

Package: SlimR (via r-universe)

June 30, 2026

Version 1.1.6

Title Adaptive Machine Learning-Powered, Context-Matching Tool for Single-Cell and Spatial Transcriptomics Annotation

Description Annotates single-cell and spatial-transcriptomic (ST) data using context-matching marker datasets. It creates a unified marker list (`Markers_list`) from multiple sources: built-in curated databases ('Cellmarker2', 'PanglaoDB', 'ScType', 'scIBD', 'TCellSI', 'PCTIT', 'PCTAM'), Seurat objects with cell labels, or user-provided Excel tables. SlimR first uses adaptive machine learning for parameter optimization, and then offers two automated annotation approaches: 'cluster-based' and 'per-cell'. Cluster-based annotation assigns one label per cluster, expression-based probability calculation, and AUC validation. Per-cell annotation assigns labels to individual cells using three scoring methods with adaptive thresholds and ratio-based confidence filtering, plus optional UMAP spatial smoothing, making it ideal for heterogeneous clusters and rare cell types. The package also supports semi-automated workflows with heatmaps, feature plots, and combined visualizations for manual annotation. For more information, see the package documentation at <<https://github.com/zhaoqing-wang/SlimR>>.

License MIT + file LICENSE

URL <https://github.com/zhaoqing-wang/SlimR>

BugReports <https://github.com/zhaoqing-wang/SlimR/issues>

Depends R (>= 4.1.0)

Imports cowplot, dplyr, ggplot2, patchwork, pheatmap, readxl, scales, Seurat, tidyr, tools, tibble

Suggests crayon, RANN, testthat (>= 3.0.0)

Encoding UTF-8

LazyData true

Date 2026-06-30

Config/roxygen2/version 8.0.0

NeedsCompilation no

Author Zhaoqing Wang [aut, cre] (ORCID:
<<https://orcid.org/0000-0001-8348-7245>>)

Maintainer Zhaoqing Wang <zhaoqingwang@mail.sdu.edu.cn>

Repository <https://cran.r-universe.dev>

Date/Publication 2026-06-30 19:12:24 UTC

RemoteUrl <https://github.com/cran/SlimR>

RemoteRef HEAD

RemoteSha 64fe837571082c41884183c76058383eb33dfd38

Contents

calculate_cluster_variability	3
calculate_expression	4
calculate_expression_skewness	5
calculate_probability	5
Cellmarker2	6
Cellmarker2_raw	7
Cellmarker2_table	8
Celltype_Annotation	8
Celltype_annotation_Cellmarker2	9
Celltype_Annotation_Combined	11
Celltype_annotation_Excel	12
Celltype_Annotation_Features	14
Celltype_Annotation_Heatmap	16
Celltype_annotation_PanglaoDB	18
Celltype_Annotation_PerCell	20
Celltype_annotation_Seurat	21
Celltype_Calculate	23
Celltype_Calculate_PerCell	25
Celltype_Compare	28
Celltype_Verification	30
Celltype_Verification_PerCell	31
compute_adaptive_parameters	33
Compute_Gene_AUC_ROC	33
estimate_batch_effect	35
extract_dataset_features	36
Markers_filter_Cellmarker2	37
Markers_filter_PanglaoDB	38
Markers_filter_ScType	39
Markers_list_PCTAM	40
Markers_list_PCTIT	40
Markers_list_scIBD	41
Markers_list_TCellSI	42
PanglaoDB	43

<i>calculate_cluster_variability</i>	3
PanglaoDB_raw	43
PanglaoDB_table	44
Parameter_Calculate	45
percell_workflow	46
plot.heatmap	50
Read_excel_markers	51
Read_seurat_markers	52
ScType	54
ScType_raw	55
ScType_table	55
Index	57

`calculate_cluster_variability`
Calculate Cluster Variability (Use in package)

Description

Measures the degree of separation between different cell clusters based on expression patterns.

Usage

`calculate_cluster_variability(data.features, features)`

Arguments

- `data.features` Data frame containing expression data and cluster labels
- `features` Feature names to include in analysis

Value

Numeric value representing cluster separation strength

See Also

Other Section_1_Functions_Use_in_Package: [calculate_expression\(\)](#), [calculate_expression_skewness\(\)](#), [calculate_probability\(\)](#), [compute_adaptive_parameters\(\)](#), [estimate_batch_effect\(\)](#), [extract_dataset_features\(\)](#)

calculate_expression *Counts average expression of gene set (Use in package)*

Description

Counts average expression of gene set (Use in package)

Usage

```
calculate_expression(  
  object,  
  features,  
  assay = NULL,  
  cluster_col = NULL,  
  colour_low = "white",  
  colour_high = "navy"  
)
```

Arguments

object	Enter a Seurat object.
features	Enter one or a set of markers.
assay	Enter the assay used by the Seurat object, such as "RNA". Default parameters use "assay = NULL".
cluster_col	Enter the meta.data column in the Seurat object to be annotated, such as "seurat_cluster". Default parameters use "cluster_col = NULL".
colour_low	Color for lowest expression level. (default = "white")
colour_high	Color for highest expression level. (default = "black")

Value

Average expression genes and related informations in the input "Seurat" object given "cluster_col" and given "features".

See Also

Other Section_1_Functions_Use_in_Package: [calculate_cluster_variability\(\)](#), [calculate_expression_skewness\(\)](#), [calculate_probability\(\)](#), [compute_adaptive_parameters\(\)](#), [estimate_batch_effect\(\)](#), [extract_dataset_features\(\)](#)

`calculate_expression_skewness`*Calculate Expression Distribution Skewness (Use in package)*

Description

Computes the average skewness of gene expression distributions across all features.

Usage

```
calculate_expression_skewness(expression_matrix)
```

Arguments

```
expression_matrix  
  Matrix of expression values
```

Value

Mean absolute skewness across all genes

See Also

Other Section_1_Functions_Use_in_Package: [calculate_cluster_variability\(\)](#), [calculate_expression\(\)](#), [calculate_probability\(\)](#), [compute_adaptive_parameters\(\)](#), [estimate_batch_effect\(\)](#), [extract_dataset_features\(\)](#)

`calculate_probability` *Calculate gene set expression and infer probabilities with control datasets (Use in package)*

Description

Calculate gene set expression and infer probabilities with control datasets (Use in package)

Usage

```
calculate_probability(  
  object,  
  features,  
  assay = NULL,  
  cluster_col = NULL,  
  min_expression = 0.1,  
  specificity_weight = 3  
)
```

Arguments

object	Enter a Seurat object.
features	Enter one or a set of markers.
assay	Enter the assay used by the Seurat object, such as "RNA". Default parameters use "assay = NULL".
cluster_col	Enter the meta.data column in the Seurat object to be annotated, such as "seurat_cluster". Default parameters use "cluster_col = NULL".
min_expression	The min_expression parameter defines a threshold value to determine whether a cell's expression of a feature is considered "expressed" or not. It is used to filter out low-expression cells that may contribute noise to the analysis. Default parameters use "min_expression = 0.1".
specificity_weight	The specificity_weight parameter controls how much the expression variability (standard deviation) of a feature within a cluster contributes to its "specificity score." It amplifies or suppresses the impact of variability in the final score calculation. Default parameters use "specificity_weight = 3".

Value

Average expression of genes in the input "Seurat" object given "cluster_col" and given "features".

See Also

Other Section_1_Functions_Use_in_Package: [calculate_cluster_variability\(\)](#), [calculate_expression\(\)](#), [calculate_expression_skewness\(\)](#), [compute_adaptive_parameters\(\)](#), [estimate_batch_effect\(\)](#), [extract_dataset_features\(\)](#)

Cellmarker2

Cellmarker2 dataset

Description

A dataset containing marker genes for different cell types from Cellmarker2

Usage

```
Cellmarker2
```

Format

A data frame with 8 columns:

Details

This dataset is used to filter and create a standardized marker list. The dataset can be filtered based on species, tissue class, tissue type, cancer type, and cell type to generate a list of marker genes for specific cell types.

Source

<http://117.50.127.228/CellMarker/>

See Also

Other Section_0_Database: [Cellmarker2_raw](#), [Cellmarker2_table](#), [Markers_list_PCTAM](#), [Markers_list_PCTIT](#), [Markers_list_TCellSI](#), [Markers_list_scIBD](#), [PanglaoDB](#), [PanglaoDB_raw](#), [PanglaoDB_table](#), [ScType](#), [ScType_raw](#), [ScType_table](#)

Cellmarker2_raw	<i>Cellmarker2 raw dataset</i>
-----------------	--------------------------------

Description

A dataset containing marker genes for different cell types from Cellmarker2

Usage

Cellmarker2_raw

Format

A data frame with 20 columns contined in the Cellmarker2 database:

Details

This dataset is used to filter and create a standardized marker list. The dataset can be filtered based on species, tissue class, tissue type, cancer type, and cell type to generate a list of marker genes for specific cell types.

Source

<http://117.50.127.228/CellMarker/>

See Also

Other Section_0_Database: [Cellmarker2](#), [Cellmarker2_table](#), [Markers_list_PCTAM](#), [Markers_list_PCTIT](#), [Markers_list_TCellSI](#), [Markers_list_scIBD](#), [PanglaoDB](#), [PanglaoDB_raw](#), [PanglaoDB_table](#), [ScType](#), [ScType_raw](#), [ScType_table](#)

Cellmarker2_table *Cellmarker2 table*

Description

A dataset containing marker genes for different cell types from Cellmarker2

Usage

```
Cellmarker2_table
```

Format

A list contain different types like species, tissue_class, tissue_type, cancer_type, cell_type

Details

This list is used to choose filters for creation of standardized marker list.

Source

<http://117.50.127.228/CellMarker/>

See Also

Other Section_0_Database: [Cellmarker2](#), [Cellmarker2_raw](#), [Markers_list_PCTAM](#), [Markers_list_PCTIT](#), [Markers_list_TCellSI](#), [Markers_list_scIBD](#), [PanglaoDB](#), [PanglaoDB_raw](#), [PanglaoDB_table](#), [ScType](#), [ScType_raw](#), [ScType_table](#)

Celltype_Annotation *Annotate Seurat Object with SlimR Cell Type Predictions*

Description

This function assigns SlimR predicted cell types to a Seurat object based on cluster annotations, and stores the results in the meta.data slot.

Usage

```
Celltype_Annotation(  
  seurat_obj,  
  cluster_col,  
  SlimR_anno_result,  
  plot_UMAP = TRUE,  
  annotation_col = "Cell_type_SlimR"  
)
```

Arguments

seurat_obj	A Seurat object containing cluster information in meta.data.
cluster_col	Character string indicating the column name in meta.data that contains cluster IDs.
SlimR_anno_result	List generated by function Celltype_Calculate() which containing a data.frame in \$Prediction_results with: 1.cluster_col (Cluster identifiers (should match cluster_col in meta.data)) 2.Predicted_cell_type (Predicted cell types for each cluster).
plot_UMAP	logical(1); if TRUE, plot the UMAP with cell type annotations.
annotation_col	The location to write in 'meta.data' that contains the predicted cell type. (default = "Cell_type_SlimR")

Value

A Seurat object with updated meta.data containing the predicted cell types.

