the top\_cell\_score\_quantile\_threshold are kept. This parameter is necessary only when group\_markers\_by is set to "score\_cell". Default is 0.90.

#### n\_top\_celltypes

Integer value specifying the number of the top cell types to be included in the

output for each cluster and cell depending on the selected annotation\_resolution parameter Default is 5.

n\_top\_markers

Integer value specifying the number of the top markers to be included in the output for each cell type, cluster or cell depending on the selected annotation\_resolution and group\_markers\_by parameters. Default is 5.

top\_marker\_score\_quantile\_threshold

numeric value in (0,1] specifying the marker score quantile threshold. For each marker a score specific for each cell is computed. To identify the n\_top\_markers for a cluster cl or a cell type ct, the top\_marker\_score\_quantile\_threshold is computed across cells belonging to that cluster or labeled as ct, and the n\_top\_markers with the maximum score are reported. Default is 0.75.

plot

Logical value indicating whether to store plots displaying detailed annotation information. This parameter can be set to TRUE only when include\_detailed\_annotation\_info is set to TRUE. If TRUE, lollipop plots displaying the top n\_top\_markers group by group\_markers\_by and op n\_top\_celltypes for each annotation\_resolution together with the cell types hierarchies based on the cell ontology structure are stored in the annotation\_name list. Default is TRUE.

#### **Details**

If a Seurat object was provided in input, the function returns the Seurat object with markers-based scaled data in the scale.data slot and cell types annotation results in the metadata. If include\_detailed\_annotation\_in and plot were set to TRUE, a list containing cell types and markers information, together with ggplot objects, is stored in the "misc@annotation\_name" slot. If a count matrix was provided in input, the function returns a list containing the following elements:

• "scaled\_matrix":normalized and scaled expression matrix;

If annotation\_resolution is set to "cell":

• "cell\_annotation":data table containing cell types annotation results for each cell;

If annotation\_resolution is set to "cluster":

• "cluster\_annotation":data table containing cell types annotation results for each cell;

If include\_detailed\_annotation\_info is set to TRUE:

• "annotation\_name":list containing detailed information of cell types annotation.

#### Value

A Seurat object or a list

accordion\_disease

Automatically annotating and interpreting aberrant single-cell populations with the built-in Cell Marker Accordion disease database

#### **Description**

This function identified aberrant populations exploiting the built-in Accordion gene marker disease database. It takes in input either a Seurat object or a raw or normalized count matrix and return in output the cell types assignment and the detailed informations of the annotation results (added to the Seurat object or as a list).

#### Usage

```
accordion_disease(
  data,
  disease = NULL,
  tissue = NULL,
  cluster_info = "seurat_clusters",
  assay = "RNA",
 NCIT_celltypes = NULL,
  species = "Human",
  include_descendants = FALSE,
  evidence_consistency_score_threshold = NULL,
  specificity_score_threshold = NULL,
  log2FC_threshold = NULL,
 malignant_quantile_threshold = 0.95,
 min_n_marker = 5,
 max_n_marker = NULL,
  combined_score_quantile_threshold = NULL,
  disease_vs_healthy = TRUE,
  annotation_resolution = "cluster",
  cluster_score_quantile_threshold = 0.75,
  allow_unknown = TRUE,
  annotation_name = "accordion_disease",
  include_detailed_annotation_info = TRUE,
  condition_group_info = NULL,
 NCIT_celltype_group_info = NULL,
  group_markers_by = "celltype_cluster",
  top_cell_score_quantile_threshold = 0.9,
 n_{top_celltypes} = 5,
  n_{top_markers} = 5,
  top_marker_score_quantile_threshold = 0.75,
 plot = TRUE,
  color_by = "cell_type"
```

## **Arguments**

data

Either a Seurat object (version 4 or 5) or a raw or normalized count matrix with genes on rows and cells on columns. If raw counts are provided, data are lognormalized exploiting the NormalizeData() function from the Seurat package.

disease Character string or character string vector specifying diseases to consider. Run

the function "list\_diseases()" to obtain the available diseases. If NULL, all dis-

eases are considered. Default is NULL.

tissue Character string or character string vector specifying the tissue. Run the func-

 $tion \ "list\_disease\_tissues()" \ to \ obtain \ the \ available \ tissues. \ If \ NULL, \ all \ tissues$ 

information are aggregated together. Default is NULL.

cluster\_info in case object is a Seurat object, cluster\_info should be need to be a charac-

ter string specifying the name of the column in the metadata that contains cluster ids; if object is a count matrix, cluster\_info should be need to be a data frame or data table containing cluster identity for each cell. The data frame or data table should contain at least two columns, one named "cell", which specifies cell id's, and one named "cluster", which specifies the clustering id's for each cell. This parameter is necessary only when the input is a count matrix and only if the annotation\_resolution parameter is set to "cluster". Default is

"seurat clusters".

assay Character string specifying the Assay of the Seurat object. This parameter is

necessary only in case data is a Seurat object. Default is "RNA".

