

# Package ‘clustifyr’

July 13, 2025

**Title** Classifier for Single-cell RNA-seq Using Cell Clusters

**Version** 1.20.0

**Description** Package designed to aid in classifying cells from single-cell RNA sequencing data using external reference data (e.g., bulk RNA-seq, scRNA-seq, microarray, gene lists). A variety of correlation based methods and gene list enrichment methods are provided to assist cell type assignment.

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**Depends** R (>= 2.10)

**Imports** cowplot, dplyr, entropy, fgsea, ggplot2, Matrix, rlang,  
scales, stringr, tibble, tidyr, stats, methods,  
SingleCellExperiment, SummarizedExperiment, SeuratObject,  
matrixStats, S4Vectors, proxy, httr, utils

**Suggests** ComplexHeatmap, covr, knitr, rmarkdown, testthat, ggrepel,  
BiocStyle, BiocManager, remotes, shiny, gprofiler2, purrr,  
data.table, R.utils

**biocViews** SingleCell, Annotation, Sequencing, Microarray,  
GeneExpression

**BugReports** <https://github.com/rnabioco/clustifyr/issues>

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<https://rnabioco.github.io/clustifyr/>

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|              |           |
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---

|                   |                                                                          |
|-------------------|--------------------------------------------------------------------------|
| clustifyr-package | <i>clustifyr: Classifier for Single-cell RNA-seq Using Cell Clusters</i> |
|-------------------|--------------------------------------------------------------------------|

---

## Description

Package designed to aid in classifying cells from single-cell RNA sequencing data using external reference data (e.g., bulk RNA-seq, scRNA-seq, microarray, gene lists). A variety of correlation based methods and gene list enrichment methods are provided to assist cell type assignment.

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## See Also

Useful links:

- <https://github.com/rnabioco/clustifyr>
- <https://rnabioco.github.io/clustifyr/>
- Report bugs at <https://github.com/rnabioco/clustifyr/issues>

---

|              |                                                                                                                                                                          |
|--------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| append_genes | <i>Given a reference matrix and a list of genes, take the union of all genes in vector and genes in reference matrix and insert zero counts for all remaining genes.</i> |
|--------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Given a reference matrix and a list of genes, take the union of all genes in vector and genes in reference matrix and insert zero counts for all remaining genes.

**Usage**

```
append_genes(gene_vector, ref_matrix)
```

**Arguments**

|             |                                                                   |
|-------------|-------------------------------------------------------------------|
| gene_vector | char vector with gene names                                       |
| ref_matrix  | Reference matrix containing cell types vs. gene expression values |

**Value**

Reference matrix with union of all genes

**Examples**

```
mat <- append_genes(  
  gene_vector = human_genes_10x,  
  ref_matrix = cbmc_ref  
)
```

---

|                  |                       |
|------------------|-----------------------|
| assess_rank_bias | <i>Find rank bias</i> |
|------------------|-----------------------|

---

**Description**

Find rank bias

**Usage**

```
assess_rank_bias(  
  avg_mat,  
  ref_mat,  
  query_genes = NULL,  
  res,  
  organism,  
  plot_name = NULL,
```

```

    rds_name = NULL,
    expand_unassigned = FALSE
  )

```

### Arguments

|                   |                                                                              |
|-------------------|------------------------------------------------------------------------------|
| avg_mat           | average expression matrix                                                    |
| ref_mat           | reference expression matrix                                                  |
| query_genes       | original vector of genes used to clustify                                    |
| res               | dataframe of idents, such as output of cor_to_call                           |
| organism          | for GO term analysis, organism name: human - 'hsapiens', mouse - 'mmusculus' |
| plot_name         | name for saved pdf, if NULL then no file is written (default)                |
| rds_name          | name for saved rds of rank_diff, if NULL then no file is written (default)   |
| expand_unassigned | test all ref clusters for unassigned results                                 |

### Value

pdf of ggplot object

### Examples

```

## Not run:
avg <- average_clusters(
  pbmc_matrix_small,
  pbmc_meta$seurat_clusters
)
res <- clustify(
  input = pbmc_matrix_small,
  metadata = pbmc_meta,
  ref_mat = cbmc_ref,
  query_genes = pbmc_vargenes,
  cluster_col = "seurat_clusters"
)
top_call <- cor_to_call(
  res,
  metadata = pbmc_meta,
  cluster_col = "seurat_clusters",
  collapse_to_cluster = FALSE,
  threshold = 0.8
)
res_rank <- assess_rank_bias(
  avg,
  cbmc_ref,
  res = top_call
)

## End(Not run)

```

---

|              |                                      |
|--------------|--------------------------------------|
| assign_ident | <i>manually change ids as needed</i> |
|--------------|--------------------------------------|

---

**Description**

manually change ids as needed

**Usage**

```
assign_ident(  
  metadata,  
  cluster_col = "cluster",  
  ident_col = "type",  
  clusters,  
  ids  
)
```

**Arguments**

- metadata            column of id
- cluster\_col        column in metadata containing cluster info
- ident\_col          column in metadata containing identity assignment
- clusters           names of clusters to change, string or vector of strings
- ids                new ids to assign, must be length of 1 or same as clusters

**Value**

new dataframe of metadata

---

|                  |                                              |
|------------------|----------------------------------------------|
| average_clusters | <i>Average expression values per cluster</i> |
|------------------|----------------------------------------------|

---

**Description**

Average expression values per cluster

**Usage**

```
average_clusters(  
  mat,  
  metadata,  
  cluster_col = "cluster",  
  if_log = TRUE,  
  cell_col = NULL,
```

```

    low_threshold = 0,
    method = "mean",
    output_log = TRUE,
    subclusterpower = 0,
    cut_n = NULL
  )

```

## Arguments

|                              |                                                                                                                                                                                  |
|------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>mat</code>             | expression matrix                                                                                                                                                                |
| <code>metadata</code>        | data.frame or vector containing cluster assignments per cell. Order must match column order in supplied matrix. If a data.frame provide the <code>cluster_col</code> parameters. |
| <code>cluster_col</code>     | column in metadata with cluster number                                                                                                                                           |
| <code>if_log</code>          | input data is natural log, averaging will be done on unlogged data                                                                                                               |
| <code>cell_col</code>        | if provided, will reorder matrix first                                                                                                                                           |
| <code>low_threshold</code>   | option to remove clusters with too few cells                                                                                                                                     |
| <code>method</code>          | whether to take mean (default), median, 10% truncated mean, or trimean, max, min                                                                                                 |
| <code>output_log</code>      | whether to report log results                                                                                                                                                    |
| <code>subclusterpower</code> | whether to get multiple averages per original cluster                                                                                                                            |
| <code>cut_n</code>           | set on a limit of genes as expressed, lower ranked genes are set to 0, considered unexpressed                                                                                    |

## Value

average expression matrix, with genes for row names, and clusters for column names

## Examples

```

mat <- average_clusters(
  mat = pbmc_matrix_small,
  metadata = pbmc_meta,
  cluster_col = "classified",
  if_log = FALSE
)
mat[1:3, 1:3]

```



---

|               |                               |
|---------------|-------------------------------|
| binarize_expr | <i>Binarize scRNAseq data</i> |
|---------------|-------------------------------|

---

**Description**

Binarize scRNAseq data

**Usage**

```
binarize_expr(mat, n = 1000, cut = 0)
```

**Arguments**

|     |                                        |
|-----|----------------------------------------|
| mat | single-cell expression matrix          |
| n   | number of top expressing genes to keep |
| cut | cut off to set to 0                    |

**Value**

matrix of 1s and 0s

**Examples**

```
pbmc_avg <- average_clusters(  
  mat = pbmc_matrix_small,  
  metadata = pbmc_meta,  
  cluster_col = "classified"  
)  
  
mat <- binarize_expr(pbmc_avg)  
mat[1:3, 1:3]
```

---

|             |                                                      |
|-------------|------------------------------------------------------|
| build_atlas | <i>Function to combine records into single atlas</i> |
|-------------|------------------------------------------------------|

---

**Description**

Function to combine records into single atlas

**Usage**

```
build_atlas(matrix_fns = NULL, genes_fn, matrix_objs = NULL, output_fn = NULL)
```

**Arguments**

|             |                                                                                                                                                                                                                                                     |
|-------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| matrix_fns  | character vector of paths to study matrices stored as .rds files. If a named character vector, then the name will be added as a suffix to the cell type name in the final matrix. If it is not named, then the filename will be used (without .rds) |
| genes_fn    | text file with a single column containing genes and the ordering desired in the output matrix                                                                                                                                                       |
| matrix_objs | Checks to see whether .rds files will be read or R objects in a local environment. A list of environmental objects can be passed to matrix_objs, and that names will be used, otherwise defaults to numbers                                         |
| output_fn   | output filename for .rds file. If NULL the matrix will be returned instead of saving                                                                                                                                                                |

**Value**

Combined matrix with all genes given

**Examples**

```
pbmc_ref_matrix <- average_clusters(
  mat = pbmc_matrix_small,
  metadata = pbmc_meta,
  cluster_col = "classified",
  if_log = TRUE # whether the expression matrix is already log transformed
)
references_to_combine <- list(pbmc_ref_matrix, cbmc_ref)
atlas <- build_atlas(NULL, human_genes_10x, references_to_combine, NULL)
```

---

calculate\_pathway\_gsea

*Convert expression matrix to GSEA pathway scores (would take a similar place in workflow before average\_clusters/binarize)*

---

**Description**

Convert expression matrix to GSEA pathway scores (would take a similar place in workflow before average\_clusters/binarize)

**Usage**

```
calculate_pathway_gsea(
  mat,
  pathway_list,
  n_perm = 1000,
  scale = TRUE,
  no_warnings = TRUE
)
```

**Arguments**

|              |                                                                       |
|--------------|-----------------------------------------------------------------------|
| mat          | expression matrix                                                     |
| pathway_list | a list of vectors, each named for a specific pathway, or dataframe    |
| n_perm       | Number of permutation for fgsea function. Defaults to 1000.           |
| scale        | convert expr_mat into zscores prior to running GSEA?, default = FALSE |
| no_warnings  | suppress warnings from gsea ties                                      |

**Value**

matrix of GSEA NES values, cell types as row names, pathways as column names

**Examples**

```
gl <- list(
  "n" = c("PPBP", "LYZ", "S100A9"),
  "a" = c("IGLL5", "GNLY", "FTL")
)

pbmc_avg <- average_clusters(
  mat = pbmc_matrix_small,
  metadata = pbmc_meta,
  cluster_col = "classified"
)

calculate_pathway_gsea(
  mat = pbmc_avg,
  pathway_list = gl
)
```

---

|               |                                                |
|---------------|------------------------------------------------|
| calc_distance | <i>Distance calculations for spatial coord</i> |
|---------------|------------------------------------------------|

---

**Description**

Distance calculations for spatial coord

**Usage**

```
calc_distance(
  coord,
  metadata,
  cluster_col = "cluster",
  collapse_to_cluster = FALSE
)
```

**Arguments**

|                     |                                                                                                                                                                     |
|---------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| coord               | dataframe or matrix of spatial coordinates, cell barcode as rownames                                                                                                |
| metadata            | data.frame or vector containing cluster assignments per cell. Order must match column order in supplied matrix. If a data.frame provide the cluster_col parameters. |
| cluster_col         | column in metadata with cluster number                                                                                                                              |
| collapse_to_cluster | instead of reporting min distance to cluster per cell, summarize to cluster level                                                                                   |

**Value**

min distance matrix

**Examples**

```
cbs <- paste0("cb_", 1:100)

spatial_coords <- data.frame(
  row.names = cbs,
  X = runif(100),
  Y = runif(100)
)
group_ids <- sample(c("A", "B"), 100, replace = TRUE)
dist_res <- calc_distance(
  spatial_coords,
  group_ids
)
```

---

|                 |                           |
|-----------------|---------------------------|
| calc_similarity | <i>compute similarity</i> |
|-----------------|---------------------------|