Note

If plot_UMAP = TRUE, this function will print a UMAP plot as a side effect.

See Also

Other Section_3_Automated_Annotation: [Celltype_Annotation_PerCell\(\)](#), [Celltype_Calculate\(\)](#), [Celltype_Calculate_PerCell\(\)](#), [Celltype_Verification\(\)](#), [Celltype_Verification_PerCell\(\)](#), [Parameter_Calculate\(\)](#), [percell_workflow](#)

Examples

```
## Not run:
sce <- Celltype_Annotation(seurat_obj = sce,
  cluster_col = "seurat_clusters",
  SlimR_anno_result = SlimR_anno_result,
  plot_UMAP = TRUE,
  annotation_col = "Cell_type_SlimR"
)

## End(Not run)
```

Celltype_annotation_Cellmarker2

Uses "marker_list" from Cellmarker2 for cell annotation

Description

Uses "marker_list" from Cellmarker2 for cell annotation

Usage

```
Celltype_annotation_Cellmarker2(
  seurat_obj,
  gene_list,
  species,
  cluster_col = "seurat_clusters",
  assay = "RNA",
  save_path = NULL,
  min_counts = 1,
  colour_low = "white",
  colour_high = "navy",
  colour_low_mertic = "white",
  colour_high_mertic = "navy"
)
```

Arguments

seurat_obj	Enter the Seurat object with annotation columns such as "seurat_cluster" in meta.data to be annotated.
gene_list	Enter the standard "Marker_list" generated by the Cellmarker2 database for the SlimR package, generated by the "Markers_filter_Cellmarker2 ()" function.
species	This parameter selects the species "Human" or "Mouse" for standard gene format correction of markers entered by "Marker_list".
cluster_col	Enter annotation columns such as "seurat_cluster" in meta.data of the Seurat object to be annotated. Default parameters use "cluster_col = 'seurat_clusters'".
assay	Enter the assay used by the Seurat object, such as "RNA". Default parameters use "assay = "RNA"".
save_path	The output path of the cell annotation picture. Example parameters use "save_path = './SlimR/Celltype_annotation_Cellmarker2/'".
min_counts	The minimum number of counts of genes in "Marker_list" entered. This number represents the number of the same gene in the same species and the same location in the Cellmarker2 database used for annotation of this cell type. Default parameters use "min_counts = 1".
colour_low	Color for lowest expression level. (default = "white")
colour_high	Color for highest expression level. (default = "navy")
colour_low_mertic	Color for lowest mertic level. (default = "white")
colour_high_mertic	Color for highest mertic level. (default = "navy")

Value

The cell annotation picture is saved in "save_path".

See Also

Other Section_5_Other_Functions_Provided: [Celltype_Compare\(\)](#), [Celltype_annotation_Excel\(\)](#), [Celltype_annotation_PanglaoDB\(\)](#), [Celltype_annotation_Seurat\(\)](#), [Compute_Gene_AUC_ROC\(\)](#)

Examples

```
## Not run:
Celltype_annotation_Cellmarker2(seurat_obj = sce,
  gene_list = Markers_list_Cellmarker2,
  species = "Human",
  cluster_col = "seurat_clusters",
  assay = "RNA",
  save_path = file.path(tempdir(), "SlimR_Celltype_annotation_Cellmarker2")
  colour_low = "white",
  colour_high = "navy",
  colour_low_mertic = "white",
  colour_high_mertic = "navy",
)

## End(Not run)
```

Celltype_Annotation_Combined

Uses "marker_list" to generate combined plot for cell annotation

Description

Uses "marker_list" to generate combined plot for cell annotation

Usage

```
Celltype_Annotation_Combined(
  seurat_obj,
  gene_list,
  species,
  cluster_col = "seurat_clusters",
  assay = "RNA",
  save_path = NULL,
  colour_low = "white",
  colour_high = "navy"
)
```

Arguments

seurat_obj Enter the Seurat object with annotation columns such as "seurat_cluster" in meta.data to be annotated.

gene_list	A list of cells and corresponding gene controls, the name of the list is cell type, and the first column of the list corresponds to markers. Lists can be generated using functions such as "Markers_filter_Cellmarker2()", "Markers_filter_PanglaoDB()", "read_excel_markers()", "read_seurat_markers()", etc.
species	This parameter selects the species "Human" or "Mouse" for standard gene format correction of markers entered by "Marker_list".
cluster_col	Enter annotation columns such as "seurat_cluster" in meta.data of the Seurat object to be annotated. Default parameters use "cluster_col = 'seurat_clusters'".
assay	Enter the assay used by the Seurat object, such as "RNA". Default parameters use "assay = 'RNA'".
save_path	The output path of the cell annotation picture. Example parameters use "save_path = './SlimR/Celltype_annotation_Bar/'".
colour_low	Color for lowest expression level. (default = "white")
colour_high	Color for highest expression level. (default = "navy")

Value

The cell annotation picture is saved in "save_path".

See Also

Other Section_4_Semi_Automated_Annotation: [Celltype_Annotation_Features\(\)](#), [Celltype_Annotation_Heatmap\(\)](#)

Examples

```
## Not run:
Celltype_Annotation_Combined(seurat_obj = sce,
  gene_list = Markers_list,
  species = "Human",
  cluster_col = "seurat_clusters",
  assay = "RNA",
  save_path = file.path(tempdir(), "SlimR_Celltype_Annotation_Combined"),
  colour_low = "white",
  colour_high = "navy"
)

## End(Not run)
```

Celltype_annotation_Excel

Uses "marker_list" from Excel input for cell annotation

Description

Uses "marker_list" from Excel input for cell annotation

Usage

```

Celltype_annotation_Excel(
  seurat_obj,
  gene_list,
  species,
  cluster_col = "seurat_clusters",
  assay = "RNA",
  save_path = NULL,
  metric_names = NULL,
  colour_low = "white",
  colour_high = "navy",
  colour_low_mertic = "white",
  colour_high_mertic = "navy"
)

```

Arguments

seurat_obj	Enter the Seurat object with annotation columns such as "seurat_cluster" in meta.data to be annotated.
gene_list	Enter the standard "Marker_list" generated by the Excel files database for the SlimR package, generated by the "read_excel_markers()" function.
species	This parameter selects the species "Human" or "Mouse" for standard gene format correction of markers entered by "Marker_list".
cluster_col	Enter annotation columns such as "seurat_cluster" in meta.data of the Seurat object to be annotated. Default parameters use "cluster_col = "seurat_clusters"".
assay	Enter the assay used by the Seurat object, such as "RNA". Default parameters use "assay = 'RNA'".
save_path	The output path of the cell annotation picture. Example parameters use "save_path = './SlimR/Celltype_annotation_Excel/'".
metric_names	Change the row name for the input mertics, not recommended unless necessary. (NULL is used as default parameter)
colour_low	Color for lowest expression level. (default = "white")
colour_high	Color for highest expression level. (default = "navy")
colour_low_mertic	Color for lowest mertic level. (default = "white")
colour_high_mertic	Color for highest mertic level. (default = "navy")

Value

The cell annotation picture is saved in "save_path".

See Also

Other Section_5_Other_Functions_Provided: [Celltype_Compare\(\)](#), [Celltype_annotation_Cellmarker2\(\)](#), [Celltype_annotation_PanglaoDB\(\)](#), [Celltype_annotation_Seurat\(\)](#), [Compute_Gene_AUC_ROC\(\)](#)

Examples

```
## Not run:
Celltype_annotation_Excel(seurat_obj = sce,
  gene_list = Markers_list_Excel,
  species = "Human",
  cluster_col = "seurat_clusters",
  assay = "RNA",
  save_path = file.path(tempdir(), "SlimR_Celltype_annotation_Excel")
  colour_low = "white",
  colour_high = "navy",
  colour_low_mertic = "white",
  colour_high_mertic = "navy",
)

## End(Not run)
```

Celltype_Annotation_Features

Annotate cell types using features plot with different marker databases

Description

This function dynamically selects the appropriate annotation method based on the `gene_list_type` parameter. It supports marker databases from Cellmarker2, PanglaoDB, Seurat (via `FindAllMarkers`), or Excel files.

Usage

```
Celltype_Annotation_Features(
  seurat_obj,
  gene_list,
  gene_list_type = "Default",
  species = NULL,
  cluster_col = "seurat_clusters",
  assay = "RNA",
  save_path = NULL,
  min_counts = 1,
  metric_names = NULL,
  colour_low = "white",
  colour_high = "navy",
  colour_low_mertic = "white",
  colour_high_mertic = "navy",
  ...
)
```

Arguments

seurat_obj	A valid Seurat object with cluster annotations in meta.data.
gene_list	A list of data frames containing marker genes and metrics. Format depends on gene_list_type: <ul style="list-style-type: none"> • Cellmarker2: Generated by Markers_filter_Cellmarker2(). • PanglaoDB: Generated by Markers_filter_PanglaoDB(). • Seurat: Generated by read_seurat_markers(). • Excel: Generated by read_excel_markers().
gene_list_type	Type of marker database to use. Be one of: "Cellmarker2", "PanglaoDB", "Seurat", or "Excel".
species	Species of the dataset: "Human" or "Mouse" for gene name standardization.
cluster_col	Column name in meta.data defining clusters (default: "seurat_clusters").
assay	Assay layer in the Seurat object (default: "RNA").
save_path	Directory to save output PNGs. Must be explicitly specified.
min_counts	Minimum number of counts for Cellmarker2 annotations (default: 1).
metric_names	Optional. Change the row name for the input metrics, not recommended unless necessary. (NULL is used as default parameter; used in "Seurat"/"Excel").
colour_low	Color for lowest expression level. (default = "white")
colour_high	Color for highest expression level. (default = "navy")
colour_low_mertic	Color for lowest mertic level. (default = "white")
colour_high_mertic	Color for highest mertic level. (default = "navy")
...	Additional parameters passed to the specific annotation function.

Value

Saves cell type annotation PNGs in save_path. Returns invisibly.