NCIT\_celltypes Character string or character string vector specifying the cell types to annotate.

Run the function "list\_aberrant\_celltypes()" to obtain the available aberrant cell types. If this parameter is not specified, all aberrant cell types are used for the

annotation. Default is NULL.

species Character string or character string vector specifying the species. Currently, either "Human" and/or "Mouse" are supported. If multiple species are selected,

marker genes are merged together. Default is "Human".

evidence\_consistency\_score\_threshold

Integer value (currently in 1,7) specifying the minimum evidence consistency (EC) score for each marker. Only markers >= this threshold are kept. If NULL,

no filter is applied. Default is NULL.

specificity\_score\_threshold

numeric value in (0,1] specifying the minimum specificity score for each marker. Only markers <= this threshold are kept. If NULL, no filter is applied. Default

is NULL.

min\_n\_marker Integer value specifying the minimum number of markers to keep for each cell

type. Only cell types with a number of markers >= this threshold are kept.

Default is 5.

max\_n\_marker Integer value specifying the maximum number of markers to keep for each cell

type. For the selection, markers are ranked according to their combined score, obtained by multiplying evidence consistency score and specificity score. If

NULL, no filter is applied. Default is NULL.

combined\_score\_quantile\_threshold

numeric value in (0,1] specifying the combined score quantile threshold. For the selection, markers are ranked according to their combined score, obtained by multiplying evidence consistency score and specificity score. Only markers > the quantile\_threshold are kept. If NULL, no filter is applied. Default is

NULL.

disease\_vs\_healthy

Logical value indicating whether to compare the markers associated with disease cell types with respect to the markers associated with the corresponding healthy cell types. If TRUE the specificity score is calculated considering if a gene is also a marker for the corresponding healthy cell type. For each cell, a specific

score for disease cell types and another score for the healthy types (named as the cell types label) are returned. Default is TRUE.

#### annotation\_resolution

Character string or character string vector specifying the resolution of the annotation. Either "cluster" and/or "cell" are supported. Default is "cluster".

#### cluster\_score\_quantile\_threshold

numeric value in 0,1 specifying the cluster score quantile threshold. For each cell a score specific for each cell type is computed. To annotate a cluster cl, for each cell type the cluster\_score\_quantile\_threshold is computed across cells belonging to that cluster and the cell type with the maximum score is then assigned to the cluster cl. Default is 0.75.

allow\_unknown

Logical value indicating whether to allow cells or clusters to be labeled as "unknown". If it is set to TRUE, cells or clusters with negative scores are assigned to the "unknown" category. Default is TRUE.

#### annotation\_name

Character string specifying the name of the column in either the metadata of the input Seurat object or in the input cluster\_info where the annotation will be stored. Per cluster and per cell annotation results will be stored in the annotation\_name\_per\_cluster and annotation\_name\_per\_cell columns respectively. If include\_detailed\_annotation\_info parameter is set to TRUE, the detailed information the stored in a list named annotation\_name. Default is "accordion disease".

#### include\_detailed\_annotation\_info

Logical value indicating whether to store information on the top cell types and markers in the output. If TRUE, a nested list named annotation\_name is created. If resolution\_annotation is set to "cluster" and/or "cell, sublists named "cluster\_resolution" and/or "cell\_resolution" are then added. Inside the sublist "detailed\_annotation\_info" the n\_top\_markers markers, group by group\_markers\_by and the n\_top\_celltypes cell types are then included. If a Seurat object is provided as input the list is stored in the misc slot of the object (object@misc@annotation\_name). If the input is a count matrix, the list is returned in the final output. Default is TRUE.

#### condition\_group\_info

in case object is a Seurat object, condition\_group\_info should be need to be a character string specifying the name of the column in the metadata that contains condition ids for each cell; if object is a count matrix, condition\_group\_info should be need to be a data frame or data table containing condition identity for each cell. The data frame or data table should contain at least two columns, one named "cell", which specifies cell id's, and one named "condition", which specifies the condition id's for each cell. Default is NULL.

## NCIT\_celltype\_group\_info

in case object is a Seurat object, NCIT\_celltype\_group\_info should be need to be a character string specifying the name of the column in the metadata that contains cell types ids for each cell; if object is a count matrix, NCIT\_celltype\_group\_info should be need to be a data frame or data table containing cell types identity for each cell. The data frame or data table should contain at least two columns, one named "cell", which specifies cell id's, and one named "NCIT\_celltype", which specifies the cell types for each cell. Default is NULL.