---

**Description**

compute similarity

**Usage**

```
calc_similarity(query_mat, ref_mat, compute_method, rm0 = FALSE, ...)
```

**Arguments**

|                |                                                      |
|----------------|------------------------------------------------------|
| query_mat      | query data matrix                                    |
| ref_mat        | reference data matrix                                |
| compute_method | method(s) for computing similarity scores            |
| rm0            | consider 0 as missing data, recommended for per_cell |
| ...            | additional parameters                                |

**Value**

matrix of numeric values

---

|                |                                                    |
|----------------|----------------------------------------------------|
| call_consensus | <i>get consensus calls for a list of cor calls</i> |
|----------------|----------------------------------------------------|

---

**Description**

get concensus calls for a list of cor calls

**Usage**

```
call_consensus(list_of_res)
```

**Arguments**

list\_of\_res      list of call dataframes from cor\_to\_call\_rank

**Value**

dataframe of cluster, new ident, and mean rank

**Examples**

```
res <- clustify(  
  input = pbmc_matrix_small,  
  metadata = pbmc_meta,  
  cluster_col = "classified",  
  ref_mat = cbmc_ref  
)  
  
res2 <- cor_to_call_rank(res, threshold = "auto")  
res3 <- cor_to_call_rank(res)  
call_consensus(list(res2, res3))
```

---

|                  |                                                  |
|------------------|--------------------------------------------------|
| call_to_metadata | <i>Insert called ident results into metadata</i> |
|------------------|--------------------------------------------------|

---

**Description**

Insert called ident results into metadata

**Usage**

```
call_to_metadata(
  res,
  metadata,
  cluster_col,
  per_cell = FALSE,
  rename_prefix = NULL
)
```

**Arguments**

|               |                                                              |
|---------------|--------------------------------------------------------------|
| res           | dataframe of ids, such as output of cor_to_call              |
| metadata      | input metadata with tsne or umap coordinates and cluster ids |
| cluster_col   | metadata column, can be cluster or cellid                    |
| per_cell      | whether the res dataframe is listed per cell                 |
| rename_prefix | prefix to add to type and r column names                     |

**Value**

new metadata with added columns

**Examples**

```
res <- clustify(
  input = pbmc_matrix_small,
  metadata = pbmc_meta,
  cluster_col = "classified",
  ref_mat = cbmc_ref
)

res2 <- cor_to_call(res, cluster_col = "classified")

call_to_metadata(
  res = res2,
  metadata = pbmc_meta,
  cluster_col = "classified",
  rename_prefix = "assigned"
)
```

---

cbmc\_m

---

*reference marker matrix from seurat citeseq CBMC tutorial*


---

**Description**

reference marker matrix from seurat citeseq CBMC tutorial

**Usage**

cbmc\_m

**Format**

An object of class `data.frame` with 3 rows and 13 columns.

**Source**

[https://satijalab.org/seurat/v3.0/multimodal\\_vignette.html#identify-differentially-expressed-proteins](https://satijalab.org/seurat/v3.0/multimodal_vignette.html#identify-differentially-expressed-proteins)

**See Also**

Other data: [cbmc\\_ref](#), [downrefs](#), [human\\_genes\\_10x](#), [mouse\\_genes\\_10x](#), [pbmc\\_markers](#), [pbmc\\_markers\\_M3Drop](#), [pbmc\\_matrix\\_small](#), [pbmc\\_meta](#), [pbmc\\_vargenes](#)

---

cbmc\_ref

*reference matrix from seurat citeseq CBMC tutorial*

---

**Description**

reference matrix from seurat citeseq CBMC tutorial

**Usage**

cbmc\_ref

**Format**

An object of class `matrix` (inherits from `array`) with 2000 rows and 13 columns.

**Source**

[https://satijalab.org/seurat/v3.0/multimodal\\_vignette.html#identify-differentially-expressed-proteins](https://satijalab.org/seurat/v3.0/multimodal_vignette.html#identify-differentially-expressed-proteins)

**See Also**

Other data: [cbmc\\_m](#), [downrefs](#), [human\\_genes\\_10x](#), [mouse\\_genes\\_10x](#), [pbmc\\_markers](#), [pbmc\\_markers\\_M3Drop](#), [pbmc\\_matrix\\_small](#), [pbmc\\_meta](#), [pbmc\\_vargenes](#)

---

|                  |                                                                                                                         |
|------------------|-------------------------------------------------------------------------------------------------------------------------|
| check_raw_counts | <i>Given a count matrix, determine if the matrix has been either log-normalized, normalized, or contains raw counts</i> |
|------------------|-------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Given a count matrix, determine if the matrix has been either log-normalized, normalized, or contains raw counts

**Usage**

```
check_raw_counts(counts_matrix, max_log_value = 50)
```

**Arguments**

counts\_matrix    Count matrix containing scRNA-seq read data  
max\_log\_value    Static value to determine if a matrix is normalized

**Value**

String either raw counts, log-normalized or normalized

**Examples**

```
check_raw_counts(pbmc_matrix_small)
```

---

|          |                                                  |
|----------|--------------------------------------------------|
| clustify | <i>Compare scRNA-seq data to reference data.</i> |
|----------|--------------------------------------------------|

---

**Description**

Compare scRNA-seq data to reference data.

**Usage**

```
clustify(input, ...)

## Default S3 method:
clustify(
  input,
  ref_mat,
  metadata = NULL,
  cluster_col = NULL,
  query_genes = NULL,
  n_genes = 1000,
  per_cell = FALSE,
```



```
n_perm = 0,
compute_method = "spearman",
pseudobulk_method = "mean",
verbose = TRUE,
lookuptable = NULL,
rm0 = FALSE,
obj_out = TRUE,
seurat_out = obj_out,
vec_out = FALSE,
rename_prefix = NULL,
threshold = "auto",
low_threshold_cell = 0,
exclude_genes = c(),
if_log = TRUE,
organism = "hsapiens",
plot_name = NULL,
rds_name = NULL,
expand_unassigned = FALSE,
...
)

## S3 method for class 'Seurat'
clustify(
  input,
  ref_mat,
  cluster_col = NULL,
  query_genes = NULL,
  n_genes = 1000,
  per_cell = FALSE,
  n_perm = 0,
  compute_method = "spearman",
  pseudobulk_method = "mean",
  use_var_genes = TRUE,
  dr = "umap",
  obj_out = TRUE,
  seurat_out = obj_out,
  vec_out = FALSE,
  threshold = "auto",
  verbose = TRUE,
  rm0 = FALSE,
  rename_prefix = NULL,
  exclude_genes = c(),
  metadata = NULL,
  organism = "hsapiens",
  plot_name = NULL,
  rds_name = NULL,
  expand_unassigned = FALSE,
  ...
)
```

```

)

## S3 method for class 'SingleCellExperiment'
clustify(
  input,
  ref_mat,
  cluster_col = NULL,
  query_genes = NULL,
  per_cell = FALSE,
  n_perm = 0,
  compute_method = "spearman",
  pseudobulk_method = "mean",
  use_var_genes = TRUE,
  dr = "umap",
  obj_out = TRUE,
  seurat_out = obj_out,
  vec_out = FALSE,
  threshold = "auto",
  verbose = TRUE,
  rm0 = FALSE,
  rename_prefix = NULL,
  exclude_genes = c(),
  metadata = NULL,
  organism = "hsapiens",
  plot_name = NULL,
  rds_name = NULL,
  expand_unassigned = FALSE,
  ...
)

```

### Arguments

|                          |                                                                                                                                                                                      |
|--------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>input</code>       | single-cell expression matrix or Seurat object                                                                                                                                       |
| <code>...</code>         | additional arguments to pass to <code>compute_method</code> function                                                                                                                 |
| <code>ref_mat</code>     | reference expression matrix                                                                                                                                                          |
| <code>metadata</code>    | cell cluster assignments, supplied as a vector or data.frame. If data.frame is supplied then <code>cluster_col</code> needs to be set. Not required if running correlation per cell. |
| <code>cluster_col</code> | column in metadata that contains cluster ids per cell. Will default to first column of metadata if not supplied. Not required if running correlation per cell.                       |
| <code>query_genes</code> | A vector of genes of interest to compare. If NULL, then common genes between the <code>expr_mat</code> and <code>ref_mat</code> will be used for comparison.                         |
| <code>n_genes</code>     | number of genes limit for Seurat variable genes, by default 1000, set to 0 to use all variable genes (generally not recommended)                                                     |
| <code>per_cell</code>    | if true run per cell, otherwise per cluster.                                                                                                                                         |
| <code>n_perm</code>      | number of permutations, set to 0 by default                                                                                                                                          |

|                    |                                                                                                                               |
|--------------------|-------------------------------------------------------------------------------------------------------------------------------|
| compute_method     | method(s) for computing similarity scores                                                                                     |
| pseudobulk_method  | method used for summarizing clusters, options are mean (default), median, truncate (10% truncated mean), or trimean, max, min |
| verbose            | whether to report certain variables chosen and steps                                                                          |
| lookuptable        | if not supplied, will look in built-in table for object parsing                                                               |
| rm0                | consider 0 as missing data, recommended for per_cell                                                                          |
| obj_out            | whether to output object instead of cor matrix                                                                                |
| seurat_out         | output cor matrix or called seurat object (deprecated, use obj_out instead)                                                   |
| vec_out            | only output a result vector in the same order as metadata                                                                     |
| rename_prefix      | prefix to add to type and r column names                                                                                      |
| threshold          | identity calling minimum correlation score threshold, only used when obj_out = TRUE                                           |
| low_threshold_cell | option to remove clusters with too few cells                                                                                  |
| exclude_genes      | a vector of gene names to throw out of query                                                                                  |
| if_log             | input data is natural log, averaging will be done on unlogged data                                                            |
| organism           | for GO term analysis, organism name: human - 'hsapiens', mouse - 'mmusculus'                                                  |
| plot_name          | name for saved pdf, if NULL then no file is written (default)                                                                 |
| rds_name           | name for saved rds of rank_diff, if NULL then no file is written (default)                                                    |
| expand_unassigned  | test all ref clusters for unassigned results                                                                                  |
| use_var_genes      | if providing a seurat object, use the variable genes (stored in seurat_object@var.genes) as the query_genes.                  |
| dr                 | stored dimension reduction                                                                                                    |

## Value

single cell object with identity assigned in metadata, or matrix of correlation values, clusters from input as row names, cell types from ref\_mat as column names

## Examples

```
# Annotate a matrix and metadata
clustify(
  input = pbmc_matrix_small,
  metadata = pbmc_meta,
  ref_mat = cbmc_ref,
  query_genes = pbmc_vargenes,
  cluster_col = "RNA_snn_res.0.5",
  verbose = TRUE
)

# Annotate using a different method
```

```

clustify(
  input = pbmc_matrix_small,
  metadata = pbmc_meta,
  ref_mat = cbmc_ref,
  query_genes = pbmc_vargenes,
  cluster_col = "RNA_snn_res.0.5",
  compute_method = "cosine"
)

# Annotate a SingleCellExperiment object
sce <- sce_pbmc()
clustify(
  sce,
  cbmc_ref,
  cluster_col = "clusters",
  obj_out = TRUE,
  per_cell = FALSE,
  dr = "umap"
)

# Annotate a Seurat object
so <- so_pbmc()
clustify(
  so,
  cbmc_ref,
  cluster_col = "seurat_clusters",
  obj_out = TRUE,
  per_cell = FALSE,
  dr = "umap"
)

# Annotate (and return) a Seurat object per-cell
clustify(
  input = so,
  ref_mat = cbmc_ref,
  cluster_col = "seurat_clusters",
  obj_out = TRUE,
  per_cell = TRUE,
  dr = "umap"
)

```

---

clustifyr\_methods

*Correlation functions available in clustifyr*


---

## Description

Correlation functions available in clustifyr

## Usage

```
clustifyr_methods
```

**Format**

An object of class character of length 5.

**Examples**

```
clustifyr_methods
```

---

|                |                                                               |
|----------------|---------------------------------------------------------------|
| clustify_lists | <i>Main function to compare scRNA-seq data to gene lists.</i> |
|----------------|---------------------------------------------------------------|

---

**Description**

Main function to compare scRNA-seq data to gene lists.