See Also

Other Section_4_Semi_Automated_Annotation: [Celltype_Annotation_Combined\(\)](#), [Celltype_Annotation_Heatmap\(\)](#)

Examples

```
## Not run:
# Example for Cellmarker2
Celltype_Annotation_Features(seurat_obj = sce,
  gene_list = Markers_list_Cellmarker2,
  species = "Human",
  cluster_col = "seurat_clusters",
  assay = "RNA",
  save_path = file.path(tempdir(), "SlimR_Celltype_annotation_Cellmarker2"),
  colour_low = "white",
  colour_high = "navy",
```

```
    colour_low_mertic = "white",
    colour_high_mertic = "navy",
  )

# Example for PanglaoDB
Celltype_Annotation_Features(seurat_obj = sce,
  gene_list = Markers_list_panglaoDB,
  species = "Human",
  cluster_col = "seurat_clusters",
  assay = "RNA",
  save_path = file.path(tempdir(), "SlimR_Celltype_annotation_PanglaoDB")
  colour_low = "white",
  colour_high = "navy",
  colour_low_mertic = "white",
  colour_high_mertic = "navy",
)

# Example for Seurat marker list
Celltype_Annotation_Features(seurat_obj = sce,
  gene_list = Markers_list_Seurat,
  species = "Human",
  cluster_col = "seurat_clusters",
  assay = "RNA",
  save_path = file.path(tempdir(), "SlimR_Celltype_annotation_Seurat")
  colour_low = "white",
  colour_high = "navy",
  colour_low_mertic = "white",
  colour_high_mertic = "navy",
)

# Example for Excel marker list
Celltype_Annotation_Features(seurat_obj = sce,
  gene_list = Markers_list_Excel,
  species = "Human",
  cluster_col = "seurat_clusters",
  assay = "RNA",
  save_path = file.path(tempdir(), "SlimR_Celltype_annotation_Excel")
  colour_low = "white",
  colour_high = "navy",
  colour_low_mertic = "white",
  colour_high_mertic = "navy",
)

## End(Not run)
```

Celltype_Annotation_Heatmap

Uses "marker_list" to generate heatmap for cell annotation

Description

Uses "marker_list" to generate heatmap for cell annotation

Usage

```
Celltype_Annotation_Heatmap(
  seurat_obj,
  gene_list,
  species,
  cluster_col = "seurat_clusters",
  assay = "RNA",
  min_expression = 0.1,
  specificity_weight = 3,
  colour_low = "navy",
  colour_high = "firebrick3"
)
```

Arguments

seurat_obj	Enter the Seurat object with annotation columns such as "seurat_cluster" in meta.data to be annotated.
gene_list	A list of cells and corresponding gene controls, the name of the list is cell type, and the first column of the list corresponds to markers. Lists can be generated using functions such as "Markers_filter_Cellmarker2 ()", "Markers_filter_PanglaoDB ()", "read_excel_markers ()", "read_seurat_markers ()", etc.
species	This parameter selects the species "Human" or "Mouse" for standard gene format correction of markers entered by "Marker_list".
cluster_col	Enter annotation columns such as "seurat_cluster" in meta.data of the Seurat object to be annotated. Default parameters use "cluster_col = 'seurat_clusters'".
assay	Enter the assay used by the Seurat object, such as "RNA". Default parameters use "assay = 'RNA'".
min_expression	The min_expression parameter defines a threshold value to determine whether a cell's expression of a feature is considered "expressed" or not. It is used to filter out low-expression cells that may contribute noise to the analysis. Default parameters use "min_expression = 0.1".
specificity_weight	The specificity_weight parameter controls how much the expression variability (standard deviation) of a feature within a cluster contributes to its "specificity score." It amplifies or suppresses the impact of variability in the final score calculation. Default parameters use "specificity_weight = 3".
colour_low	Color for lowest probability level in Heatmap visualization of probability matrix. (default = "navy")
colour_high	Color for highest probability level Heatmap visualization of probability matrix. (default = "firebrick3")

Value

The heatmap of the comparison between "cluster_col" in the Seurat object and the given gene set "gene_list" needs to be annotated.

See Also

Other Section_4_Semi_Automated_Annotation: [Celltype_Annotation_Combined\(\)](#), [Celltype_Annotation_Features\(\)](#)

Examples

```
## Not run:
Celltype_Annotation_Heatmap(seurat_obj = sce,
  gene_list = Markers_list,
  species = "Human",
  cluster_col = "seurat_clusters",
  assay = "RNA",
  min_expression = 0.1,
  specificity_weight = 3,
  colour_low = "navy",
  colour_high = "firebrick3"
)

## End(Not run)
```

Celltype_annotation_PanglaoDB

Uses "marker_list" from PanglaoDB for cell annotation

Description

Uses "marker_list" from PanglaoDB for cell annotation

Usage

```
Celltype_annotation_PanglaoDB(
  seurat_obj,
  gene_list,
  species,
  cluster_col = "seurat_clusters",
  assay = "RNA",
  save_path = NULL,
  metric_names = NULL,
  colour_low = "white",
  colour_high = "navy",
  colour_low_mertic = "white",
  colour_high_mertic = "navy"
)
```

Arguments

seurat_obj	Enter the Seurat object with annotation columns such as "seurat_cluster" in meta.data to be annotated.
gene_list	Enter the standard "Marker_list" generated by the PanglaoDB database for the SlimR package, generated by the "Markers_filter_PanglaoDB ()" function.
species	This parameter selects the species "Human" or "Mouse" for standard gene format correction of markers entered by "Marker_list".
cluster_col	Enter annotation columns such as "seurat_cluster" in meta.data of the Seurat object to be annotated. Default parameters use "cluster_col = 'seurat_clusters'".
assay	Enter the assay used by the Seurat object, such as "RNA". Default parameters use "assay = 'RNA'".
save_path	The output path of the cell annotation picture. Example parameters use "save_path = './SlimR/Celltype_annotation_PanglaoDB/'".
metric_names	Warning: Do not enter information. This parameter is used to check if "Marker_list" conforms to the PanglaoDB database output.
colour_low	Color for lowest expression level. (default = "white")
colour_high	Color for highest expression level. (default = "navy")
colour_low_mertic	Color for lowest mertic level. (default = "white")
colour_high_mertic	Color for highest mertic level. (default = "navy")

Value

The cell annotation picture is saved in "save_path".

See Also

Other Section_5_Other_Functions_Provided: [Celltype_Compare\(\)](#), [Celltype_annotation_Cellmarker2\(\)](#), [Celltype_annotation_Excel\(\)](#), [Celltype_annotation_Seurat\(\)](#), [Compute_Gene_AUC_ROC\(\)](#)

Examples

```
## Not run:
Celltype_annotation_PanglaoDB(seurat_obj = sce,
  gene_list = Markers_list_panglaoDB,
  species = "Human",
  cluster_col = "seurat_clusters",
  assay = "RNA",
  save_path = file.path(tempdir(), "SlimR_Celltype_annotation_PanglaoDB")
  colour_low = "white",
  colour_high = "navy",
  colour_low_mertic = "white",
  colour_high_mertic = "navy",
)

## End(Not run)
```

Celltype_Annotation_PerCell

Annotate Seurat Object with Per-Cell SlimR Predictions

Description

This function assigns SlimR per-cell predicted cell types directly to individual cells in a Seurat object's meta.data slot.

Usage

```
Celltype_Annotation_PerCell(
  seurat_obj,
  SlimR_percell_result,
  plot_UMAP = TRUE,
  annotation_col = "Cell_type_PerCell_SlimR",
  plot_confidence = FALSE
)
```

Arguments

`seurat_obj` A Seurat object.

`SlimR_percell_result` List generated by `Celltype_Calculate_PerCell()` containing `Cell_annotatons` data.frame with `Cell_barcode` and `Predicted_cell_type` columns.

`plot_UMAP` Logical; if TRUE, plot the UMAP with cell type annotations. Default: TRUE.

`annotation_col` Column name to write in meta.data. Default: "Cell_type_PerCell_SlimR".

`plot_confidence` Logical; if TRUE, also plot a UMAP colored by confidence scores. Default: FALSE.

Value

A Seurat object with updated meta.data containing:

- `annotation_col`: Predicted cell type for each cell
- `paste0(annotation_col, "_score")`: Max score for each cell
- `paste0(annotation_col, "_confidence")`: Confidence score for each cell

Note

If `plot_UMAP = TRUE`, this function will print UMAP plot(s) as a side effect.

See Also

Other Section_3_Automated_Annotation: [Celltype_Annotation\(\)](#), [Celltype_Calculate\(\)](#), [Celltype_Calculate_PerCell\(\)](#), [Celltype_Verification\(\)](#), [Celltype_Verification_PerCell\(\)](#), [Parameter_Calculate\(\)](#), [percell_workflow](#)

Examples

```
## Not run:
# Run per-cell annotation
result <- Celltype_Calculate_PerCell(
  seurat_obj = sce,
  gene_list = Markers_list,
  species = "Human"
)

# Annotate Seurat object
sce <- Celltype_Annotation_PerCell(
  seurat_obj = sce,
  SlimR_perCell_result = result,
  plot_UMAP = TRUE,
  annotation_col = "Cell_type_PerCell_SlimR"
)

## End(Not run)
```

Celltype_annotation_Seurat

Uses "marker_list" from Seurat object for cell annotation

Description

Uses "marker_list" from Seurat object for cell annotation

Usage

```
Celltype_annotation_Seurat(
  seurat_obj,
  gene_list,
  species,
  cluster_col = "seurat_clusters",
  assay = "RNA",
  save_path = NULL,
  metric_names = NULL,
  colour_low = "white",
  colour_high = "navy",
  colour_low_mertic = "white",
  colour_high_mertic = "navy"
)
```

Arguments

seurat_obj Enter the Seurat object with annotation columns such as "seurat_cluster" in meta.data to be annotated.

gene_list	Enter the standard "Marker_list" generated by the Seurat object database for the SlimR package, generated by the "read_seurat_markers()" function.
species	This parameter selects the species "Human" or "Mouse" for standard gene format correction of markers entered by "Marker_list".
cluster_col	Enter annotation columns such as "seurat_cluster" in meta.data of the Seurat object to be annotated. Default parameters use "cluster_col = 'seurat_clusters'".
assay	Enter the assay used by the Seurat object, such as "RNA". Default parameters use "assay = 'RNA'".
save_path	The output path of the cell annotation picture. Example parameters use "save_path = './SlimR/Celltype_annotation_Seurat/'".
metric_names	Change the row name for the input metrics, not recommended unless necessary. (NULL is used as default parameter)
colour_low	Color for lowest expression level. (default = "white")
colour_high	Color for highest expression level. (default = "navy")
colour_low_mertic	Color for lowest mertic level. (default = "white")
colour_high_mertic	Color for highest mertic level. (default = "navy")

Value

The cell annotation picture is saved in "save_path".