#### group\_markers\_by

Character string or character string vector specifying the classification of marker genes. It possible to retrieve n\_top\_markers marker genes for each cell type

> identified with cluster ("celltype cluster") or cell ("celltype cell") resolution; n\_top\_markers marker genes per cluster ("cluster") or per cell ("cell") can be also obtained. Additionally, by setting group\_markers\_by to "score\_cell", the n\_top\_markers marker genes only for cells with a score greater than top\_cell\_score\_quantile\_th are retrieved. Either "celltype\_cluster", "celltype\_cell", "cluster", "cell" or "score\_cell". Default is "celltype\_cluster".

top\_cell\_score\_quantile\_threshold

numeric value in (0,1] specifying the cell score quantile threshold. For each cell type a score specific for each cell is computed. The top\_cell\_score\_quantile\_threshold is computed across cells belonging to the same cell type, and only cells with a score greater than the top\_cell\_score\_quantile\_threshold are kept. This parameter is necessary only when group\_markers\_by is set to "score\_cell". Default is 0.90.

n\_top\_celltypes

Integer value specifying the number of the top cell types to be included in the output for each cluster and cell depending on the selected annotation\_resolution parameter Default is 5.

Integer value specifying the number of the top markers to be included in the outn\_top\_markers put for each cell type, cluster or cell depending on the selected annotation\_resolution and group\_markers\_by parameters. Default is 5.

top\_marker\_score\_quantile\_threshold

numeric value in (0,1) specifying the marker score quantile threshold. For each marker a score specific for each cell is computed. To identify the n\_top\_markers for a cluster cl or a cell type ct, the top\_marker\_score\_quantile\_threshold is computed across cells belonging to that cluster or labeled as ct, and the n\_top\_markers with the maximum score are reported. Default is 0.75.

Logical value indicating whether to store plots displaying detailed annotation in-

formation. This parameter can be set to TRUE only when include\_detailed\_annotation\_info is set to TRUE. If TRUE, lollipop plots displaying the top n\_top\_markers group by group\_markers\_by and op n\_top\_celltypes for each annotation\_resolution together with the cell types hierarchies based on the cell ontology structure are

stored in the "accordion" list. Default is TRUE.

#### **Details**

If a Seurat object was provided in input, the function returns the Seurat object with markers-based scaled data in the scale.data slot and cell types annotation results in the metadata. If include\_detailed\_annotation\_in and plot were set to TRUE, a list containing cell types and markers information, together with ggplot objects, is stored in the "misc@annotation\_name" slot. If a count matrix was provided in input, the function returns a list containing the following elements:

"scaled\_matrix":normalized and scaled expression matrix;

If annotation\_resolution is set to "cell":

• "cell\_annotation":data table containing cell types annotation results for each cell;

If annotation\_resolution is set to "cluster":

• "cluster\_annotation":data table containing cell types annotation results for each cell;

If include\_detailed\_annotation\_info is set to TRUE:

• "annotation\_name":list containing detailed information of cell types annotation.

plot

accordion\_plot 17

#### Value

A Seurat object or a list

accordion\_plot

Interpreting annotation results

## **Description**

This function generates lollipop plots displaying the detailed annotation results obtained with the accordion\_annotation, accordion\_disease\_annotation and accordion\_custom\_annotation functions.

## Usage

```
accordion_plot(
  data,
  info_to_plot = "accordion",
  resolution = "cluster",
  group_markers_by = "celltype_cluster",
  color_by = "CL_celltype"
)
```

#### **Arguments**

data

A Seurat object or a list containing "detailed\_annotation\_info", from either accordion(), accordion\_disease() or accordion\_custom() functions, run with include detailed annotation info parameter set to TRUE.

info\_to\_plot

Character string or character string vector specifying the list from which extract the detailed annotation information, either "accordion", "accordion\_disease" or "accordion\_custom", for which returns the plot, either "accordion", "accordion\_disease" or "accordion\_custom".

resolution

Character string or character string vector specifying the annotation resolution for which provided the plots. Either "cluster" and/or "cell" are supported. Default is "cluster".

group\_markers\_by

Character string or character string vector specifying the classification of marker genes. It is possible to retrieve top marker genes for each cell type identified with cluster ("celltype\_cluster") or cell ("celltype\_cell") resolution; top marker genes per cluster ("cluster") or per cell ("cell") can be also obtained. Additionally, by setting group\_markers\_by to "score\_cell", the n\_top\_markers marker genes only for cells with a score greater than top\_cell\_score\_quantile\_threshold are retrieved. Either "celltype\_cluster", "celltype\_cell", "cluster", "cell" or "score\_cell". Default is "celltype cluster".

color\_by

Character string specifying if the plot reporting the top cell types for each cluster/cell is colored based on the assigned cell type ("CL\_celltype") or on cluster id ("cluster"). Default is "CL\_celltype.