**Usage**

```
clustify_lists(input, ...)

## Default S3 method:
clustify_lists(
  input,
  marker,
  marker_inmatrix = TRUE,
  metadata = NULL,
  cluster_col = NULL,
  if_log = TRUE,
  per_cell = FALSE,
  topn = 800,
  cut = 0,
  genome_n = 30000,
  metric = "hyper",
  output_high = TRUE,
  lookuptable = NULL,
  obj_out = TRUE,
  seurat_out = obj_out,
  vec_out = FALSE,
  rename_prefix = NULL,
  threshold = 0,
  low_threshold_cell = 0,
  verbose = TRUE,
  input_markers = FALSE,
  details_out = FALSE,
  ...
)

## S3 method for class 'Seurat'
clustify_lists(
```

```

    input,
    metadata = NULL,
    cluster_col = NULL,
    if_log = TRUE,
    per_cell = FALSE,
    topn = 800,
    cut = 0,
    marker,
    marker_inmatrix = TRUE,
    genome_n = 30000,
    metric = "hyper",
    output_high = TRUE,
    dr = "umap",
    obj_out = TRUE,
    seurat_out = obj_out,
    vec_out = FALSE,
    threshold = 0,
    rename_prefix = NULL,
    verbose = TRUE,
    details_out = FALSE,
    ...
)

## S3 method for class 'SingleCellExperiment'
clustify_lists(
  input,
  metadata = NULL,
  cluster_col = NULL,
  if_log = TRUE,
  per_cell = FALSE,
  topn = 800,
  cut = 0,
  marker,
  marker_inmatrix = TRUE,
  genome_n = 30000,
  metric = "hyper",
  output_high = TRUE,
  dr = "umap",
  obj_out = TRUE,
  seurat_out = obj_out,
  vec_out = FALSE,
  threshold = 0,
  rename_prefix = NULL,
  verbose = TRUE,
  details_out = FALSE,
  ...
)

```

**Arguments**

|                    |                                                                                                                                                                         |
|--------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| input              | single-cell expression matrix, Seurat object, or SingleCellExperiment                                                                                                   |
| ...                | passed to matrixize_markers                                                                                                                                             |
| marker             | matrix or dataframe of candidate genes for each cluster                                                                                                                 |
| marker_inmatrix    | whether markers genes are already in preprocessed matrix form                                                                                                           |
| metadata           | cell cluster assignments, supplied as a vector or data.frame. If data.frame is supplied then cluster_col needs to be set. Not required if running correlation per cell. |
| cluster_col        | column in metadata with cluster number                                                                                                                                  |
| if_log             | input data is natural log, averaging will be done on unlogged data                                                                                                      |
| per_cell           | compare per cell or per cluster                                                                                                                                         |
| topn               | number of top expressing genes to keep from input matrix                                                                                                                |
| cut                | expression cut off from input matrix                                                                                                                                    |
| genome_n           | number of genes in the genome                                                                                                                                           |
| metric             | adjusted p-value for hypergeometric test, or jaccard index                                                                                                              |
| output_high        | if true (by default to fit with rest of package), -log10 transform p-value                                                                                              |
| lookuptable        | if not supplied, will look in built-in table for object parsing                                                                                                         |
| obj_out            | whether to output object instead of cor matrix                                                                                                                          |
| seurat_out         | output cor matrix or called seurat object (deprecated, use obj_out instead)                                                                                             |
| vec_out            | only output a result vector in the same order as metadata                                                                                                               |
| rename_prefix      | prefix to add to type and r column names                                                                                                                                |
| threshold          | identity calling minimum correlation score threshold, only used when obj_out = T                                                                                        |
| low_threshold_cell | option to remove clusters with too few cells                                                                                                                            |
| verbose            | whether to report certain variables chosen and steps                                                                                                                    |
| input_markers      | whether input is marker data.frame of 0 and 1s (output of pos_neg_marker), and uses alternate enrichment mode                                                           |
| details_out        | whether to also output shared gene list from jaccard                                                                                                                    |
| dr                 | stored dimension reduction                                                                                                                                              |

**Value**

matrix of numeric values, clusters from input as row names, cell types from marker\_mat as column names

**Examples**

```
# Annotate a matrix and metadata

# Annotate using a different method
clustify_lists(
  input = pbmc_matrix_small,
  marker = cbmc_m,
  metadata = pbmc_meta,
  cluster_col = "classified",
  verbose = TRUE,
  metric = "jaccard"
)
```

---

|                |                                                                                         |
|----------------|-----------------------------------------------------------------------------------------|
| clustify_nudge | <i>Combined function to compare scRNA-seq data to bulk RNA-seq data and marker list</i> |
|----------------|-----------------------------------------------------------------------------------------|

---

**Description**

Combined function to compare scRNA-seq data to bulk RNA-seq data and marker list

**Usage**

```
clustify_nudge(input, ...)

## Default S3 method:
clustify_nudge(
  input,
  ref_mat,
  marker,
  metadata = NULL,
  cluster_col = NULL,
  query_genes = NULL,
  compute_method = "spearman",
  weight = 1,
  threshold = -Inf,
  dr = "umap",
  norm = "diff",
  call = TRUE,
  marker_inmatrix = TRUE,
  mode = "rank",
  obj_out = FALSE,
  seurat_out = obj_out,
  rename_prefix = NULL,
  lookuptable = NULL,
  ...
)
```



```
## S3 method for class 'Seurat'
clustify_nudge(
  input,
  ref_mat,
  marker,
  cluster_col = NULL,
  query_genes = NULL,
  compute_method = "spearman",
  weight = 1,
  obj_out = TRUE,
  seurat_out = obj_out,
  threshold = -Inf,
  dr = "umap",
  norm = "diff",
  marker_inmatrix = TRUE,
  mode = "rank",
  rename_prefix = NULL,
  ...
)
```

### Arguments

|                 |                                                                                                                                                                |
|-----------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------|
| input           | express matrix or object                                                                                                                                       |
| ...             | passed to <code>matrixize_markers</code>                                                                                                                       |
| ref_mat         | reference expression matrix                                                                                                                                    |
| marker          | matrix of markers                                                                                                                                              |
| metadata        | cell cluster assignments, supplied as a vector or data.frame. If data.frame is supplied then <code>cluster_col</code> needs to be set.                         |
| cluster_col     | column in metadata that contains cluster ids per cell. Will default to first column of metadata if not supplied. Not required if running correlation per cell. |
| query_genes     | A vector of genes of interest to compare. If NULL, then common genes between the <code>expr_mat</code> and <code>ref_mat</code> will be used for comparison.   |
| compute_method  | method(s) for computing similarity scores                                                                                                                      |
| weight          | relative weight for the gene list scores, when added to correlation score                                                                                      |
| threshold       | identity calling minimum score threshold, only used when <code>obj_out = T</code>                                                                              |
| dr              | stored dimension reduction                                                                                                                                     |
| norm            | whether and how the results are normalized                                                                                                                     |
| call            | make call or just return score matrix                                                                                                                          |
| marker_inmatrix | whether markers genes are already in preprocessed matrix form                                                                                                  |
| mode            | use marker expression pct or ranked cor score for nudging                                                                                                      |
| obj_out         | whether to output object instead of cor matrix                                                                                                                 |
| seurat_out      | output cor matrix or called seurat object (deprecated, use <code>obj_out</code> )                                                                              |
| rename_prefix   | prefix to add to type and r column names                                                                                                                       |
| lookuptable     | if not supplied, will look in built-in table for object parsing                                                                                                |

**Value**

single cell object, or matrix of numeric values, clusters from input as row names, cell types from ref\_mat as column names

**Examples**

```
# Seurat
so <- so_pbmc()
clustify_nudge(
  input = so,
  ref_mat = cbmc_ref,
  marker = cbmc_m,
  cluster_col = "seurat_clusters",
  threshold = 0.8,
  obj_out = FALSE,
  mode = "pct",
  dr = "umap"
)

# Matrix
clustify_nudge(
  input = pbmc_matrix_small,
  ref_mat = cbmc_ref,
  metadata = pbmc_meta,
  marker = as.matrix(cbmc_m),
  query_genes = pbmc_vargenes,
  cluster_col = "classified",
  threshold = 0.8,
  call = FALSE,
  marker_inmatrix = FALSE,
  mode = "pct"
)
```

---

|                     |                                                                    |
|---------------------|--------------------------------------------------------------------|
| collapse_to_cluster | <i>From per-cell calls, take highest freq call in each cluster</i> |
|---------------------|--------------------------------------------------------------------|

---

**Description**

From per-cell calls, take highest freq call in each cluster

**Usage**

```
collapse_to_cluster(res, metadata, cluster_col, threshold = 0)
```

**Arguments**

|             |                                                              |
|-------------|--------------------------------------------------------------|
| res         | dataframe of idents, such as output of cor_to_call           |
| metadata    | input metadata with tsne or umap coordinates and cluster ids |
| cluster_col | metadata column for cluster                                  |
| threshold   | minimum correlation coefficient cutoff for calling clusters  |

**Value**

new metadata with added columns

**Examples**

```
res <- clustify(
  input = pbmc_matrix_small,
  metadata = pbmc_meta,
  cluster_col = "classified",
  ref_mat = cbmc_ref,
  per_cell = TRUE
)

res2 <- cor_to_call(res)

collapse_to_cluster(
  res2,
  metadata = pbmc_meta,
  cluster_col = "classified",
  threshold = 0
)
```

---

|               |                                                                                           |
|---------------|-------------------------------------------------------------------------------------------|
| compare_lists | <i>Calculate adjusted p-values for hypergeometric test of gene lists or jaccard index</i> |
|---------------|-------------------------------------------------------------------------------------------|

---

**Description**

Calculate adjusted p-values for hypergeometric test of gene lists or jaccard index

**Usage**

```
compare_lists(
  bin_mat,
  marker_mat,
  n = 30000,
  metric = "hyper",
  output_high = TRUE,
  details_out = FALSE
)
```

**Arguments**

|             |                                                                             |
|-------------|-----------------------------------------------------------------------------|
| bin_mat     | binarized single-cell expression matrix, feed in by_cluster mat, if desired |
| marker_mat  | matrix or dataframe of candidate genes for each cluster                     |
| n           | number of genes in the genome                                               |
| metric      | adjusted p-value for hypergeometric test, or jaccard index                  |
| output_high | if true (by default to fit with rest of package), -log10 transform p-value  |
| details_out | whether to also output shared gene list from jaccard                        |

**Value**

matrix of numeric values, clusters from expr\_mat as row names, cell types from marker\_mat as column names

**Examples**

```
pbmc_mm <- matrixize_markers(pbmc_markers)

pbmc_avg <- average_clusters(
  pbmc_matrix_small,
  pbmc_meta,
  cluster_col = "classified"
)

pbmc_avgb <- binarize_expr(pbmc_avg)

compare_lists(
  pbmc_avgb,
  pbmc_mm,
  metric = "spearman"
)
```

---

|             |                                        |
|-------------|----------------------------------------|
| cor_to_call | <i>get best calls for each cluster</i> |
|-------------|----------------------------------------|

---

**Description**

get best calls for each cluster

**Usage**

```
cor_to_call(
  cor_mat,
  metadata = NULL,
  cluster_col = "cluster",
  collapse_to_cluster = FALSE,
  threshold = 0,
  rename_prefix = NULL,
  carry_r = FALSE
)
```

**Arguments**

|             |                                                              |
|-------------|--------------------------------------------------------------|
| cor_mat     | input similarity matrix                                      |
| metadata    | input metadata with tsne or umap coordinates and cluster ids |
| cluster_col | metadata column, can be cluster or cellid                    |

collapse\_to\_cluster  
                     if a column name is provided, takes the most frequent call of entire cluster to color in plot

threshold           minimum correlation coefficient cutoff for calling clusters

rename\_prefix       prefix to add to type and r column names

carry\_r             whether to include threshold in unassigned names

**Value**

dataframe of cluster, new ident, and r info

**Examples**

```
res <- clustify(
  input = pbmc_matrix_small,
  metadata = pbmc_meta,
  cluster_col = "classified",
  ref_mat = cbmc_ref
)

cor_to_call(res)
```