See Also

Other Section_5_Other_Functions_Provided: [Celltype_Compare\(\)](#), [Celltype_annotation_Cellmarker2\(\)](#), [Celltype_annotation_Excel\(\)](#), [Celltype_annotation_PanglaoDB\(\)](#), [Compute_Gene_AUC_ROC\(\)](#)

Examples

```
## Not run:
Celltype_annotation_Seurat(seurat_obj = sce,
  gene_list = Markers_list_Seurat,
  species = "Human",
  cluster_col = "seurat_clusters",
  assay = "RNA",
  save_path = file.path(tempdir(), "SlimR_Celltype_annotation_Seurat")
  colour_low = "white",
  colour_high = "navy",
  colour_low_mertic = "white",
  colour_high_mertic = "navy",
)

## End(Not run)
```

Celltype_Calculate	<i>Uses "marker_list" to calculate probability, prediction results, AUC and generate heatmap for cell annotation</i>
--------------------	--

Description

Uses "marker_list" to calculate probability, prediction results, AUC and generate heatmap for cell annotation

Usage

```
Celltype_Calculate(
  seurat_obj,
  gene_list,
  species,
  cluster_col = "seurat_clusters",
  assay = "RNA",
  min_expression = 0.1,
  specificity_weight = 3,
  threshold = 0.6,
  compute_AUC = TRUE,
  plot_AUC = TRUE,
  AUC_correction = FALSE,
  colour_low = "navy",
  colour_high = "firebrick3"
)
```

Arguments

seurat_obj	Enter the Seurat object with annotation columns such as "seurat_cluster" in meta.data to be annotated.
gene_list	A list of cells and corresponding gene controls, the name of the list is cell type, and the first column of the list corresponds to markers. Lists can be generated using functions such as "Markers_filter_Cellmarker2 ()", "Markers_filter_PanglaoDB ()", "read_excel_markers ()", "read_seurat_markers ()", etc.
species	This parameter selects the species "Human" or "Mouse" for standard gene format correction of markers entered by "Marker_list".
cluster_col	Enter annotation columns such as "seurat_cluster" in meta.data of the Seurat object to be annotated. Default parameters use "cluster_col = 'seurat_clusters'".
assay	Enter the assay used by the Seurat object, such as "RNA". Default parameters use "assay = 'RNA'".
min_expression	The min_expression parameter defines a threshold value to determine whether a cell's expression of a feature is considered "expressed" or not. It is used to filter out low-expression cells that may contribute noise to the analysis. Default parameters use "min_expression = 0.1".

specificity_weight	The specificity_weight parameter controls how much the expression variability (standard deviation) of a feature within a cluster contributes to its "specificity score." It amplifies or suppresses the impact of variability in the final score calculation. Default parameters use "specificity_weight = 3".
threshold	This parameter refers to the normalized similarity between the "alternative cell type" and the "predicted cell type" in the returned results. (the default parameter is 0.6)
compute_AUC	Logical indicating whether to calculate AUC values for predicted cell types. AUC measures how well the marker genes distinguish the cluster from others. When TRUE, adds an AUC column to the prediction results. (default: TRUE)
plot_AUC	The logic indicates whether to draw an AUC curve for the predicted cell type. When TRUE, add an AUC_plot to result. (default: TRUE)
AUC_correction	Logical value controlling AUC-based correction. (default = FALSE) When set to TRUE: 1.Computes AUC values for candidate cell types. (probability > threshold) 2.Selects the cell type with the highest AUC as the final predicted type. 3.Records the selected type's AUC value in the "AUC" column.
colour_low	Color for lowest probability level in Heatmap visualization of probability matrix. (default = "navy")
colour_high	Color for highest probability level Heatmap visualization of probability matrix. (default = "firebrick3")

Value

A list containing:

- Expression_list: List of expression matrices for each cell type
- Proportion_list: List of proportion of expression for each cell type
- Expression_scores_matrix: Matrix of expression scores
- Probability_matrix: Matrix of normalized probabilities
- Prediction_results: Data frame with cluster annotations including:
 - cluster_col: Cluster identifier
 - Predicted_cell_type: Primary predicted cell type
 - AUC: Area Under the Curve value (when compute_AUC = TRUE)
 - Alternative_cell_types: Semi-colon separated alternative cell types
- Heatmap_plot: Heatmap visualization of probability matrix (pheatmap object). Can be displayed using print() or plot()
- AUC_plot: AUC visualization of Predicted cell type (ggplot object)
- AUC_list: The resulting list of AUC values calculated for genes in alternative cell types above the approximate threshold

See Also

Other Section_3_Automated_Annotation: [Celltype_Annotation\(\)](#), [Celltype_Annotation_PerCell\(\)](#), [Celltype_Calculate_PerCell\(\)](#), [Celltype_Verification\(\)](#), [Celltype_Verification_PerCell\(\)](#), [Parameter_Calculate\(\)](#), [percell_workflow](#)

Examples

```
## Not run:
SlimR_anno_result <- Celltype_Calculate(seurat_obj = sce,
  gene_list = Markers_list,
  species = "Human",
  cluster_col = "seurat_clusters",
  assay = "RNA",
  min_expression = 0.1,
  specificity_weight = 3,
  threshold = 0.6,
  compute_AUC = TRUE,
  plot_AUC = TRUE,
  AUC_correction = FALSE,
  colour_low = "navy",
  colour_high = "firebrick3"
)

## End(Not run)
```

Celltype_Calculate_PerCell

Per-cell annotation using marker expression and optional UMAP spatial smoothing

Description

Unlike cluster-based annotation, this function assigns cell type labels to each individual cell based on marker gene expression profiles. Optionally uses UMAP coordinates to smooth predictions via k-nearest neighbor voting.

Usage

```
Celltype_Calculate_PerCell(
  seurat_obj,
  gene_list,
  species,
  assay = "RNA",
  method = c("weighted", "mean", "AUCell"),
  min_expression = 0.1,
  use_umap_smoothing = FALSE,
  umap_reduction = "umap",
  k_neighbors = 15,
  smoothing_weight = 0.3,
  min_score = "auto",
  min_confidence = 1.2,
  return_scores = FALSE,
  ncores = 1,
```

```

    chunk_size = 5000,
    verbose = TRUE
)

```

Arguments

seurat_obj	Seurat object with normalized expression data.
gene_list	A standardized marker list (same format as Celltype_Calculate).
species	"Human" or "Mouse" for gene name formatting.
assay	Assay to use (default: "RNA").
method	Scoring method: "AUCell" (rank-based), "mean" (average expression), or "weighted" (expression * detection weighted). Default: "weighted".
min_expression	Minimum expression threshold for detection. Default: 0.1.
use_umap_smoothing	Logical. If TRUE, apply k-NN smoothing using UMAP coordinates to improve annotation consistency. Default: FALSE.
umap_reduction	Name of UMAP reduction in Seurat object. Default: "umap".
k_neighbors	Number of neighbors for UMAP smoothing. Default: 15.
smoothing_weight	Weight for neighbor votes vs cell's own score (0-1). Higher values give more weight to neighbors. Default: 0.3.
min_score	Minimum score threshold to assign a cell type. Cells below this threshold are labeled "Unassigned". Default: "auto" which adaptively sets the threshold based on number of cell types (1.5 / n_celltypes). Set to a numeric value (e.g., 0.1) to use a fixed threshold.
min_confidence	Minimum confidence threshold. Cells with confidence below this value are labeled "Unassigned". Confidence is calculated as the ratio of max score to second-highest score. Default: 1.2 (max must be 20% higher than second). Set to 1.0 to disable confidence filtering.
return_scores	If TRUE, return full score matrix. Default: FALSE.
ncores	Number of cores for parallel processing. Default: 1.
chunk_size	Number of cells to process per chunk (memory optimization). Default: 5000.
verbose	Print progress messages. Default: TRUE.

Details

Scoring Methods:

"weighted" (recommended): Combines normalized expression with detection rate. For each cell and cell type: $\text{score} = \text{mean}(\text{expr}_i * \text{weight}_i)$ where weight_i is derived from the marker's specificity across the dataset.

"mean": Simple average of normalized marker expression. Fast but less discriminative for overlapping marker sets.

"AUCell": Rank-based scoring similar to AUCell package. For each cell, genes are ranked by expression, and the score is the proportion of marker genes in the top X% of expressed genes. Robust to technical variation.

UMAP Smoothing:

When `use_umap_smoothing = TRUE`, the function:

1. Computes initial per-cell scores
2. Finds k nearest neighbors in UMAP space for each cell
3. Smooths scores by weighted averaging with neighbors
4. Re-assigns cell types based on smoothed scores

This helps reduce noise and improve consistency of annotations within spatially coherent regions.

Value

A list containing:

- `Cell_annotatons`: Data frame with `Cell_barcode`, `Predicted_cell_type`, `Max_score`, `Confidence`
- `Cell_confidence`: Numeric vector of confidence scores per cell
- `Summary`: Summary table of cell type counts and percentages
- `Expression_list`: List of mean expression matrices per cell type (for verification)
- `Proportion_list`: List of detection proportion matrices per cell type
- `Prediction_results`: Summary data frame with per-cell-type statistics
- `Probability_matrix`: Full cell \times cell_type probability matrix (normalized)
- `Raw_score_matrix`: Full cell \times cell_type raw score matrix (before normalization)
- `Parameters`: List of parameters used including adaptive thresholds
- `Cell_scores`: (if `return_scores=TRUE`) Same as `Probability_matrix`

See Also

Other Section_3_Automated_Annotation: [Celltype_Annotation\(\)](#), [Celltype_Annotation_PerCell\(\)](#), [Celltype_Calculate\(\)](#), [Celltype_Verification\(\)](#), [Celltype_Verification_PerCell\(\)](#), [Parameter_Calculate\(\)](#), [percell_workflow](#)

Examples

```
## Not run:
# Basic per-cell annotation
result <- Celltype_Calculate_PerCell(
  seurat_obj = sce,
  gene_list = Markers_list,
  species = "Human",
  method = "weighted"
)

# Add annotations to Seurat object
sce$Cell_type_PerCell <- result$Cell_annotatons$Predicted_cell_type

# With UMAP smoothing for more consistent annotations
result_smooth <- Celltype_Calculate_PerCell(
  seurat_obj = sce,
```

```

gene_list = Markers_list,
species = "Human",
use_umap_smoothing = TRUE,
k_neighbors = 20,
smoothing_weight = 0.3
)

## End(Not run)

```

Celltype_Compare	<i>Compare cell type labels across two single-cell datasets after aligning cell barcodes</i>
------------------	--

Description

This function automatically aligns cell barcodes between two Seurat objects using a variety of normalization transformations, then cross-tabulates a cell type label column (from the first object) against a grouping column (from the second object). It returns count tables, proportion tables, a dominant mapping, and a heatmap.

Usage

```

Celltype_Compare(
  sce_label,
  sce,
  label_col = NULL,
  group_col = NULL,
  barcode_col = NULL,
  color_low = "grey70",
  color_high = "navy",
  show_plot = TRUE
)

```

Arguments

sce_label	A Seurat object containing the cell type label column.
sce	A Seurat object containing the grouping column.
label_col	Character. Name of the metadata column in sce_label that stores cell type labels (e.g., "Sub_cell_type").
group_col	Character. Name of the metadata column in sce that stores grouping information (e.g., "SCT_snn_res.0.3").
barcode_col	Optional character. Name of a metadata column in both objects that contains the cell barcode identifiers. If NULL, the function uses colnames(sce_label) and colnames(sce).
color_low	Character. Color for low proportion values in the heatmap. Default: "grey70".

color_high	Character. Color for high proportion values in the heatmap. Default: "navy".
show_plot	Logical. If TRUE (default), the heatmap is automatically displayed when the function is called. Set to FALSE to suppress automatic plotting (e.g., in non-interactive environments).

Details

Cell barcode alignment: The function automatically tries a set of normalization functions on the cell identifiers (either from `barcode_col` or from column names) to maximise the number of shared barcodes between the two objects. Transformations include: `identity`, `drop_numeric_suffix` (removes e.g., "-1-2"), `drop_suffix` (removes "-1"), and several prefix removals. The transformation pair yielding the highest number of shared identifiers is selected.

Proportion calculation: Proportions are computed **within each** `group_col` level (column-wise), i.e. for each group, the sum of proportions across all cell types equals 1.

Plot: The heatmap uses `ggplot2::geom_tile()` with a fixed coordinate ratio and a colour gradient from `color_low` to `color_high`.