#### **Details**

top cell types (or pathways) and top markers It takes in input either a Seurat object or a raw or normalized count matrix and a table of marker genes associated to cell types or even to pathways and return in output the cell types/pathways assignment and the detailed informations of the annotation results (added to the Seurat object or as a list).

If a Seurat object was provided in input, the function returns the Seurat object with a list of ggplot objects added to the "misc" slot in the info\_to\_plot list. If a list was provided in input, the function returns the same list with the addition of the ggplot objects.

### Value

A Seurat object or a list.

```
list_aberrant_celltypes
```

List aberrant cell types available in the Cell Marker Accordion disease database

#### **Description**

List aberrant cell types available in the Cell Marker Accordion disease database

## Usage

```
list_aberrant_celltypes(
  species = c("Human", "Mouse"),
  disease = NULL,
  tissue = NULL
)
```

## **Arguments**

species

Character string or character string vector specifying the species for which to extract the associate list of available cell types. Currently, either "Human" and/or "Mouse" are supported. Default is c("Mouse", "Human"). @param disease Character string or character string vector specifying diseases to consider. If NULL, information from all diseases are considered. Default is NULL. @param tissue Character string or character string vector specifying the tissue for which to extract the associate list of available cell types. If NULL, information from all tissues are retrieved.

## Value

List of aberrant cell types available in the Cell Marker Accordion disease database

list\_celltypes 19

list\_celltypes

List cell types available in the Cell Marker Accordion database

#### **Description**

List cell types available in the Cell Marker Accordion database

#### Usage

```
list_celltypes(species = c("Human", "Mouse"), tissue = NULL)
```

#### **Arguments**

species

Character string or character string vector specifying the species for which to extract the associate list of available cell types. Currently, either "Human" and/or "Mouse" are supported. Default is c("Mouse", "Human"). @param tissue Character string or character string vector specifying the tissue for which to extract the associate list of available cell types. If NULL, information from all tissues are retrieved.

#### Value

List of cell types available in the Cell Marker Accordion database

list\_diseases

List diseases available in the Cell Marker Accordion disease database

## **Description**

List diseases available in the Cell Marker Accordion disease database

#### Usage

```
list_diseases(
  species = c("Human", "Mouse"),
  tissue = NULL,
  aberrant_celltype = NULL
)
```

#### **Arguments**

species

Character string or character string vector specifying the species for which to extract the associate list of available diseases Currently, either "Human" and/or "Mouse" are supported. Default is c("Mouse", "Human"). @param tissue Character string or character string vector specifying the tissue for which to extract the associate list of available diseases. If NULL, information from all tissues are retrieved. @param aberrant\_celltype Character string or character string vector specifying the aberrant celltype for which to extract the associate list of available tissues. If NULL, information from all aberrant cell types are retrieved. @return List of diseases available in the Cell Marker Accordion disease database

20 list\_tissues

#### **Description**

List tissues available in the Cell Marker Accordion disease database

## Usage

```
list_disease_tissues(
  species = c("Human", "Mouse"),
  disease = NULL,
  aberrant_celltype = NULL
)
```

#### **Arguments**

species

Character string or character string vector specifying the species for which to extract the associate list of available tissues. Currently, either "Human" and/or "Mouse" are supported. Default is c("Mouse", "Human"). @param disease Character string or character string vector specifying diseases to consider. If NULL, information from all diseases are considered. Default is NULL. @param celltype Character string or character string vector specifying the celltype for which to extract the associate list of available tissues. If NULL, information from all cell types are retrieved.

## Value

List of tissues available in the Cell Marker Accordion disease database

list\_tissues

List tissues available in the Cell Marker Accordion database

## **Description**

List tissues available in the Cell Marker Accordion database

## Usage

```
list_tissues(species = c("Human", "Mouse"), celltype = NULL)
```

#### Arguments

species

Character string or character string vector specifying the species for which to extract the associate list of available tissues. Currently, either "Human" and/or "Mouse" are supported. Default is c("Mouse", "Human"). @param celltype Character string or character string vector specifying the celltype for which to extract the associate list of available tissues. If NULL, information from all cell types are retrieved.

list\_tissues 21

## Value

List of tissues available in the Cell Marker Accordion database

# **Index**

```
0,1,4,7,10,15
1,7,3,14

accordion,2
accordion_cell_cycle,6
accordion_disease,13
accordion_plot,17

list_aberrant_celltypes,18
list_celltypes,19
list_disease_tissues,20
list_diseases,19
list_tissues,20
```