---

|                  |                                          |
|------------------|------------------------------------------|
| cor_to_call_rank | <i>get ranked calls for each cluster</i> |
|------------------|------------------------------------------|

---

**Description**

get ranked calls for each cluster

**Usage**

```
cor_to_call_rank(
  cor_mat,
  metadata = NULL,
  cluster_col = "cluster",
  collapse_to_cluster = FALSE,
  threshold = 0,
  rename_prefix = NULL,
  top_n = NULL
)
```

**Arguments**

cor\_mat             input similarity matrix

metadata           input metadata with tsne or umap coordinates and cluster ids

cluster\_col         metadata column, can be cluster or cellid

`collapse_to_cluster` if a column name is provided, takes the most frequent call of entire cluster to color in plot

`threshold` minimum correlation coefficient cutoff for calling clusters

`rename_prefix` prefix to add to type and r column names

`top_n` the number of ranks to keep, the rest will be set to 100

**Value**

dataframe of cluster, new ident, and r info

**Examples**

```
res <- clustify(
  input = pbmc_matrix_small,
  metadata = pbmc_meta,
  cluster_col = "classified",
  ref_mat = cbmc_ref
)

cor_to_call_rank(res, threshold = "auto")
```

---

|                               |                                       |
|-------------------------------|---------------------------------------|
| <code>cor_to_call_topn</code> | <i>get top calls for each cluster</i> |
|-------------------------------|---------------------------------------|

---

**Description**

get top calls for each cluster

**Usage**

```
cor_to_call_topn(
  cor_mat,
  metadata = NULL,
  col = "cluster",
  collapse_to_cluster = FALSE,
  threshold = 0,
  topn = 2
)
```

**Arguments**

`cor_mat` input similarity matrix

`metadata` input metadata with tsne or umap coordinates and cluster ids

`col` metadata column, can be cluster or cellid

collapse\_to\_cluster

threshold

topn

if a column name is provided, takes the most frequent call of entire cluster to color in plot

minimum correlation coefficient cutoff for calling clusters

number of calls for each cluster

Value

dataframe of cluster, new potential ident, and r info

Examples

```
res <- clustify(
  input = pbmc_matrix_small,
  metadata = pbmc_meta,
  ref_mat = cbmc_ref,
  query_genes = pbmc_vargenes,
  cluster_col = "classified"
)

cor_to_call_topn(
  cor_mat = res,
  metadata = pbmc_meta,
  col = "classified",
  collapse_to_cluster = FALSE,
  threshold = 0.5
)
```

---

|        |                        |
|--------|------------------------|
| cosine | <i>Cosine distance</i> |
|--------|------------------------|

---

Description

Cosine distance

Usage

```
cosine(vec1, vec2)
```

Arguments

vec1

vec2

test vector

reference vector

Value

numeric value of cosine distance between the vectors

---

|          |                                                    |
|----------|----------------------------------------------------|
| downrefs | <i>table of references stored in clustifyrdata</i> |
|----------|----------------------------------------------------|

---

**Description**

table of references stored in clustifyrdata

**Usage**

downrefs

**Format**

An object of class `tbl_df` (inherits from `tbl`, `data.frame`) with 9 rows and 6 columns.

**Source**

various packages

**See Also**

Other data: [cbmc\\_m](#), [cbmc\\_ref](#), [human\\_genes\\_10x](#), [mouse\\_genes\\_10x](#), [pbmc\\_markers](#), [pbmc\\_markers\\_M3Drop](#), [pbmc\\_matrix\\_small](#), [pbmc\\_meta](#), [pbmc\\_vargenes](#)

---

|                   |                                                          |
|-------------------|----------------------------------------------------------|
| downsample_matrix | <i>downsample matrix by cluster or completely random</i> |
|-------------------|----------------------------------------------------------|

---

**Description**

downsample matrix by cluster or completely random

**Usage**

```
downsample_matrix(  
  mat,  
  n = 1,  
  keep_cluster_proportions = TRUE,  
  metadata = NULL,  
  cluster_col = "cluster"  
)
```



**Arguments**

|                          |                                                                                                                                                                     |
|--------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| mat                      | expression matrix                                                                                                                                                   |
| n                        | number per cluster or fraction to keep                                                                                                                              |
| keep_cluster_proportions | whether to subsample                                                                                                                                                |
| metadata                 | data.frame or vector containing cluster assignments per cell. Order must match column order in supplied matrix. If a data.frame provide the cluster_col parameters. |
| cluster_col              | column in metadata with cluster number                                                                                                                              |

**Value**

new smaller mat with less cell\_id columns

**Examples**

```
set.seed(42)
mat <- downsample_matrix(
  mat = pbmc_matrix_small,
  metadata = pbmc_meta$classified,
  n = 10,
  keep_cluster_proportions = TRUE
)
mat[1:3, 1:3]
```

---

|                    |                                                      |
|--------------------|------------------------------------------------------|
| feature_select_PCA | <i>Returns a list of variable genes based on PCA</i> |
|--------------------|------------------------------------------------------|

---

**Description**

Extract genes, i.e. "features", based on the top loadings of principal components formed from the bulk expression data set

**Usage**

```
feature_select_PCA(
  mat = NULL,
  pcs = NULL,
  n_pcs = 10,
  percentile = 0.99,
  if_log = TRUE
)
```

**Arguments**

|            |                                                                                                                                                                    |
|------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| mat        | Expression matrix. Rownames are genes, colnames are single cell cluster name, and values are average single cell expression (log transformed).                     |
| pcs        | Precalculated pcs if available, will skip over processing on mat.                                                                                                  |
| n_pcs      | Number of PCs to selected gene loadings from. See the explore_PCA_corr.Rmd vignette for details.                                                                   |
| percentile | Select the percentile of absolute values of PCA loadings to select genes from. E.g. 0.999 would select the top point 1 percent of genes with the largest loadings. |
| if_log     | whether the data is already log transformed                                                                                                                        |

**Value**

vector of genes

**Examples**

```
feature_select_PCA(
  cbmc_ref,
  if_log = FALSE
)
```

---

|                   |                                                                                                             |
|-------------------|-------------------------------------------------------------------------------------------------------------|
| file_marker_parse | <i>takes files with positive and negative markers, as described in garnett, and returns list of markers</i> |
|-------------------|-------------------------------------------------------------------------------------------------------------|

---

**Description**

takes files with positive and negative markers, as described in garnett, and returns list of markers

**Usage**

```
file_marker_parse(filename)
```

**Arguments**

|          |                  |
|----------|------------------|
| filename | txt file to load |
|----------|------------------|

**Value**

list of positive and negative gene markers

**Examples**

```
marker_file <- system.file(
  "extdata",
  "hsPBMC_markers.txt",
  package = "clustifyr"
)

file_marker_parse(marker_file)
```

---

**find\_rank\_bias***Find rank bias*

---

**Description**

Find rank bias

**Usage**

```
find_rank_bias(avg_mat, ref_mat, query_genes = NULL)
```

**Arguments**

|             |                                           |
|-------------|-------------------------------------------|
| avg_mat     | average expression matrix                 |
| ref_mat     | reference expression matrix               |
| query_genes | original vector of genes used to clustify |

**Value**

list of matrix of rank diff values

**Examples**

```
avg <- average_clusters(
  mat = pbmc_matrix_small,
  metadata = pbmc_meta,
  cluster_col = "classified",
  if_log = FALSE
)

rankdiff <- find_rank_bias(
  avg,
  cbmc_ref,
  query_genes = pbmc_vargenes
)
```

---

|          |                                                           |
|----------|-----------------------------------------------------------|
| gene_pct | <i>pct of cells in each cluster that express genelist</i> |
|----------|-----------------------------------------------------------|

---

**Description**

pct of cells in each cluster that express genelist

**Usage**

```
gene_pct(matrix, genelist, clusters, returning = "mean")
```

**Arguments**

|           |                                                                      |
|-----------|----------------------------------------------------------------------|
| matrix    | expression matrix                                                    |
| genelist  | vector of marker genes for one identity                              |
| clusters  | vector of cluster identities                                         |
| returning | whether to return mean, min, or max of the gene pct in the gene list |

**Value**

vector of numeric values

---

|                  |                                                                         |
|------------------|-------------------------------------------------------------------------|
| gene_pct_markerm | <i>pct of cells in every cluster that express a series of genelists</i> |
|------------------|-------------------------------------------------------------------------|

---

**Description**

pct of cells in every cluster that express a series of genelists

**Usage**

```
gene_pct_markerm(matrix, marker_m, metadata, cluster_col = NULL, norm = NULL)
```

**Arguments**

|             |                                                                                                                                                                     |
|-------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| matrix      | expression matrix                                                                                                                                                   |
| marker_m    | matrixized markers                                                                                                                                                  |
| metadata    | data.frame or vector containing cluster assignments per cell. Order must match column order in supplied matrix. If a data.frame provide the cluster_col parameters. |
| cluster_col | column in metadata with cluster number                                                                                                                              |
| norm        | whether and how the results are normalized                                                                                                                          |

**Value**

matrix of numeric values, clusters from mat as row names, cell types from marker\_m as column names

**Examples**

```
gene_pct_markerm(
  matrix = pbmc_matrix_small,
  marker_m = cbmc_m,
  metadata = pbmc_meta,
  cluster_col = "classified"
)
```

---

get\_best\_match\_matrix    *Function to make best call from correlation matrix*

---

**Description**

Function to make best call from correlation matrix

**Usage**

```
get_best_match_matrix(cor_mat)
```

**Arguments**

cor\_mat                      correlation matrix

**Value**

matrix of 1s and 0s

---

get\_best\_str                      *Function to make call and attach score*

---

**Description**

Function to make call and attach score

**Usage**

```
get_best_str(name, best_mat, cor_mat, carry_cor = TRUE)
```

**Arguments**

|           |                                             |
|-----------|---------------------------------------------|
| name      | name of row to query                        |
| best_mat  | binarized call matrix                       |
| cor_mat   | correlation matrix                          |
| carry_cor | whether the correlation score gets reported |

**Value**

string with ident call and possibly cor value

---

|                     |                                           |
|---------------------|-------------------------------------------|
| get_common_elements | <i>Find entries shared in all vectors</i> |
|---------------------|-------------------------------------------|

---

**Description**

return entries found in all supplied vectors. If the vector supplied is NULL or NA, then it will be excluded from the comparison.