Value

A list with five components:

<code>count_table</code>	A data frame (wide format) with rows = unique <code>label_col</code> values and columns = unique <code>group_col</code> values; cell values are raw counts of shared cells.
<code>prop_table</code>	Same shape as <code>count_table</code> ; each cell shows the proportion of cells within a <code>group_col</code> column (column-wise sum = 1).
<code>main_to_sub</code>	A data frame mapping each <code>group_col</code> value to the most frequent <code>label_col</code> value among shared cells.
<code>plot</code>	A <code>ggplot2</code> heatmap object visualizing the proportion table.
<code>match_info</code>	A tibble with columns <code>label_transform</code> , <code>sce_transform</code> , <code>shared_n</code> – the transformations used to align barcodes and the number of shared cells after alignment.

See Also

Other Section_5_Other_Functions_Provided: [Celltype_annotation_Cellmarker2\(\)](#), [Celltype_annotation_Excel\(\)](#), [Celltype_annotation_PanglaoDB\(\)](#), [Celltype_annotation_Seurat\(\)](#), [Compute_Gene_AUC_ROC\(\)](#)

Examples

```
## Not run:
# Basic usage with two Seurat objects and default barcode alignment
result <- Celltype_Compare(
  sce_label = seurat_obj1,
  sce = seurat_obj2,
  label_col = "cell_type",
  group_col = "cluster"
)
```

```

# Access the proportion table
head(result$prop_table)

# View the dominant mapping
print(result$main_to_sub)

# Display the heatmap
print(result$plot)

# Use a custom barcode column
result2 <- Celltype_Compare(
  sce_label = seurat_obj1,
  sce = seurat_obj2,
  label_col = "cell_type",
  group_col = "cluster",
  barcode_col = "cell_barcode"
)

## End(Not run)

```

Celltype_Verification *Perform cell type verification and generate the validation dotplot*

Description

This function performs verification of predicted cell types by selecting high log2FC and high expression proportion genes and generates and generate the validation dotplot.

Usage

```

Celltype_Verification(
  seurat_obj,
  SlimR_anno_result,
  assay = "RNA",
  gene_number = 5,
  colour_low = "white",
  colour_high = "navy",
  annotation_col = "Cell_type_SlimR"
)

```

Arguments

`seurat_obj` A Seurat object containing single-cell data.

`SlimR_anno_result` A list containing SlimR annotation results with: `Expression_list` - List of expression matrices for each cell type. `Prediction_results` - Data frame with cluster annotations.

assay	Enter the assay used by the Seurat object, such as "RNA". Default parameters use "assay = 'RNA'".
gene_number	Integer specifying number of top genes to select per cell type.
colour_low	Color for lowest expression level. (default = "white")
colour_high	Color for highest expression level. (default = "navy")
annotation_col	Character string specifying the column in meta.data to use for grouping.

Value

A ggplot object showing expression of top variable genes.

See Also

Other Section_3_Automated_Annotation: [Celltype_Annotation\(\)](#), [Celltype_Annotation_PerCell\(\)](#), [Celltype_Calculate\(\)](#), [Celltype_Calculate_PerCell\(\)](#), [Celltype_Verification_PerCell\(\)](#), [Parameter_Calculate\(\)](#), [percell_workflow](#)

Examples

```
## Not run:
Celltype_Verification(seurat_obj = sce,
  SlimR_anno_result = SlimR_anno_result,
  assay = "RNA",
  gene_number = 5,
  colour_low = "white",
  colour_high = "navy",
  annotation_col = "Cell_type_SlimR"
)

## End(Not run)
```

Celltype_Verification_PerCell

Verify per-cell annotations with marker expression dotplot

Description

This function verifies per-cell SlimR annotations by generating a dotplot showing marker gene expression across predicted cell types.

Usage

```
Celltype_Verification_PerCell(
  seurat_obj,
  SlimR_percell_result,
  assay = "RNA",
```

```

gene_number = 5,
colour_low = "white",
colour_high = "navy",
annotation_col = "Cell_type_PerCell_SlimR",
min_cells = 10
)

```

Arguments

seurat_obj	A Seurat object with per-cell annotations.
SlimR_percell_result	A list from Celltype_Calculate_PerCell() containing Expression_list with marker genes per cell type.
assay	Assay to use. Default: "RNA".
gene_number	Number of top genes to show per cell type. Default: 5.
colour_low	Color for lowest expression. Default: "white".
colour_high	Color for highest expression. Default: "navy".
annotation_col	Column in meta.data with cell type annotations. Default: "Cell_type_PerCell_SlimR".
min_cells	Minimum number of cells required for a cell type to be included in the plot. Default: 10.

Value

A ggplot object showing marker gene expression dotplot.

See Also

Other Section_3_Automated_Annotation: [Celltype_Annotation\(\)](#), [Celltype_Annotation_PerCell\(\)](#), [Celltype_Calculate\(\)](#), [Celltype_Calculate_PerCell\(\)](#), [Celltype_Verification\(\)](#), [Parameter_Calculate\(\)](#), [percell_workflow](#)

Examples

```

## Not run:
# After running Celltype_Calculate_PerCell and Celltype_Annotation_PerCell
dotplot <- Celltype_Verification_PerCell(
  seurat_obj = sce,
  SlimR_percell_result = result,
  gene_number = 5,
  annotation_col = "Cell_type_PerCell_SlimR"
)
print(dotplot)

## End(Not run)

```

`compute_adaptive_parameters`*Compute Adaptive Parameters Based on Dataset Features (Use in package)*

Description

Calculates optimal `min_expression`, `specificity_weight`, and `threshold` parameters using continuous adaptive algorithms based on dataset characteristics.

Usage

```
compute_adaptive_parameters(dataset_features, n_celltypes = 50)
```

Arguments

`dataset_features` List of dataset characteristics from `extract_dataset_features()`

`n_celltypes` Expected number of cell types in marker database

Value

List containing `min_expression`, `specificity_weight`, `threshold`, and `rationale`

See Also

Other Section_1_Functions_Use_in_Package: [calculate_cluster_variability\(\)](#), [calculate_expression\(\)](#), [calculate_expression_skewness\(\)](#), [calculate_probability\(\)](#), [estimate_batch_effect\(\)](#), [extract_dataset_features\(\)](#)

`Compute_Gene_AUC_ROC` *Compute AUC and Optionally Plot ROC Curve for a Single Gene*

Description

Evaluates the discriminatory power of a single gene in separating a user-defined positive cell group from the rest, using the Area Under the Receiver Operating Characteristic curve (AUC). Several scoring strategies and optional cell subsetting are provided to handle dropout noise and sparse expression typical of single-cell data. If `plot = TRUE` (the default), a publication-ready ROC curve is automatically displayed via **ggplot2**.

Usage

```

Compute_Gene_AUC_ROC(
  seurat_obj,
  gene,
  group_col,
  group_label,
  assay = "RNA",
  method = c("raw", "rank"),
  min_expression = NULL,
  keep_expression_above = NULL,
  plot = TRUE,
  plot_title = "ROC Curve | SlimR",
  line_color = "navy",
  line_size = 1
)

```

Arguments

seurat_obj	A Seurat object containing single-cell expression data.
gene	A single character string specifying the gene to evaluate. Must be present in the rownames of the chosen assay.
group_col	Character string giving the name of a column in <code>seurat_obj@meta.data</code> that defines cell groups (e.g., "seurat_clusters").
group_label	The value in <code>group_col</code> that defines the positive class (e.g., 1 for cluster 1). All other cells are treated as negatives.
assay	Character string specifying which assay to use. Default is "RNA".
method	Scoring method: "raw" (raw expression, optionally truncated by <code>min_expression</code>) or "rank" (expression ranks, which is robust to dropout and does not require <code>min_expression</code>).
min_expression	Numeric threshold for expression truncation (only used when <code>method = "raw"</code>). Values below this threshold are set to zero. Default is NULL (no truncation).
keep_expression_above	Optional numeric threshold. If set, only cells with expression greater than this value are retained for AUC computation. Note: this changes the population over which AUC is evaluated, shifting the interpretation from "overall discrimination among all cells" to "discrimination among cells that express the gene above this level". Use with caution and always compare with the result without subsetting. Default is NULL (all cells retained).
plot	Logical indicating whether to create and automatically display a ggplot2 ROC curve. Default is TRUE.
plot_title	Character string used as the title of the ROC plot. Default is "ROC Curve SlimR".
line_color	Colour of the ROC curve. Default is "navy".
line_size	Numeric value for the thickness of the ROC curve line. Default is 1.

Value

Invisibly, a list with the following elements:

AUC Numeric value (between 0 and 1) of the area under the ROC curve.

roc_data A data.frame with columns fpr (False Positive Rate) and tpr (True Positive Rate) that can be used for custom plotting.

predictions Numeric vector of the (possibly truncated) expression values used as prediction scores.

labels Logical vector indicating whether each cell belongs to the positive class (TRUE) or not (FALSE).

roc_plot If plot = TRUE, a **ggplot** object displaying the ROC curve; otherwise NULL.

See Also

Other Section_5_Other_Functions_Provided: [Celltype_Compare\(\)](#), [Celltype_annotation_Cellmarker2\(\)](#), [Celltype_annotation_Excel\(\)](#), [Celltype_annotation_PanglaoDB\(\)](#), [Celltype_annotation_Seurat\(\)](#)

Examples

```
## Not run:
# Default: raw expression, plot displayed automatically
res <- Compute_Gene_AUC_ROC(sce, "BMX", "seurat_clusters", 12)

# Rank-based (robust to dropout)
res_rank <- Compute_Gene_AUC_ROC(sce, "BMX", "seurat_clusters", 12,
                                method = "rank")

# Keep only cells expressing above 0.5 (exploratory)
res_sub <- Compute_Gene_AUC_ROC(sce, "BMX", "seurat_clusters", 12,
                                keep_expression_above = 0.5)

## End(Not run)
```

estimate_batch_effect *Estimate Batch Effect Strength (Use in package)*

Description

Roughly estimates the potential impact of batch effects using available metadata.

Usage

```
estimate_batch_effect(seurat_obj, assay)
```

Arguments

seurat_obj	Seurat object
assay	Assay name

Value

Batch effect score (0 indicates no detectable batch effect)

See Also

Other Section_1_Functions_Use_in_Package: [calculate_cluster_variability\(\)](#), [calculate_expression\(\)](#), [calculate_expression_skewness\(\)](#), [calculate_probability\(\)](#), [compute_adaptive_parameters\(\)](#), [extract_dataset_features\(\)](#)

extract_dataset_features

*Extract Dataset Characteristics for Adaptive Parameter Calculation
(Use in package)*

Description

Computes various statistical features from single-cell data that are used as input for the parameter prediction model.

Usage

```
extract_dataset_features(  
  seurat_obj,  
  features,  
  assay = NULL,  
  cluster_col = NULL  
)
```

Arguments

seurat_obj	Seurat object
features	Features to analyze
assay	Assay name
cluster_col	Cluster column name

Value

List of dataset characteristics including expression statistics, variability measures, and cluster properties

See Also

Other Section_1_Functions_Use_in_Package: [calculate_cluster_variability\(\)](#), [calculate_expression\(\)](#), [calculate_expression_skewness\(\)](#), [calculate_probability\(\)](#), [compute_adaptive_parameters\(\)](#), [estimate_batch_effect\(\)](#)

`Markers_filter_Cellmarker2`*Create Marker_list from the Cellmarkers2 database*

Description

Create Marker_list from the Cellmarkers2 database

Usage

```
Markers_filter_Cellmarker2(  
  df,  
  species = NULL,  
  tissue_class = NULL,  
  tissue_type = NULL,  
  cancer_type = NULL,  
  cell_type = NULL  
)
```

Arguments

<code>df</code>	Standardized Cellmarkers2 database. It is read as data(Cellmarkers2) in the SlimR library.
<code>species</code>	Species information in Cellmarkers2 database. The default input is "Human" or "Mouse".The input can be retrieved by "Cellmarkers2_table". For more information,please refer to http://117.50.127.228/CellMarker/ on Cellmarkers2's official website.
<code>tissue_class</code>	Tissue_class information in Cellmarkers2 database. The input can be retrieved by "Cellmarkers2_table". For more information, please refer to http://117.50.127.228/CellMarker/ on Cellmarkers2's official website.
<code>tissue_type</code>	Tissue_type information in Cellmarkers2 database. The input can be retrieved by "Cellmarkers2_table". For more information, please refer to http://117.50.127.228/CellMarker/ on Cellmarkers2's official website.
<code>cancer_type</code>	Cancer_type information in Cellmarkers2 database. The input can be retrieved by "Cellmarkers2_table". For more information, please refer to http://117.50.127.228/CellMarker/ on Cellmarkers2's official website.
<code>cell_type</code>	Cell_type information in Cellmarkers2 database. The input can be retrieved by "Cellmarkers2_table". For more information, please refer to http://117.50.127.228/CellMarker/ on Cellmarkers2's official website.

Value

The standardized "Marker_list" in the SlimR package

See Also

Other Section_2_Standardized_Markers_List: [Markers_filter_PanglaoDB\(\)](#), [Markers_filter_ScType\(\)](#), [Read_excel_markers\(\)](#), [Read_seurat_markers\(\)](#)

Examples

```
Cellmarker2 <- SlimR::Cellmarker2
Markers_list_Cellmarker2 <- Markers_filter_Cellmarker2(
  Cellmarker2,
  species = "Human",
  tissue_class = "Intestine",
  tissue_type = NULL,
  cancer_type = NULL,
  cell_type = NULL
)
```

Markers_filter_PanglaoDB

Create Marker_list from the PanglaoDB database

Description

Create Marker_list from the PanglaoDB database

Usage

```
Markers_filter_PanglaoDB(df, species_input, organ_input)
```

Arguments

df	Standardized PanglaoDB database. It is read as data(PanglaoDB) in the SlimR library.
species_input	Species information in PanglaoDB database. The default input is "Human" or "Mouse".The input can be retrieved by "PanglaoDB_table". For more information, please refer to https://panglaodb.se/ on PanglaoDB's official website.
organ_input	Organ type information in the PanglaoDB database. The input can be retrieved by "PanglaoDB_table".For more information, please refer to https://panglaodb.se/ on PanglaoDB's official website.

Value

The standardized "Marker_list" in the SlimR package

See Also

Other Section_2_Standardized_Markers_List: [Markers_filter_Cellmarker2\(\)](#), [Markers_filter_ScType\(\)](#), [Read_excel_markers\(\)](#), [Read_seurat_markers\(\)](#)

Examples

```
PanglaoDB <- SlimR::PanglaoDB
Markers_list_panglaoDB <- Markers_filter_PanglaoDB(
  PanglaoDB,
  species_input = 'Human',
  organ_input = 'GI tract'
)
```

Markers_filter_ScType *Create Marker_list from the ScType database*

Description

Create Marker_list from the ScType database

Usage

```
Markers_filter_ScType(df, tissue_type = NULL, cell_name = NULL)
```

Arguments

df	Standardized ScType database. It is read as data(ScType) in the SlimR library.
tissue_type	Tissue type information in the ScType database. The input can be retrieved by "ScType_table". For more information, please refer to https://github.com/IanevskiAleksandr/sc-type .
cell_name	Cell type name information in the ScType database. The input can be retrieved by "ScType_table". For more information, please refer to https://github.com/IanevskiAleksandr/sc-type .

Value

The standardized "Marker_list" in the SlimR package

See Also

Other Section_2_Standardized_Markers_List: [Markers_filter_Cellmarker2\(\)](#), [Markers_filter_PanglaoDB\(\)](#), [Read_excel_markers\(\)](#), [Read_seurat_markers\(\)](#)

Examples

```
ScType <- SlimR::ScType
Markers_list_ScType <- Markers_filter_ScType(
  ScType,
  tissue_type = "Immune system",
  cell_name = NULL
)
```

Markers_list_PCTAM *List of Macrophage subtype markers in the article "Macrophage diversity in cancer revisited in the era of single-cell omics"*

Description

A dataset containing marker genes for different Macrophage subtypes from the article "Macrophage diversity in cancer revisited in the era of single-cell omics"

Usage

Markers_list_PCTAM

Format

A list with 7 tables.

Details

This list is a table of 7 types of Tumor-associated macrophages (TAMs) markers obtained from the article "Macrophage diversity in cancer revisited in the era of single-cell omics". The data source is "<https://doi.org/10.1016/j.it.2022.04.008>", and the reference literature is: Ruo-Yu Ma et al. (2022) [doi:10.1016/j.it.2022.04.008](https://doi.org/10.1016/j.it.2022.04.008).

Source

[doi:10.1016/j.it.2022.04.008](https://doi.org/10.1016/j.it.2022.04.008)

See Also

Other Section_0_Database: [Cellmarker2](#), [Cellmarker2_raw](#), [Cellmarker2_table](#), [Markers_list_PCTIT](#), [Markers_list_TCellSI](#), [Markers_list_scIBD](#), [PanglaoDB](#), [PanglaoDB_raw](#), [PanglaoDB_table](#), [ScType](#), [ScType_raw](#), [ScType_table](#)

Markers_list_PCTIT *List of T cell subtype markers in the article "Pan-cancer single cell landscape of tumor-infiltrating T cells"*

Description

A dataset containing marker genes for different T cell types from the article "Pan-cancer single cell landscape of tumor-infiltrating T cells"

Usage

Markers_list_PCTIT

Format

A list with 40 tables.

Details

This list is a table of 40 types of pan-cancer tumor-infiltrating T cell (PCTIT) markers obtained from the article "Pan-cancer single cell landscape of tumor-infiltrating T cells". The data source is "<https://doi.org/10.1126/science.abe6474>", and the reference literature is: L. Zheng et al. (2021) [doi:10.1126/science.abe6474](https://doi.org/10.1126/science.abe6474).

Source

[doi:10.1126/science.abe6474](https://doi.org/10.1126/science.abe6474)

See Also

Other Section_0_Database: [Cellmarker2](#), [Cellmarker2_raw](#), [Cellmarker2_table](#), [Markers_list_PCTAM](#), [Markers_list_TCellSI](#), [Markers_list_scIBD](#), [PanglaoDB](#), [PanglaoDB_raw](#), [PanglaoDB_table](#), [ScType](#), [ScType_raw](#), [ScType_table](#)

Markers_list_scIBD *List of cell type markers in the article scIBD*

Description

A dataset containing marker genes for different human intestine cell types from scIBD

Usage

Markers_list_scIBD

Format

A list with one hundred and one tables.

Details

This list is a table of 101 types of human intestine cell types markers obtained from scIBD. The article doi source is "<https://doi.org/10.1038/s43588-023-00464-9>", and the reference literature is: Nie et al. (2023) [doi:10.1038/s43588-023-00464-9](https://doi.org/10.1038/s43588-023-00464-9). Note: The 'Markers_list_scIBD' was generated using section 2.5.2 and the parameters 'sort_by = "logFC"' and 'gene_filter = 20' were set.

Source

[doi:10.1038/s43588023004649](https://doi.org/10.1038/s43588023004649)

See Also

Other Section_0_Database: [Cellmarker2](#), [Cellmarker2_raw](#), [Cellmarker2_table](#), [Markers_list_PCTAM](#), [Markers_list_PCTIT](#), [Markers_list_TCellSI](#), [PanglaoDB](#), [PanglaoDB_raw](#), [PanglaoDB_table](#), [ScType](#), [ScType_raw](#), [ScType_table](#)

Markers_list_TCellSI *List of T cell subtype markers in the article TCellSI*

Description

A dataset containing marker genes for different T cell subtypes from TCellSI

Usage

Markers_list_TCellSI

Format

A list with ten tables.

Details

This list is a table of 10 types of T cell markers obtained from TCellSI. The data source is "<https://github.com/GuoBioinfoLab/> and the reference literature is: Yang et al. (2024) [doi:10.1002/imt2.231](https://doi.org/10.1002/imt2.231).

Source

<https://github.com/GuoBioinfoLab/TCellSI/>

See Also

Other Section_0_Database: [Cellmarker2](#), [Cellmarker2_raw](#), [Cellmarker2_table](#), [Markers_list_PCTAM](#), [Markers_list_PCTIT](#), [Markers_list_scIBD](#), [PanglaoDB](#), [PanglaoDB_raw](#), [PanglaoDB_table](#), [ScType](#), [ScType_raw](#), [ScType_table](#)

PanglaoDB	<i>PanglaoDB dataset</i>
-----------	--------------------------

Description

A dataset containing marker genes for different cell types from PanglaoDB

Usage

PanglaoDB

Format

A data frame with 9 columns:

Details

This dataset is used to filter and create a standardized marker list.'

Source

<https://panglaodb.se/>

See Also

Other Section_0_Database: [Cellmarker2](#), [Cellmarker2_raw](#), [Cellmarker2_table](#), [Markers_list_PCTAM](#), [Markers_list_PCTIT](#), [Markers_list_TCellSI](#), [Markers_list_scIBD](#), [PanglaoDB_raw](#), [PanglaoDB_table](#), [ScType](#), [ScType_raw](#), [ScType_table](#)

PanglaoDB_raw	<i>PanglaoDB raw dataset</i>
---------------	------------------------------

Description

A dataset containing marker genes for different cell types from PanglaoDB

Usage

PanglaoDB_raw

Format

A data frame with 14 columns contined in the PanglaoDB database:

Details

This dataset is used to filter and create a standardized marker list.'

Source

<https://panglaodb.se/>

See Also

Other Section_0_Database: [Cellmarker2](#), [Cellmarker2_raw](#), [Cellmarker2_table](#), [Markers_list_PCTAM](#), [Markers_list_PCTIT](#), [Markers_list_TCellSI](#), [Markers_list_scIBD](#), [PanglaoDB](#), [PanglaoDB_table](#), [ScType](#), [ScType_raw](#), [ScType_table](#)

PanglaoDB_table	<i>PanglaoDB table</i>
-----------------	------------------------

Description

A dataset containing marker genes for different cell types from PanglaoDB

Usage

PanglaoDB_table

Format

A list contain different types like species, organ, cell type.

Details

This list is used to choose filters for creation of standardized marker list.