**Usage**

```
get_common_elements(...)
```

**Arguments**

...                    vectors

**Value**

vector of shared elements

---

|                |                                       |
|----------------|---------------------------------------|
| get_similarity | <i>Compute similarity of matrices</i> |
|----------------|---------------------------------------|

---

**Description**

Compute similarity of matrices

**Usage**

```

get_similarity(
  expr_mat,
  ref_mat,
  cluster_ids,
  compute_method,
  pseudobulk_method = "mean",
  per_cell = FALSE,
  rm0 = FALSE,
  if_log = TRUE,
  low_threshold = 0,
  ...
)

```

**Arguments**

|                   |                                                                                                                               |
|-------------------|-------------------------------------------------------------------------------------------------------------------------------|
| expr_mat          | single-cell expression matrix                                                                                                 |
| ref_mat           | reference expression matrix                                                                                                   |
| cluster_ids       | vector of cluster ids for each cell                                                                                           |
| compute_method    | method(s) for computing similarity scores                                                                                     |
| pseudobulk_method | method used for summarizing clusters, options are mean (default), median, truncate (10% truncated mean), or trimean, max, min |
| per_cell          | run per cell?                                                                                                                 |
| rm0               | consider 0 as missing data, recommended for per_cell                                                                          |
| if_log            | input data is natural log, averaging will be done on unlogged data                                                            |
| low_threshold     | option to remove clusters with too few cells                                                                                  |
| ...               | additional parameters not used yet                                                                                            |

**Value**

matrix of numeric values, clusters from expr\_mat as row names, cell types from ref\_mat as column names

---

|                    |                                                                |
|--------------------|----------------------------------------------------------------|
| get_ucsc_reference | <i>Build reference atlases from external UCSC cellbrowsers</i> |
|--------------------|----------------------------------------------------------------|

---

**Description**

Build reference atlases from external UCSC cellbrowsers

**Usage**

```
get_ucsc_reference(cb_url, cluster_col, ...)
```

**Arguments**

|             |                                                                                                                                                                                        |
|-------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| cb_url      | URL of cellbrowser dataset (e.g. <a href="http://cells.ucsc.edu/?ds=cortex-dev">http://cells.ucsc.edu/?ds=cortex-dev</a> ). Note that the URL must contain the ds=dataset-name suffix. |
| cluster_col | annotation field for summarizing gene expression (e.g. clustering, cell-type name, samples, etc.)                                                                                      |
| ...         | additional args passed to average_clusters                                                                                                                                             |

**Value**

reference matrix

**Examples**

```
## Not run:

# many datasets hosted by UCSC have UMI counts in the expression matrix
# set if_log = FALSE if the expression matrix has not been natural log transformed

get_ucsc_reference(
  cb_url = "https://cells.ucsc.edu/?ds=evocell+mus-musculus+marrow",
  cluster_col = "Clusters", if_log = FALSE
)

get_ucsc_reference(
  cb_url = "http://cells.ucsc.edu/?ds=muscle-cell-atlas",
  cluster_col = "cell_annotation",
  if_log = FALSE
)

## End(Not run)
```

---

|                   |                                                    |
|-------------------|----------------------------------------------------|
| get_unique_column | <i>Generate a unique column id for a dataframe</i> |
|-------------------|----------------------------------------------------|

---

**Description**

Generate a unique column id for a dataframe

**Usage**

```
get_unique_column(df, id = NULL)
```

**Arguments**

|    |                             |
|----|-----------------------------|
| df | dataframe with column names |
| id | desired id if unique        |



**Value**

character

---

|              |                                                       |
|--------------|-------------------------------------------------------|
| get_vargenes | <i>Generate variable gene list from marker matrix</i> |
|--------------|-------------------------------------------------------|

---

**Description**

Variable gene list is required for clustify main function. This function parses variables genes from a matrix input.

**Usage**

```
get_vargenes(marker_mat)
```

**Arguments**

marker\_mat      matrix or dataframe of candidate genes for each cluster

**Value**

vector of marker gene names

**Examples**

```
get_vargenes(cbmc_m)
```

---

|             |                                                          |
|-------------|----------------------------------------------------------|
| gmt_to_list | <i>convert gmt format of pathways to list of vectors</i> |
|-------------|----------------------------------------------------------|

---

**Description**

convert gmt format of pathways to list of vectors

**Usage**

```
gmt_to_list(
  path,
  cutoff = 0,
  sep = "\thttp://www.broadinstitute.org/gsea/msigdb/cards/.*?\t"
)
```

**Arguments**

|        |                                                  |
|--------|--------------------------------------------------|
| path   | gmt file path                                    |
| cutoff | remove pathways with less genes than this cutoff |
| sep    | sep used in file to split path and genes         |

**Value**

list of genes in each pathway

**Examples**

```
gmt_file <- system.file(
  "extdata",
  "c2.cp.reactome.v6.2.symbols.gmt.gz",
  package = "clustifyr"
)

gene.lists <- gmt_to_list(path = gmt_file)
length(gene.lists)
```

---

|                 |                                                          |
|-----------------|----------------------------------------------------------|
| human_genes_10x | <i>Vector of human genes for 10x cellranger pipeline</i> |
|-----------------|----------------------------------------------------------|

---

**Description**

Vector of human genes for 10x cellranger pipeline

**Usage**

```
human_genes_10x
```

**Format**

An object of class character of length 33514.

**Source**

<https://support.10xgenomics.com/single-cell-gene-expression/software/downloads/latest>

**See Also**

Other data: [cbmc\\_m](#), [cbmc\\_ref](#), [downrefs](#), [mouse\\_genes\\_10x](#), [pbmc\\_markers](#), [pbmc\\_markers\\_M3Drop](#), [pbmc\\_matrix\\_small](#), [pbmc\\_meta](#), [pbmc\\_vargenes](#)

---

|                    |                                                             |
|--------------------|-------------------------------------------------------------|
| insert_meta_object | <i>more flexible metadata update of single cell objects</i> |
|--------------------|-------------------------------------------------------------|

---

**Description**

more flexible metadata update of single cell objects

**Usage**

```
insert_meta_object(
  input,
  new_meta,
  type = class(input),
  meta_loc = NULL,
  lookuptable = NULL
)
```

**Arguments**

|             |                                                                 |
|-------------|-----------------------------------------------------------------|
| input       | input object                                                    |
| new_meta    | new metadata table to insert back into object                   |
| type        | look up predefined slots/loc                                    |
| meta_loc    | metadata location                                               |
| lookuptable | if not supplied, will look in built-in table for object parsing |

**Value**

new object with new metadata inserted

**Examples**

```
so <- so_pbmc()
insert_meta_object(so, seurat_meta(so, dr = "umap"))
```

---

|               |                      |
|---------------|----------------------|
| kl_divergence | <i>KL divergence</i> |
|---------------|----------------------|

---

**Description**

Use package entropy to compute Kullback-Leibler divergence. The function first converts each vector's reads to pseudo-number of transcripts by normalizing the total reads to total\_reads. The normalized read for each gene is then rounded to serve as the pseudo-number of transcripts. Function entropy::KL.shrink is called to compute the KL-divergence between the two vectors, and the maximal allowed divergence is set to max\_KL. Finally, a linear transform is performed to convert the KL divergence, which is between 0 and max\_KL, to a similarity score between -1 and 1.

**Usage**

```
kl_divergence(vec1, vec2, if_log = FALSE, total_reads = 1000, max_KL = 1)
```

**Arguments**

|             |                                                                                                                  |
|-------------|------------------------------------------------------------------------------------------------------------------|
| vec1        | Test vector                                                                                                      |
| vec2        | Reference vector                                                                                                 |
| if_log      | Whether the vectors are log-transformed. If so, the raw count should be computed before computing KL-divergence. |
| total_reads | Pseudo-library size                                                                                              |
| max_KL      | Maximal allowed value of KL-divergence.                                                                          |

**Value**

numeric value, with additional attributes, of kl divergence between the vectors

---

|               |                                                          |
|---------------|----------------------------------------------------------|
| make_comb_ref | <i>make combination ref matrix to assess intermixing</i> |
|---------------|----------------------------------------------------------|

---

**Description**

make combination ref matrix to assess intermixing

**Usage**

```
make_comb_ref(ref_mat, if_log = TRUE, sep = "_and_")
```

**Arguments**

|         |                                 |
|---------|---------------------------------|
| ref_mat | reference expression matrix     |
| if_log  | whether input data is natural   |
| sep     | separator for name combinations |

**Value**

expression matrix

**Examples**

```
ref <- make_comb_ref(
  cbmc_ref,
  sep = "_+_")
ref[1:3, 1:3]
```

---

|               |                                                                           |
|---------------|---------------------------------------------------------------------------|
| marker_select | <i>decide for one gene whether it is a marker for a certain cell type</i> |
|---------------|---------------------------------------------------------------------------|

---

**Description**

decide for one gene whether it is a marker for a certain cell type

**Usage**

```
marker_select(row1, cols, cut = 1, compto = 1)
```

**Arguments**

|        |                                                       |
|--------|-------------------------------------------------------|
| row1   | a numeric vector of expression values (row)           |
| cols   | a vector of cell types (column)                       |
| cut    | an expression minimum cutoff                          |
| compto | compare max expression to the value of next 1 or more |

**Value**

vector of cluster name and ratio value

**Examples**

```
pbmc_avg <- average_clusters(
  mat = pbmc_matrix_small,
  metadata = pbmc_meta,
  cluster_col = "classified",
  if_log = FALSE
)

marker_select(
  row1 = pbmc_avg["PPBP", ],
  cols = names(pbmc_avg["PPBP", ])
)
```

---

|                   |                                                 |
|-------------------|-------------------------------------------------|
| matrixize_markers | <i>Convert candidate genes list into matrix</i> |
|-------------------|-------------------------------------------------|

---

**Description**

Convert candidate genes list into matrix

Usage

```
matrixize_markers(  
  marker_df,  
  ranked = FALSE,  
  n = NULL,  
  step_weight = 1,  
  background_weight = 0,  
  unique = FALSE,  
  metadata = NULL,  
  cluster_col = "classified",  
  remove_rp = FALSE  
)
```

Arguments

|                   |                                                                                                                         |
|-------------------|-------------------------------------------------------------------------------------------------------------------------|
| marker_df         | dataframe of candidate genes, must contain "gene" and "cluster" columns, or a matrix of gene names to convert to ranked |
| ranked            | unranked gene list feeds into hyperp, the ranked gene list feeds into regular corr_coef                                 |
| n                 | number of genes to use                                                                                                  |
| step_weight       | ranked genes are tranformed into pseudo expression by descending weight                                                 |
| background_weight | ranked genes are tranformed into pseudo expression with added weight                                                    |
| unique            | whether to use only unique markers to 1 cluster                                                                         |
| metadata          | vector or dataframe of cluster names, should have column named cluster                                                  |
| cluster_col       | column for cluster names to replace original cluster, if metadata is dataframe                                          |
| remove_rp         | do not include rps, rpl, rp1-9 in markers                                                                               |

Value

matrix of unranked gene marker names, or matrix of ranked expression

Examples

```
matrixize_markers(pbmc_markers)
```

---

|                 |                                                          |
|-----------------|----------------------------------------------------------|
| mouse_genes_10x | <i>Vector of mouse genes for 10x cellranger pipeline</i> |
|-----------------|----------------------------------------------------------|

---

Description

Vector of mouse genes for 10x cellranger pipeline

Usage

```
mouse_genes_10x
```

**Format**

An object of class character of length 31017.

**Source**

<https://support.10xgenomics.com/single-cell-gene-expression/software/downloads/latest>

**See Also**

Other data: [cbmc\\_m](#), [cbmc\\_ref](#), [downrefs](#), [human\\_genes\\_10x](#), [pbmc\\_markers](#), [pbmc\\_markers\\_M3Drop](#), [pbmc\\_matrix\\_small](#), [pbmc\\_meta](#), [pbmc\\_vargenes](#)

---

|                    |                                                                 |
|--------------------|-----------------------------------------------------------------|
| not_pretty_palette | <i>black and white palette for plotting continous variables</i> |
|--------------------|-----------------------------------------------------------------|

---

**Description**

black and white palette for plotting continous variables

**Usage**

```
not_pretty_palette
```

**Format**

An object of class character of length 9.

**Value**

vector of colors

---

|             |                                       |
|-------------|---------------------------------------|
| object_data | <i>Function to access object data</i> |
|-------------|---------------------------------------|

---

**Description**

Function to access object data

**Usage**

```
object_data(object, ...)
```

```
## S3 method for class 'Seurat'
```

```
object_data(object, slot, n_genes = 1000, ...)
```

```
## S3 method for class 'SingleCellExperiment'
```

```
object_data(object, slot, ...)
```

**Arguments**

|         |                                                                                                                                  |
|---------|----------------------------------------------------------------------------------------------------------------------------------|
| object  | object after tsne or umap projections and clustering                                                                             |
| ...     | additional arguments                                                                                                             |
| slot    | data to access                                                                                                                   |
| n_genes | number of genes limit for Seurat variable genes, by default 1000, set to 0 to use all variable genes (generally not recommended) |

**Value**

expression matrix, with genes as row names, and cell types as column names

**Examples**

```
so <- so_pbmc()
mat <- object_data(
  object = so,
  slot = "data"
)
mat[1:3, 1:3]
sce <- sce_pbmc()
mat <- object_data(
  object = sce,
  slot = "data"
)
mat[1:3, 1:3]
```

---

|                   |                                                       |
|-------------------|-------------------------------------------------------|
| object_loc_lookup | <i>lookup table for single cell object structures</i> |
|-------------------|-------------------------------------------------------|

---

**Description**

lookup table for single cell object structures

**Usage**

```
object_loc_lookup()
```

**Value**

A list populated with standardized functions to access relevant data structures in multiple single cell data formats.