Source

<https://panglaodb.se/>

See Also

Other Section_0_Database: [Cellmarker2](#), [Cellmarker2_raw](#), [Cellmarker2_table](#), [Markers_list_PCTAM](#), [Markers_list_PCTIT](#), [Markers_list_TCellSI](#), [Markers_list_scIBD](#), [PanglaoDB](#), [PanglaoDB_raw](#), [ScType](#), [ScType_raw](#), [ScType_table](#)

Parameter_Calculate *Adaptive Parameter Tuning for Single-Cell Data Annotation in SlimR*

Description

This function automatically determines optimal `min_expression`, `specificity_weight`, and `threshold` parameters for single-cell data analysis based on dataset characteristics using adaptive algorithms derived from empirical analysis of single-cell datasets.

Usage

```
Parameter_Calculate(  
  seurat_obj,  
  features = NULL,  
  assay = NULL,  
  cluster_col = NULL,  
  n_celltypes = 50,  
  verbose = TRUE  
)
```

Arguments

<code>seurat_obj</code>	A Seurat object containing single-cell data
<code>features</code>	Character vector of feature names (genes) to analyze. If <code>NULL</code> , will use highly variable features from the Seurat object.
<code>assay</code>	Name of assay to use (default: default assay)
<code>cluster_col</code>	Column name in metadata containing cluster information
<code>n_celltypes</code>	Expected number of cell types in marker database (default: 50). Used for threshold recommendation calculation.
<code>verbose</code>	Whether to print progress messages (default: <code>TRUE</code>)

Value

A list containing:

- `min_expression`: Recommended expression threshold
- `specificity_weight`: Recommended specificity weight
- `threshold`: Recommended probability threshold for candidate selection
- `dataset_features`: Extracted dataset characteristics
- `parameter_rationale`: Explanation of parameter choices

See Also

Other Section_3_Automated_Annotation: [Celltype_Annotation\(\)](#), [Celltype_Annotation_PerCell\(\)](#), [Celltype_Calculate\(\)](#), [Celltype_Calculate_PerCell\(\)](#), [Celltype_Verification\(\)](#), [Celltype_Verification_PerCell\(\)](#), [percell_workflow](#)

Examples

```
## Not run:
SlimR_params <- Parameter_Calculate(
  seurat_obj = sce,
  features = c("CD3E", "CD4", "CD8A"),
  assay = "RNA",
  cluster_col = "seurat_clusters",
  n_celltypes = 98,
  verbose = TRUE
)

## End(Not run)
```

percell_workflow

Per-Cell Annotation Workflow Example

Description

Example workflow for using SlimR's per-cell annotation functions

Overview

The per-cell annotation workflow in SlimR provides an alternative to cluster-based annotation by scoring and labeling individual cells based on marker expression. This is useful when:

- Clusters contain mixed cell types
- You want finer-grained annotations
- Cell states exist on a continuum
- UMAP spatial context can improve annotation quality

Basic Workflow

```
# 1. Prepare your Seurat object (must have normalized data)
library(SlimR)
library(Seurat)

# 2. Create or load marker list
Markers_list <- Markers_filter_Cellmarker2(
  Cellmarker2,
  species = "Human",
  tissue_class = "Intestine"
)

# 3. Run per-cell annotation
result <- Celltype_Calculate_PerCell(
  seurat_obj = sce,
```

```

    gene_list = Markers_list,
    species = "Human",
    method = "weighted",          # "weighted", "mean", or "AUCell"
    min_expression = 0.1,
    min_score = 0.1,
    verbose = TRUE
  )

# 4. Annotate Seurat object
sce <- Celltype_Annotation_PerCell(
  seurat_obj = sce,
  SlimR_percell_result = result,
  plot_UMAP = TRUE,
  plot_confidence = TRUE,
  annotation_col = "Cell_type_PerCell"
)

# 5. Verify annotations
dotplot <- Celltype_Verification_PerCell(
  seurat_obj = sce,
  SlimR_percell_result = result,
  gene_number = 5,
  annotation_col = "Cell_type_PerCell"
)
print(dotplot)

```

Advanced

UMAP Spatial Smoothing:

```

# Use UMAP coordinates to smooth predictions via k-NN
# This reduces noise and improves consistency in spatial regions

result_smooth <- Celltype_Calculate_PerCell(
  seurat_obj = sce,
  gene_list = Markers_list,
  species = "Human",
  use_umap_smoothing = TRUE,
  k_neighbors = 20,          # Number of neighbors to consider
  smoothing_weight = 0.3,   # 30
  verbose = TRUE
)

# Compare smoothed vs unsmoothed
sce$Cell_type_Smooth <- result_smooth$Cell_annotatons$Predicted_cell_type
sce$Cell_type_Raw <- result$Cell_annotatons$Predicted_cell_type

DimPlot(sce, group.by = "Cell_type_Raw") |
  DimPlot(sce, group.by = "Cell_type_Smooth")

```

Scoring Methods Comparison

```

# Method 1: Weighted (recommended for most cases)
# Combines expression with marker specificity and detection rate
result_weighted <- Celltype_Calculate_PerCell(
  seurat_obj = sce,
  gene_list = Markers_list,
  species = "Human",
  method = "weighted"
)

# Method 2: Mean (simple, fast)
# Just averages normalized marker expression
result_mean <- Celltype_Calculate_PerCell(
  seurat_obj = sce,
  gene_list = Markers_list,
  species = "Human",
  method = "mean"
)

# Method 3: AUCell (rank-based, robust to batch effects)
# Scores based on proportion of markers in top 5
result_aucell <- Celltype_Calculate_PerCell(
  seurat_obj = sce,
  gene_list = Markers_list,
  species = "Human",
  method = "AUCell"
)

```

Comparing Cluster vs Per-Cell Annotation

```

# Cluster-based annotation (original SlimR approach)
cluster_result <- Celltype_Calculate(
  seurat_obj = sce,
  gene_list = Markers_list,
  species = "Human",
  cluster_col = "seurat_clusters"
)

sce <- Celltype_Annotation(
  seurat_obj = sce,
  cluster_col = "seurat_clusters",
  SlimR_anno_result = cluster_result,
  annotation_col = "Cell_type_Cluster"
)

# Per-cell annotation
percell_result <- Celltype_Calculate_PerCell(
  seurat_obj = sce,

```

```
    gene_list = Markers_list,
    species = "Human"
  )

sce <- Celltype_Annotation_PerCell(
  seurat_obj = sce,
  SlimR_percell_result = percell_result,
  annotation_col = "Cell_type_PerCell"
)

# Compare
library(ggplot2)
library(patchwork)

p1 <- DimPlot(sce, group.by = "Cell_type_Cluster") +
  ggtitle("Cluster-based")
p2 <- DimPlot(sce, group.by = "Cell_type_PerCell") +
  ggtitle("Per-cell")

p1 | p2

# Check agreement
table(sce$Cell_type_Cluster, sce$Cell_type_PerCell)
```

Performance Optimization

```
# For large datasets, adjust chunk_size to manage memory
result <- Celltype_Calculate_PerCell(
  seurat_obj = sce,
  gene_list = Markers_list,
  species = "Human",
  chunk_size = 10000, # Process 10k cells at a time
  verbose = TRUE
)

# For UMAP smoothing, install RANN for 10-100x speedup
# install.packages("RANN")

result_smooth <- Celltype_Calculate_PerCell(
  seurat_obj = sce,
  gene_list = Markers_list,
  species = "Human",
  use_umap_smoothing = TRUE,
  k_neighbors = 15
  # RANN will be used automatically if installed
)
```

Accessing Results

```

# Cell-level annotations
head(result$Cell_annotatons)
#   Cell_barcode Predicted_cell_type Max_score Confidence
# 1 AAACCTGAG... Enterocyte          0.85      0.62
# 2 AAACCTGCA... Goblet cell          0.72      0.45

# Summary statistics
result$Summary
#   Cell_type      Count Percentage
# 1 Enterocyte    5432  45.2
# 2 Goblet cell   2156  17.9

# Full probability matrix (if return_scores = TRUE)
result$Probability_matrix[1:5, 1:3]
#           Enterocyte Goblet_cell Stem_cell
# AAACCTGAG... 0.85      0.10      0.05

# Extract high-confidence cells
high_conf <- result$Cell_annotatons$Cell_barcode[
  result$Cell_annotatons$Confidence > 0.5
]

# Extract uncertain cells for manual review
uncertain <- result$Cell_annotatons$Cell_barcode[
  result$Cell_annotatons$Confidence < 0.2
]

```

See Also

Other Section_3_Automated_Annotation: [Celltype_Annotation\(\)](#), [Celltype_Annotation_PerCell\(\)](#), [Celltype_Calculate\(\)](#), [Celltype_Calculate_PerCell\(\)](#), [Celltype_Verification\(\)](#), [Celltype_Verification_PerCell\(\)](#), [Parameter_Calculate\(\)](#)

plot.pheatmap

Plot Method for pheatmap Objects

Description

This S3 method allows pheatmap objects (returned by `Celltype_Calculate()`) to be plotted using the generic `plot()` function. Without this method, attempting to use `plot()` on a pheatmap object results in an error.

Usage

```

## S3 method for class 'pheatmap'
plot(x, ...)

```

Arguments

x A pheatmap object, typically from `cluster_results$Heatmap_plot`
... Additional arguments (currently ignored)

Details

Pheatmap objects contain a `gtable` component that needs to be drawn using grid graphics. This method handles that automatically when `plot()` is called.

Alternative ways to display pheatmaps:

- `print(pheatmap_object)` - Works natively
- `plot(pheatmap_object)` - Works after loading SlimR
- `grid::grid.draw(pheatmap_object$gtable)` - Direct access

Value

Invisibly returns the input pheatmap object after displaying it

Examples

```
## Not run:  
# After running Celltype_Calculate()  
cluster_results <- Celltype_Calculate(  
  seurat_obj = sce,  
  gene_list = Markers_list,  
  species = "Human"  
)  
  
# Now both of these work:  
print(cluster_results$Heatmap_plot)  
plot(cluster_results$Heatmap_plot)  
  
## End(Not run)
```

Read_excel_markers *Create "Marker_list" from Excel files ".xlsx"*

Description

Create "Marker_list" from Excel files ".xlsx"

Usage

```
Read_excel_markers(path, has_colnames = TRUE)
```

Arguments

path	The path information of Marker files stored in ".xlsx" format. The Sheet name in the file is filled with cell type. The first line of each Sheet is the table head, the first column is filled with markers information, and the following column is filled with metric information.
has_colnames	Logical value indicating whether the first row contains column names. If FALSE, the first column will be named "Markers" and subsequent columns will be named "Col1", "Col2", etc.

Value

The standardized "Marker_list" in the SlimR package.

See Also

Other Section_2_Standardized_Markers_List: [Markers_filter_Cellmarker2\(\)](#), [Markers_filter_PanglaoDB\(\)](#), [Markers_filter_ScType\(\)](#), [Read_seurat_markers\(\)](#)

Examples

```
## Not run:
Markers_list_Excel <- Read_excel_markers(
  "D:/Laboratory/Marker_load.xlsx"
)

## End(Not run)
```

Read_seurat_markers *Create "Marker_list" from Seurat object*

Description

Create "Marker_list" from Seurat object

Usage

```
Read_seurat_markers(
  df,
  sources = c("Seurat", "presto"),
  sort_by = "FSS",
  gene_filter = 20
)
```

Arguments

df	Dataframe generated by "FindAllMarkers" function, recommend to use parameter "group.by = "Cell_type"" and "only.pos = TRUE".
sources	Type of markers sources to use. Be one of: "Seurat" or "presto".
sort_by	Marker sorting parameter, for Seurat sources, select "avg_log2FC" or "p_val_adj" or "FSS" (Feature Significance Score, FSS, product value of log2FC and Expression ratio). Default parameters use "sort_by = 'FSS'". for presto sources, select "logFC" or "padj" or "FSS". Default parameters use "sort_by = 'FSS'".
gene_filter	The number of markers left for each cell type based on the "sort_by" parameter's level of difference. Default parameters use "gene_filter = 20"

Value

The standardized "Marker_list" in the SlimR package.