---

|            |                                                                     |
|------------|---------------------------------------------------------------------|
| object_ref | <i>Function to convert labelled object to avg expression matrix</i> |
|------------|---------------------------------------------------------------------|

---

**Description**

Function to convert labelled object to avg expression matrix

**Usage**

```
object_ref(input, ...)

## Default S3 method:
object_ref(
  input,
  cluster_col = NULL,
  var_genes_only = FALSE,
  assay_name = NULL,
  method = "mean",
  lookupable = NULL,
  if_log = TRUE,
  ...
)

## S3 method for class 'Seurat'
object_ref(
  input,
  cluster_col = NULL,
  var_genes_only = FALSE,
  assay_name = NULL,
  method = "mean",
  lookupable = NULL,
  if_log = TRUE,
  ...
)

## S3 method for class 'SingleCellExperiment'
object_ref(
  input,
  cluster_col = NULL,
  var_genes_only = FALSE,
  assay_name = NULL,
  method = "mean",
  lookupable = NULL,
  if_log = TRUE,
  ...
)
```

**Arguments**

|                |                                                                                                  |
|----------------|--------------------------------------------------------------------------------------------------|
| input          | object after tsne or umap projections and clustering                                             |
| ...            | additional arguments                                                                             |
| cluster_col    | column name where classified cluster names are stored in seurat meta data, cannot be "rn"        |
| var_genes_only | whether to keep only var.genes in the final matrix output, could also look up genes used for PCA |
| assay_name     | any additional assay data, such as ADT, to include. If more than 1, pass a vector of names       |
| method         | whether to take mean (default) or median                                                         |
| lookuptable    | if not supplied, will look in built-in table for object parsing                                  |
| if_log         | input data is natural log, averaging will be done on unlogged data                               |

**Value**

reference expression matrix, with genes as row names, and cell types as column names

**Examples**

```
so <- so_pbmc()
object_ref(
  so,
  cluster_col = "seurat_clusters"
)
```

---

overcluster

*Overcluster by kmeans per cluster*


---

**Description**

Overcluster by kmeans per cluster

**Usage**

```
overcluster(mat, cluster_id, power = 0.15)
```

**Arguments**

|            |                                           |
|------------|-------------------------------------------|
| mat        | expression matrix                         |
| cluster_id | list of ids per cluster                   |
| power      | decides the number of clusters for kmeans |

**Value**

new cluster\_id list of more clusters

**Examples**

```
res <- overcluster(
  mat = pbmc_matrix_small,
  cluster_id = split(colnames(pbmc_matrix_small), pbmc_meta$classified)
)
length(res)
```

---

|                  |                                                                  |
|------------------|------------------------------------------------------------------|
| overcluster_test | <i>compare clustering parameters and classification outcomes</i> |
|------------------|------------------------------------------------------------------|

---

**Description**

compare clustering parameters and classification outcomes

**Usage**

```
overcluster_test(
  expr,
  metadata,
  ref_mat,
  cluster_col,
  x_col = "UMAP_1",
  y_col = "UMAP_2",
  n = 5,
  ngenes = NULL,
  query_genes = NULL,
  threshold = 0,
  do_label = TRUE,
  do_legend = FALSE,
  newclustering = NULL,
  combine = TRUE
)
```

**Arguments**

|             |                                                                     |
|-------------|---------------------------------------------------------------------|
| expr        | expression matrix                                                   |
| metadata    | metadata including cluster info and dimension reduction plotting    |
| ref_mat     | reference matrix                                                    |
| cluster_col | column of clustering from metadata                                  |
| x_col       | column of metadata for x axis plotting                              |
| y_col       | column of metadata for y axis plotting                              |
| n           | expand n-fold for over/under clustering                             |
| ngenes      | number of genes to use for feature selection, use all genes if NULL |
| query_genes | vector, otherwise genes with be recalculated                        |

|               |                                                                                                  |
|---------------|--------------------------------------------------------------------------------------------------|
| threshold     | type calling threshold                                                                           |
| do_label      | whether to label each cluster at median center                                                   |
| do_legend     | whether to draw legend                                                                           |
| newclustering | use kmeans if NULL on dr or col name for second column of clustering                             |
| combine       | if TRUE return a single plot with combined panels, if FALSE return list of plots (default: TRUE) |

**Value**

faceted ggplot object

**Examples**

```
set.seed(42)
overcluster_test(
  expr = pbmc_matrix_small,
  metadata = pbmc_meta,
  ref_mat = cbmc_ref,
  cluster_col = "classified",
  x_col = "UMAP_1",
  y_col = "UMAP_2"
)
```

---

|                  |                                                     |
|------------------|-----------------------------------------------------|
| parse_loc_object | <i>more flexible parsing of single cell objects</i> |
|------------------|-----------------------------------------------------|

---

**Description**

more flexible parsing of single cell objects

**Usage**

```
parse_loc_object(
  input,
  type = class(input),
  expr_loc = NULL,
  meta_loc = NULL,
  var_loc = NULL,
  cluster_col = NULL,
  lookupable = NULL
)
```

**Arguments**

|             |                                                            |
|-------------|------------------------------------------------------------|
| input       | input object                                               |
| type        | look up predefined slots/loc                               |
| expr_loc    | function that extracts expression matrix                   |
| meta_loc    | function that extracts metadata                            |
| var_loc     | function that extracts variable genes                      |
| cluster_col | column of clustering from metadata                         |
| lookuptable | if not supplied, will use object_loc_lookup() for parsing. |

**Value**

list of expression, metadata, vargenes, cluster\_col info from object

**Examples**

```
so <- so_pbmc()
obj <- parse_loc_object(so)
length(obj)
```

---

|              |                                                                          |
|--------------|--------------------------------------------------------------------------|
| pbmc_markers | <i>Marker genes identified by Seurat from single-cell RNA-seq PBMCs.</i> |
|--------------|--------------------------------------------------------------------------|

---

**Description**

Dataframe of markers from Seurat FindAllMarkers function

**Usage**

```
pbmc_markers
```

**Format**

An object of class `data.frame` with 2304 rows and 7 columns.

**Source**

[pbmc\_matrix] processed by Seurat

**See Also**

Other data: [cbmc\\_m](#), [cbmc\\_ref](#), [downrefs](#), [human\\_genes\\_10x](#), [mouse\\_genes\\_10x](#), [pbmc\\_markers\\_M3Drop](#), [pbmc\\_matrix\\_small](#), [pbmc\\_meta](#), [pbmc\\_vargenes](#)

---

|                     |                                                                          |
|---------------------|--------------------------------------------------------------------------|
| pbmc_markers_M3Drop | <i>Marker genes identified by M3Drop from single-cell RNA-seq PBMCs.</i> |
|---------------------|--------------------------------------------------------------------------|

---

**Description**

Selected features of 3k pbmcs from Seurat3 tutorial

**Usage**

```
pbmc_markers_M3Drop
```

**Format**

A data frame with 3 variables:

**Source**

[pbmc\_matrix] processed by [M3Drop]

**See Also**

Other data: [cbmc\\_m](#), [cbmc\\_ref](#), [downrefs](#), [human\\_genes\\_10x](#), [mouse\\_genes\\_10x](#), [pbmc\\_markers](#), [pbmc\\_matrix\\_small](#), [pbmc\\_meta](#), [pbmc\\_vargenes](#)

---

|                   |                                             |
|-------------------|---------------------------------------------|
| pbmc_matrix_small | <i>Matrix of single-cell RNA-seq PBMCs.</i> |
|-------------------|---------------------------------------------|

---

**Description**

Count matrix of 3k pbmcs from Seurat3 tutorial, with only var.features

**Usage**

```
pbmc_matrix_small
```

**Format**

A sparseMatrix with genes as rows and cells as columns.

**Source**

[https://satijalab.org/seurat/v3.0/pbmc3k\\_tutorial.html](https://satijalab.org/seurat/v3.0/pbmc3k_tutorial.html)

**See Also**

Other data: [cbmc\\_m](#), [cbmc\\_ref](#), [downrefs](#), [human\\_genes\\_10x](#), [mouse\\_genes\\_10x](#), [pbmc\\_markers](#), [pbmc\\_markers\\_M3Drop](#), [pbmc\\_meta](#), [pbmc\\_vargenes](#)

---

|           |                                                 |
|-----------|-------------------------------------------------|
| pbmc_meta | <i>Meta-data for single-cell RNA-seq PBMCs.</i> |
|-----------|-------------------------------------------------|

---

**Description**

Metadata, including umap, of 3k pbmcs from Seurat3 tutorial

**Usage**

```
pbmc_meta
```

**Format**

An object of class `data.frame` with 2638 rows and 9 columns.

**Source**

[`pbmc_matrix`] processed by Seurat

**See Also**

Other data: [cbmc\\_m](#), [cbmc\\_ref](#), [downrefs](#), [human\\_genes\\_10x](#), [mouse\\_genes\\_10x](#), [pbmc\\_markers](#), [pbmc\\_markers\\_M3Drop](#), [pbmc\\_matrix\\_small](#), [pbmc\\_vargenes](#)

---

|               |                                                                            |
|---------------|----------------------------------------------------------------------------|
| pbmc_vargenes | <i>Variable genes identified by Seurat from single-cell RNA-seq PBMCs.</i> |
|---------------|----------------------------------------------------------------------------|

---

**Description**

Top 2000 variable genes from 3k pbmcs from Seurat3 tutorial

**Usage**

```
pbmc_vargenes
```

**Format**

An object of class `character` of length 2000.

**Source**

[`pbmc_matrix`] processed by Seurat

**See Also**

Other data: [cbmc\\_m](#), [cbmc\\_ref](#), [downrefs](#), [human\\_genes\\_10x](#), [mouse\\_genes\\_10x](#), [pbmc\\_markers](#), [pbmc\\_markers\\_M3Drop](#), [pbmc\\_matrix\\_small](#), [pbmc\\_meta](#)

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|                  |                                        |
|------------------|----------------------------------------|
| percent_clusters | <i>Percentage detected per cluster</i> |
|------------------|----------------------------------------|

---

**Description**

Percentage detected per cluster

**Usage**

```
percent_clusters(mat, metadata, cluster_col = "cluster", cut_num = 0.5)
```

**Arguments**

|             |                                        |
|-------------|----------------------------------------|
| mat         | expression matrix                      |
| metadata    | data.frame with cells                  |
| cluster_col | column in metadata with cluster number |
| cut_num     | binary cutoff for detection            |

**Value**

matrix of numeric values, with genes for row names, and clusters for column names

---

|                    |                                                           |
|--------------------|-----------------------------------------------------------|
| permute_similarity | <i>Compute a p-value for similarity using permutation</i> |
|--------------------|-----------------------------------------------------------|

---

**Description**

Permute cluster labels to calculate empirical p-value

**Usage**

```
permute_similarity(
  expr_mat,
  ref_mat,
  cluster_ids,
  n_perm,
  per_cell = FALSE,
  compute_method,
  pseudobulk_method = "mean",
  rm0 = FALSE,
  ...
)
```



**Arguments**

|                   |                                                                                                                               |
|-------------------|-------------------------------------------------------------------------------------------------------------------------------|
| expr_mat          | single-cell expression matrix                                                                                                 |
| ref_mat           | reference expression matrix                                                                                                   |
| cluster_ids       | clustering info of single-cell data assume that genes have ALREADY BEEN filtered                                              |
| n_perm            | number of permutations                                                                                                        |
| per_cell          | run per cell?                                                                                                                 |
| compute_method    | method(s) for computing similarity scores                                                                                     |
| pseudobulk_method | method used for summarizing clusters, options are mean (default), median, truncate (10% truncated mean), or trimean, max, min |
| rm0               | consider 0 as missing data, recommended for per_cell                                                                          |
| ...               | additional parameters                                                                                                         |

**Value**

matrix of numeric values

---

|                |                                                           |
|----------------|-----------------------------------------------------------|
| plot_best_call | <i>Plot best calls for each cluster on a tSNE or umap</i> |
|----------------|-----------------------------------------------------------|

---

**Description**

Plot best calls for each cluster on a tSNE or umap

**Usage**

```
plot_best_call(
  cor_mat,
  metadata,
  cluster_col = "cluster",
  collapse_to_cluster = FALSE,
  threshold = 0,
  x = "UMAP_1",
  y = "UMAP_2",
  plot_r = FALSE,
  per_cell = FALSE,
  ...
)
```

**Arguments**

|                     |                                                                                               |
|---------------------|-----------------------------------------------------------------------------------------------|
| cor_mat             | input similarity matrix                                                                       |
| metadata            | input metadata with tsne or umap coordinates and cluster ids                                  |
| cluster_col         | metadata column, can be cluster or cellid                                                     |
| collapse_to_cluster | if a column name is provided, takes the most frequent call of entire cluster to color in plot |
| threshold           | minimum correlation coefficient cutoff for calling clusters                                   |
| x                   | x variable                                                                                    |
| y                   | y variable                                                                                    |
| plot_r              | whether to include second plot of cor eff for best call                                       |
| per_cell            | whether the cor_mat was generate per cell or per cluster                                      |
| ...                 | passed to plot_dims                                                                           |

**Value**

ggplot object, cells projected by dr, colored by cell type classification

**Examples**

```
res <- clustify(
  input = pbmc_matrix_small,
  metadata = pbmc_meta,
  ref_mat = cbmc_ref,
  query_genes = pbmc_vargenes,
  cluster_col = "classified"
)

plot_best_call(
  cor_mat = res,
  metadata = pbmc_meta,
  cluster_col = "classified"
)
```

---

|           |                                                                                 |
|-----------|---------------------------------------------------------------------------------|
| plot_call | <i>Plot called clusters on a tSNE or umap, for each reference cluster given</i> |
|-----------|---------------------------------------------------------------------------------|

---

**Description**

Plot called clusters on a tSNE or umap, for each reference cluster given

**Usage**

```
plot_call(cor_mat, metadata, data_to_plot = colnames(cor_mat), ...)
```

**Arguments**

|              |                                                              |
|--------------|--------------------------------------------------------------|
| cor_mat      | input similarity matrix                                      |
| metadata     | input metadata with tsne or umap coordinates and cluster ids |
| data_to_plot | colname of data to plot, defaults to all                     |
| ...          | passed to plot_dims                                          |

**Value**

list of ggplot object, cells projected by dr, colored by cell type classification

---

|          |                                                   |
|----------|---------------------------------------------------|
| plot_cor | <i>Plot similarity measures on a tSNE or umap</i> |
|----------|---------------------------------------------------|

---

**Description**

Plot similarity measures on a tSNE or umap

**Usage**

```
plot_cor(
  cor_mat,
  metadata,
  data_to_plot = colnames(cor_mat),
  cluster_col = NULL,
  x = "UMAP_1",
  y = "UMAP_2",
  scale_legends = FALSE,
  ...
)
```

**Arguments**

|               |                                                                                                                                                                                                                 |
|---------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| cor_mat       | input similarity matrix                                                                                                                                                                                         |
| metadata      | input metadata with per cell tsne or umap coordinates and cluster ids                                                                                                                                           |
| data_to_plot  | colname of data to plot, defaults to all                                                                                                                                                                        |
| cluster_col   | colname of clustering data in metadata, defaults to rownames of the metadata if not supplied.                                                                                                                   |
| x             | metadata column name with 1st axis dimension. defaults to "UMAP_1".                                                                                                                                             |
| y             | metadata column name with 2nd axis dimension. defaults to "UMAP_2".                                                                                                                                             |
| scale_legends | if TRUE scale all legends to maximum values in entire correlation matrix. if FALSE scale legends to maximum for each plot. A two-element numeric vector can also be passed to supply custom values i.e. c(0, 1) |
| ...           | passed to plot_dims                                                                                                                                                                                             |

**Value**

list of ggplot objects, cells projected by dr, colored by cor values

**Examples**

```
res <- clustify(
  input = pbmc_matrix_small,
  metadata = pbmc_meta,
  ref_mat = cbmc_ref,
  query_genes = pbmc_vargenes,
  cluster_col = "classified"
)

plot_cor(
  cor_mat = res,
  metadata = pbmc_meta,
  data_to_plot = colnames(res)[1:2],
  cluster_col = "classified",
  x = "UMAP_1",
  y = "UMAP_2"
)
```

---

plot\_cor\_heatmap

*Plot similarity measures on heatmap*


---

**Description**

Plot similarity measures on heatmap

**Usage**

```
plot_cor_heatmap(
  cor_mat,
  metadata = NULL,
  cluster_col = NULL,
  col = not_pretty_palette,
  legend_title = NULL,
  ...
)
```

**Arguments**

|              |                                                                                               |
|--------------|-----------------------------------------------------------------------------------------------|
| cor_mat      | input similarity matrix                                                                       |
| metadata     | input metadata with per cell tsne or umap coordinates and cluster ids                         |
| cluster_col  | colname of clustering data in metadata, defaults to rownames of the metadata if not supplied. |
| col          | color ramp to use                                                                             |
| legend_title | legend title to pass to Heatmap                                                               |
| ...          | passed to Heatmap                                                                             |

**Value**

complexheatmap object

**Examples**

```
res <- clustify(
  input = pbmc_matrix_small,
  metadata = pbmc_meta,
  ref_mat = cbmc_ref,
  query_genes = pbmc_vargenes,
  cluster_col = "classified",
  per_cell = FALSE
)

plot_cor_heatmap(res)
```

---

plot\_dims

---

*Plot a tSNE or umap colored by feature.*


---

**Description**

Plot a tSNE or umap colored by feature.

**Usage**

```
plot_dims(
  data,
  x = "UMAP_1",
  y = "UMAP_2",
  feature = NULL,
  legend_name = "",
  c_cols = pretty_palette2,
  d_cols = NULL,
  pt_size = 0.25,
  alpha_col = NULL,
  group_col = NULL,
  scale_limits = NULL,
  do_label = FALSE,
  do_legend = TRUE,
  do_repel = TRUE
)
```

**Arguments**

|      |            |
|------|------------|
| data | input data |
| x    | x variable |
| y    | y variable |

|              |                                                                                                                      |
|--------------|----------------------------------------------------------------------------------------------------------------------|
| feature      | feature to color by                                                                                                  |
| legend_name  | legend name to display, defaults to no name                                                                          |
| c_cols       | character vector of colors to build color gradient for continuous values, defaults to <a href="#">pretty_palette</a> |
| d_cols       | character vector of colors for discrete values. defaults to RColorBrewer paired palette                              |
| pt_size      | point size                                                                                                           |
| alpha_col    | whether to refer to data column for alpha values                                                                     |
| group_col    | group by another column instead of feature, useful for labels                                                        |
| scale_limits | defaults to min = 0, max = max(data\$x), otherwise a two-element numeric vector indicating min and max to plot       |
| do_label     | whether to label each cluster at median center                                                                       |
| do_legend    | whether to draw legend                                                                                               |
| do_repel     | whether to use ggrepel on labels                                                                                     |

**Value**

ggplot object, cells projected by dr, colored by feature

**Examples**

```
plot_dims(
  pbmc_meta,
  feature = "classified"
)
```

---

|           |                                                |
|-----------|------------------------------------------------|
| plot_gene | <i>Plot gene expression on to tSNE or umap</i> |
|-----------|------------------------------------------------|

---

**Description**

Plot gene expression on to tSNE or umap

**Usage**

```
plot_gene(expr_mat, metadata, genes, cell_col = NULL, ...)
```

**Arguments**

|          |                                                                                   |
|----------|-----------------------------------------------------------------------------------|
| expr_mat | input single cell matrix                                                          |
| metadata | data.frame with tSNE or umap coordinates                                          |
| genes    | gene(s) to color tSNE or umap                                                     |
| cell_col | column name in metadata containing cell ids, defaults to rownames if not supplied |
| ...      | additional arguments passed to <code>[clustifyr::plot_dims()]</code>              |

**Value**

list of ggplot object, cells projected by dr, colored by gene expression

**Examples**

```
genes <- c(
  "RP11-314N13.3",
  "ARF4"
)

plot_gene(
  expr_mat = pbmc_matrix_small,
  metadata = tibble::rownames_to_column(pbmc_meta, "rn"),
  genes = genes,
  cell_col = "rn"
)
```

---

|                   |                                                                                         |
|-------------------|-----------------------------------------------------------------------------------------|
| plot_pathway_gsea | <i>plot GSEA pathway scores as heatmap, returns a list containing results and plot.</i> |
|-------------------|-----------------------------------------------------------------------------------------|

---

**Description**

plot GSEA pathway scores as heatmap, returns a list containing results and plot.

**Usage**

```
plot_pathway_gsea(
  mat,
  pathway_list,
  n_perm = 1000,
  scale = TRUE,
  topn = 5,
  returning = "both"
)
```

**Arguments**

|              |                                                                      |
|--------------|----------------------------------------------------------------------|
| mat          | expression matrix                                                    |
| pathway_list | a list of vectors, each named for a specific pathway, or dataframe   |
| n_perm       | Number of permutation for fgsea function. Defaults to 1000.          |
| scale        | convert expr_mat into zscores prior to running GSEA?, default = TRUE |
| topn         | number of top pathways to plot                                       |
| returning    | to return "both" list and plot, or either one                        |

**Value**

list of matrix and plot, or just plot, matrix of GSEA NES values, cell types as row names, pathways as column names

**Examples**

```
gl <- list(
  "n" = c("PPBP", "LYZ", "S100A9"),
  "a" = c("IGLL5", "GNLY", "FTL")
)

pbmc_avg <- average_clusters(
  mat = pbmc_matrix_small,
  metadata = pbmc_meta,
  cluster_col = "classified"
)

plot_pathway_gsea(
  pbmc_avg,
  gl,
  5
)
```

---

|                |                                |
|----------------|--------------------------------|
| plot_rank_bias | <i>Query rank bias results</i> |
|----------------|--------------------------------|

---

**Description**

Query rank bias results

**Usage**

```
plot_rank_bias(bias_df, organism = "hsapiens")
```

**Arguments**

|          |                                                                              |
|----------|------------------------------------------------------------------------------|
| bias_df  | data.frame of rank diff matrix between cluster and reference cell types      |
| organism | for GO term analysis, organism name: human - 'hsapiens', mouse - 'mmusculus' |

**Value**

ggplot object of distribution and annotated GO terms



**Examples**

```
## Not run:
avg <- average_clusters(
  mat = pbmc_matrix_small,
  metadata = pbmc_meta,
  cluster_col = "classified",
  if_log = FALSE
)

rankdiff <- find_rank_bias(
  avg,
  cbmc_ref,
  query_genes = pbmc_vargenes
)

qres <- query_rank_bias(
  rankdiff,
  "CD14+ Mono",
  "CD14+ Mono"
)

g <- plot_rank_bias(
  qres
)

## End(Not run)
```

---

|                |                                                                                                     |
|----------------|-----------------------------------------------------------------------------------------------------|
| pos_neg_marker | <i>generate pos and negative marker expression matrix from a list/dataframe of positive markers</i> |
|----------------|-----------------------------------------------------------------------------------------------------|

---

**Description**

generate pos and negative marker expression matrix from a list/dataframe of positive markers

**Usage**

```
pos_neg_marker(mat)
```

**Arguments**

mat                      matrix or dataframe of markers

**Value**

matrix of gene expression

**Examples**

```
m1 <- pos_neg_marker(cbmc_m)
```

---

|                |                                                              |
|----------------|--------------------------------------------------------------|
| pos_neg_select | <i>adapt clustify to tweak score for pos and neg markers</i> |
|----------------|--------------------------------------------------------------|

---

## Description

adapt clustify to tweak score for pos and neg markers

## Usage

```
pos_neg_select(
  input,
  ref_mat,
  metadata,
  cluster_col = "cluster",
  cutoff_n = 0,
  cutoff_score = 0.5
)
```

## Arguments

|              |                                                                                                                                                                         |
|--------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| input        | single-cell expression matrix                                                                                                                                           |
| ref_mat      | reference expression matrix with positive and negative markers(set expression at 0)                                                                                     |
| metadata     | cell cluster assignments, supplied as a vector or data.frame. If data.frame is supplied then cluster_col needs to be set. Not required if running correlation per cell. |
| cluster_col  | column in metadata that contains cluster ids per cell. Will default to first column of metadata if not supplied. Not required if running correlation per cell.          |
| cutoff_n     | expression cutoff where genes ranked below n are considered non-expressing                                                                                              |
| cutoff_score | positive score lower than this cutoff will be considered as 0 to not influence scores                                                                                   |

## Value

matrix of numeric values, clusters from input as row names, cell types from ref\_mat as column names

## Examples

```
pn_ref <- data.frame(
  "Myeloid" = c(1, 0.01, 0),
  row.names = c("CD74", "clustifyr0", "CD79A")
)

pos_neg_select(
  input = pbmc_matrix_small,
```

```
    ref_mat = pn_ref,  
    metadata = pbmc_meta,  
    cluster_col = "classified",  
    cutoff_score = 0.8  
  )
```

---

|                |                                                       |
|----------------|-------------------------------------------------------|
| pretty_palette | <i>Color palette for plotting continous variables</i> |
|----------------|-------------------------------------------------------|

---

**Description**

Color palette for plotting continous variables

**Usage**

```
pretty_palette
```

**Format**

An object of class character of length 6.