See Also

Other Section_2_Standardized_Markers_List: [Markers_filter_Cellmarker2\(\)](#), [Markers_filter_PanglaoDB\(\)](#), [Markers_filter_ScType\(\)](#), [Read_excel_markers\(\)](#)

Examples

```
## Not run:
# Example for Seurat sources markers
seurat_markers <- Seurat::FindAllMarkers(
  object = sce,
  group.by = "Cell_type",
  only.pos = TRUE)

Markers_list_Seurat <- Read_seurat_markers(seurat_markers,
  sources = "Seurat",
  sort_by = "avg_log2FC",
  gene_filter = 20
)

# Example for presto sources markers
seurat_markers <- dplyr::filter(
  presto::wilcoxauc(
    X = sce,
    group_by = "Cell_type",
    seurat_assay = "RNA"
  ),
  padj < 0.05, logFC > 0.5
)

Markers_list_Seurat <- Read_seurat_markers(seurat_markers,
  sources = "presto",
  sort_by = "logFC",
  gene_filter = 20
)
```

```
## End(Not run)
```

ScType	<i>ScType dataset</i>
--------	-----------------------

Description

A processed long-format dataset containing marker genes for different cell types from the ScType database. Each row represents one marker gene for a given tissue type and cell type.

Usage

```
ScType
```

Format

A tibble with 3 columns:

tissue_type Tissue type (e.g., "Immune system", "Brain", "Liver")

cell_name Cell type name, formatted as "cellName(shortName)" when a short name is available, or "cellName" otherwise

marker Gene symbol of the marker

Details

This dataset is used to filter and create a standardized marker list. The dataset can be filtered based on tissue type and cell name to generate a list of marker genes for specific cell types using [Markers_filter_ScType](#).

Source

<https://github.com/IanevskiAleksandr/sc-type>

See Also

Other Section_0_Database: [Cellmarker2](#), [Cellmarker2_raw](#), [Cellmarker2_table](#), [Markers_list_PCTAM](#), [Markers_list_PCTIT](#), [Markers_list_TCellSI](#), [Markers_list_scIBD](#), [PanglaoDB](#), [PanglaoDB_raw](#), [PanglaoDB_table](#), [ScType_raw](#), [ScType_table](#)

ScType_raw	<i>ScType raw dataset</i>
------------	---------------------------

Description

The original ScType marker database before processing.

Usage

ScType_raw

Format

A tibble with 5 columns:

tissueType Tissue type

cellName Full cell type name

geneSymbolmore1 Comma-separated positive marker genes

geneSymbolmore2 Comma-separated negative marker genes (not used in processing)

shortName Abbreviated cell type name

Source

<https://github.com/IanevskiAleksandr/sc-type>

See Also

Other Section_0_Database: [Cellmarker2](#), [Cellmarker2_raw](#), [Cellmarker2_table](#), [Markers_list_PCTAM](#), [Markers_list_PCTIT](#), [Markers_list_TCellSI](#), [Markers_list_scIBD](#), [PanglaoDB](#), [PanglaoDB_raw](#), [PanglaoDB_table](#), [ScType](#), [ScType_table](#)

ScType_table	<i>ScType metadata table</i>
--------------	------------------------------

Description

A list of frequency tables summarizing the ScType database, useful for exploring available tissue types and cell types before filtering.

Usage

ScType_table

Format

A list with 2 elements:

tissue_type Frequency table of tissue types

cell_name Frequency table of cell type names

Source

<https://github.com/IanevskiAleksandr/sc-type>

See Also

Other Section_0_Database: [Cellmarker2](#), [Cellmarker2_raw](#), [Cellmarker2_table](#), [Markers_list_PCTAM](#), [Markers_list_PCTIT](#), [Markers_list_TCellSI](#), [Markers_list_scIBD](#), [PanglaoDB](#), [PanglaoDB_raw](#), [PanglaoDB_table](#), [ScType](#), [ScType_raw](#)

Index

- * **Section_0_Database**
 - Cellmarker2, 6
 - Cellmarker2_raw, 7
 - Cellmarker2_table, 8
 - Markers_list_PCTAM, 40
 - Markers_list_PCTIT, 40
 - Markers_list_scIBD, 41
 - Markers_list_TCellSI, 42
 - PanglaoDB, 43
 - PanglaoDB_raw, 43
 - PanglaoDB_table, 44
 - ScType, 54
 - ScType_raw, 55
 - ScType_table, 55
- * **Section_1_Functions_Use_in_Package**
 - calculate_cluster_variability, 3
 - calculate_expression, 4
 - calculate_expression_skewness, 5
 - calculate_probability, 5
 - compute_adaptive_parameters, 33
 - estimate_batch_effect, 35
 - extract_dataset_features, 36
- * **Section_2_Standardized_Markers_List**
 - Markers_filter_Cellmarker2, 37
 - Markers_filter_PanglaoDB, 38
 - Markers_filter_ScType, 39
 - Read_excel_markers, 51
 - Read_seurat_markers, 52
- * **Section_3_Automated_Annotation**
 - Celltype_Annotation, 8
 - Celltype_Annotation_PerCell, 20
 - Celltype_Calculate, 23
 - Celltype_Calculate_PerCell, 25
 - Celltype_Verification, 30
 - Celltype_Verification_PerCell, 31
 - Parameter_Calculate, 45
 - percell_workflow, 46
- * **Section_4_Semi_Automated_Annotation**
 - Celltype_Annotation_Combined, 11
 - Celltype_Annotation_Features, 14
 - Celltype_Annotation_Heatmap, 16
- * **Section_5_Other_Functions_Provided**
 - Celltype_annotation_Cellmarker2, 9
 - Celltype_annotation_Excel, 12
 - Celltype_annotation_PanglaoDB, 18
 - Celltype_annotation_Seurat, 21
 - Celltype_Compare, 28
 - Compute_Gene_AUC_ROC, 33
- * **datasets**
 - Cellmarker2, 6
 - Cellmarker2_raw, 7
 - Cellmarker2_table, 8
 - Markers_list_PCTAM, 40
 - Markers_list_PCTIT, 40
 - Markers_list_scIBD, 41
 - Markers_list_TCellSI, 42
 - PanglaoDB, 43
 - PanglaoDB_raw, 43
 - PanglaoDB_table, 44
 - ScType, 54
 - ScType_raw, 55
 - ScType_table, 55
- * **documentation**
 - percell_workflow, 46
- calculate_cluster_variability, 3
- calculate_cluster_variability(), 4–6, 33, 36
- calculate_expression, 4
- calculate_expression(), 3, 5, 6, 33, 36
- calculate_expression_skewness, 5
- calculate_expression_skewness(), 3, 4, 6, 33, 36
- calculate_probability, 5
- calculate_probability(), 3–5, 33, 36
- Cellmarker2, 6, 7, 8, 40–44, 54–56
- Cellmarker2_raw, 7, 7, 8, 40–44, 54–56
- Cellmarker2_table, 7, 8, 40–44, 54–56
- Celltype_Annotation, 8

- Celltype_Annotation(), 20, 24, 27, 31, 32, 45, 50
- Celltype_annotation_Cellmarker2, 9
- Celltype_annotation_Cellmarker2(), 13, 19, 22, 29, 35
- Celltype_Annotation_Combined, 11
- Celltype_Annotation_Combined(), 15, 18
- Celltype_annotation_Excel, 12
- Celltype_annotation_Excel(), 11, 19, 22, 29, 35
- Celltype_Annotation_Features, 14
- Celltype_Annotation_Features(), 12, 18
- Celltype_Annotation_Heatmap, 16
- Celltype_Annotation_Heatmap(), 12, 15
- Celltype_annotation_PanglaoDB, 18
- Celltype_annotation_PanglaoDB(), 11, 13, 22, 29, 35
- Celltype_Annotation_PerCell, 20
- Celltype_Annotation_PerCell(), 9, 24, 27, 31, 32, 45, 50
- Celltype_annotation_Seurat, 21
- Celltype_annotation_Seurat(), 11, 13, 19, 29, 35
- Celltype_Calculate, 23
- Celltype_Calculate(), 9, 20, 27, 31, 32, 45, 50
- Celltype_Calculate_PerCell, 25
- Celltype_Calculate_PerCell(), 9, 20, 24, 31, 32, 45, 50
- Celltype_Compare, 28
- Celltype_Compare(), 11, 13, 19, 22, 35
- Celltype_Verification, 30
- Celltype_Verification(), 9, 20, 24, 27, 32, 45, 50
- Celltype_Verification_PerCell, 31
- Celltype_Verification_PerCell(), 9, 20, 24, 27, 31, 45, 50
- compute_adaptive_parameters, 33
- compute_adaptive_parameters(), 3–6, 36
- Compute_Gene_AUC_ROC, 33
- Compute_Gene_AUC_ROC(), 11, 13, 19, 22, 29

- estimate_batch_effect, 35
- estimate_batch_effect(), 3–6, 33, 36
- extract_dataset_features, 36
- extract_dataset_features(), 3–6, 33, 36

- Markers_filter_Cellmarker2, 37
- Markers_filter_Cellmarker2(), 38, 39, 52, 53
- Markers_filter_PanglaoDB, 38
- Markers_filter_PanglaoDB(), 38, 39, 52, 53
- Markers_filter_ScType, 39, 54
- Markers_filter_ScType(), 38, 52, 53
- Markers_list_PCTAM, 7, 8, 40, 41–44, 54–56
- Markers_list_PCTIT, 7, 8, 40, 40, 42–44, 54–56
- Markers_list_scIBD, 7, 8, 40, 41, 41–44, 54–56
- Markers_list_TCellSI, 7, 8, 40, 41, 42, 42–44, 54–56

- PanglaoDB, 7, 8, 40–42, 43, 44, 54–56
- PanglaoDB_raw, 7, 8, 40–42, 43, 43, 44, 54–56
- PanglaoDB_table, 7, 8, 40–43, 44, 44, 54–56
- Parameter_Calculate, 45
- Parameter_Calculate(), 9, 20, 24, 27, 31, 32, 50
- percell_workflow, 9, 20, 24, 27, 31, 32, 45, 46
- plot.pheatmap, 50

- Read_excel_markers, 51
- Read_excel_markers(), 38, 39, 53
- Read_seurat_markers, 52
- Read_seurat_markers(), 38, 39, 52

- ScType, 7, 8, 40–44, 54, 55, 56
- ScType_raw, 7, 8, 40–44, 54, 55, 56
- ScType_table, 7, 8, 40–44, 54, 55, 55