**Value**

vector of colors

---

|                 |                                                                         |
|-----------------|-------------------------------------------------------------------------|
| pretty_palette2 | <i>Color palette for plotting continous variables, starting at gray</i> |
|-----------------|-------------------------------------------------------------------------|

---

**Description**

Color palette for plotting continous variables, starting at gray

**Usage**

```
pretty_palette2
```

**Format**

An object of class character of length 9.

**Value**

vector of colors

---

|                       |                                                                    |
|-----------------------|--------------------------------------------------------------------|
| pretty_palette_ramp_d | <i>Expanded color palette ramp for plotting discrete variables</i> |
|-----------------------|--------------------------------------------------------------------|

---

**Description**

Expanded color palette ramp for plotting discrete variables

**Usage**

```
pretty_palette_ramp_d(n)
```

**Arguments**

n                      number of colors to use

**Value**

color ramp

---

|                 |                                |
|-----------------|--------------------------------|
| query_rank_bias | <i>Query rank bias results</i> |
|-----------------|--------------------------------|

---

**Description**

Query rank bias results

**Usage**

```
query_rank_bias(bias_list, id_mat, id_ref)
```

**Arguments**

bias\_list              list of rank diff matrix between cluster and reference cell types  
id\_mat                  name of cluster from average cluster matrix  
id\_ref                  name of cell type in reference matrix

**Value**

data.frame rank diff values

**Examples**

```
avg <- average_clusters(  
  mat = pbmc_matrix_small,  
  metadata = pbmc_meta,  
  cluster_col = "classified",  
  if_log = FALSE  
)  
  
rankdiff <- find_rank_bias(  
  avg,  
  cbmc_ref,  
  query_genes = pbmc_vargenes  
)  
  
qres <- query_rank_bias(  
  rankdiff,  
  "CD14+ Mono",  
  "CD14+ Mono"  
)
```

---

|                    |                                             |
|--------------------|---------------------------------------------|
| ref_feature_select | <i>feature select from reference matrix</i> |
|--------------------|---------------------------------------------|

---

**Description**

feature select from reference matrix

**Usage**

```
ref_feature_select(mat, n = 3000, mode = "var", rm.lowvar = TRUE)
```

**Arguments**

|           |                                               |
|-----------|-----------------------------------------------|
| mat       | reference matrix                              |
| n         | number of genes to return                     |
| mode      | the method of selecting features              |
| rm.lowvar | whether to remove lower variation genes first |

**Value**

vector of genes

**Examples**

```
pbmc_avg <- average_clusters(
  mat = pbmc_matrix_small,
  metadata = pbmc_meta,
  cluster_col = "classified"
)

ref_feature_select(
  mat = pbmc_avg[1:100, ],
  n = 5
)
```

---

|                   |                                               |
|-------------------|-----------------------------------------------|
| ref_marker_select | <i>marker selection from reference matrix</i> |
|-------------------|-----------------------------------------------|

---

**Description**

marker selection from reference matrix

**Usage**

```
ref_marker_select(mat, cut = 0.5, arrange = TRUE, compto = 1)
```

**Arguments**

|         |                                                       |
|---------|-------------------------------------------------------|
| mat     | reference matrix                                      |
| cut     | an expression minimum cutoff                          |
| arrange | whether to arrange (lower means better)               |
| compto  | compare max expression to the value of next 1 or more |

**Value**

dataframe, with gene, cluster, ratio columns

**Examples**

```
ref_marker_select(
  cbmc_ref,
  cut = 2
)
```

---

|                       |                                                                            |
|-----------------------|----------------------------------------------------------------------------|
| reverse_marker_matrix | <i>generate negative markers from a list of exclusive positive markers</i> |
|-----------------------|----------------------------------------------------------------------------|

---

**Description**

generate negative markers from a list of exclusive positive markers

**Usage**

```
reverse_marker_matrix(mat)
```

**Arguments**

|     |                                |
|-----|--------------------------------|
| mat | matrix or dataframe of markers |
|-----|--------------------------------|

**Value**

matrix of gene names

**Examples**

```
reverse_marker_matrix(cbmc_m)
```

---

|                   |                                                                                                                         |
|-------------------|-------------------------------------------------------------------------------------------------------------------------|
| run_clustifyr_app | <i>Launch Shiny app version of clustifyr; may need to run install_clustifyr_app() at first time to install packages</i> |
|-------------------|-------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Launch Shiny app version of clustifyr, may need to run install\_clustifyr\_app() at first time to install packages

**Usage**

```
run_clustifyr_app()
```

**Value**

instance of shiny app

**Examples**

```
## Not run:  
run_clustifyr_app()  
  
## End(Not run)
```

---

|          |                                                                                      |
|----------|--------------------------------------------------------------------------------------|
| run_gsea | <i>Run GSEA to compare a gene list(s) to per cell or per cluster expression data</i> |
|----------|--------------------------------------------------------------------------------------|

---

## Description

Use fgsea algorithm to compute normalized enrichment scores and pvalues for gene set overlap

## Usage

```
run_gsea(  
  expr_mat,  
  query_genes,  
  cluster_ids = NULL,  
  n_perm = 1000,  
  per_cell = FALSE,  
  scale = FALSE,  
  no_warnings = TRUE  
)
```

## Arguments

|             |                                                                                                                                                              |
|-------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------|
| expr_mat    | single-cell expression matrix or Seurat object                                                                                                               |
| query_genes | A vector or named list of vectors of genesets of interest to compare via GSEA. If supplying a named list, then the gene set names will appear in the output. |
| cluster_ids | vector of cell cluster assignments, supplied as a vector with order that matches columns in expr_mat. Not required if running per cell.                      |
| n_perm      | Number of permutation for fgsea function. Defaults to 1000.                                                                                                  |
| per_cell    | if true run per cell, otherwise per cluster.                                                                                                                 |
| scale       | convert expr_mat into zscores prior to running GSEA?, default = FALSE                                                                                        |
| no_warnings | suppress warnings from gsea ties                                                                                                                             |

## Value

dataframe of gsea scores (pval, NES), with clusters as rownames



---

|          |                                               |
|----------|-----------------------------------------------|
| sce_pbmc | <i>An example SingleCellExperiment object</i> |
|----------|-----------------------------------------------|

---

**Description**

An example SingleCellExperiment object

**Usage**

```
sce_pbmc()
```

**Value**

a SingleCellExperiment object populated with data from the [pbmc\\_matrix\\_small](#) scRNA-seq dataset, additionally annotated with cluster assignments.

---

|             |                                                                              |
|-------------|------------------------------------------------------------------------------|
| seurat_meta | <i>Function to convert labelled seurat object to fully prepared metadata</i> |
|-------------|------------------------------------------------------------------------------|

---

**Description**

Function to convert labelled seurat object to fully prepared metadata

**Usage**

```
seurat_meta(seurat_object, ...)
```

```
## S3 method for class 'Seurat'
```

```
seurat_meta(seurat_object, dr = "umap", ...)
```

**Arguments**

|               |                                                             |
|---------------|-------------------------------------------------------------|
| seurat_object | seurat_object after tsne or umap projections and clustering |
| ...           | additional arguments                                        |
| dr            | dimension reduction method                                  |

**Value**

dataframe of metadata, including dimension reduction plotting info

**Examples**

```
so <- so_pbmc()
m <- seurat_meta(so)
```

---

|            |                                                                            |
|------------|----------------------------------------------------------------------------|
| seurat_ref | <i>Function to convert labelled seurat object to avg expression matrix</i> |
|------------|----------------------------------------------------------------------------|

---

## Description

Function to convert labelled seurat object to avg expression matrix

## Usage

```
seurat_ref(seurat_object, ...)

## S3 method for class 'Seurat'
seurat_ref(
  seurat_object,
  cluster_col = "classified",
  var_genes_only = FALSE,
  assay_name = NULL,
  method = "mean",
  subclusterpower = 0,
  if_log = TRUE,
  ...
)
```

## Arguments

|                 |                                                                                                  |
|-----------------|--------------------------------------------------------------------------------------------------|
| seurat_object   | seurat_object after tsne or umap projections and clustering                                      |
| ...             | additional arguments                                                                             |
| cluster_col     | column name where classified cluster names are stored in seurat meta data, cannot be "rn"        |
| var_genes_only  | whether to keep only var_genes in the final matrix output, could also look up genes used for PCA |
| assay_name      | any additional assay data, such as ADT, to include. If more than 1, pass a vector of names       |
| method          | whether to take mean (default) or median                                                         |
| subclusterpower | whether to get multiple averages per original cluster                                            |
| if_log          | input data is natural log, averaging will be done on unlogged data                               |

## Value

reference expression matrix, with genes as row names, and cell types as column names

## Examples

```
so <- so_pbmc()
ref <- seurat_ref(so, cluster_col = "seurat_clusters")
```

---

|         |                                 |
|---------|---------------------------------|
| so_pbmc | <i>An example Seurat object</i> |
|---------|---------------------------------|

---

**Description**

An example Seurat object

**Usage**

```
so_pbmc()
```

**Value**

a Seurat object populated with data from the [pbmc\\_matrix\\_small](#) scRNA-seq dataset, additionally annotated with cluster assignments.

---

|                   |                                               |
|-------------------|-----------------------------------------------|
| vector_similarity | <i>Compute similarity between two vectors</i> |
|-------------------|-----------------------------------------------|

---

**Description**

Compute the similarity score between two vectors using a customized scoring function Two vectors may be from either scRNA-seq or bulk RNA-seq data. The lengths of vec1 and vec2 must match, and must be arranged in the same order of genes. Both vectors should be provided to this function after pre-processing, feature selection and dimension reduction.

**Usage**

```
vector_similarity(vec1, vec2, compute_method, ...)
```

**Arguments**

|                |                                              |
|----------------|----------------------------------------------|
| vec1           | test vector                                  |
| vec2           | reference vector                             |
| compute_method | method to run i.e. corr_coef                 |
| ...            | arguments to pass to compute_method function |

**Value**

numeric value of desired correlation or distance measurement

---

|            |                                             |
|------------|---------------------------------------------|
| write_meta | <i>Function to write metadata to object</i> |
|------------|---------------------------------------------|

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**Description**

Function to write metadata to object

**Usage**

```
write_meta(object, ...)  
  
## S3 method for class 'Seurat'  
write_meta(object, meta, ...)  
  
## S3 method for class 'SingleCellExperiment'  
write_meta(object, meta, ...)
```

**Arguments**

|        |                                                      |
|--------|------------------------------------------------------|
| object | object after tsne or umap projections and clustering |
| ...    | additional arguments                                 |
| meta   | new metadata dataframe                               |

**Value**

object with newly inserted metadata columns

**Examples**

```
so <- so_pbmc()  
obj <- write_meta(  
  object = so,  
  meta = seurat_meta(so)  
)  
sce <- sce_pbmc()  
obj <- write_meta(  
  object = sce,  
  meta = object_data(sce, "meta.data")  
)
```

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