

# Package ‘ASURAT’

May 8, 2024

**Type** Package

**Title** Functional annotation-driven unsupervised clustering for single-cell data

**Version** 1.9.0

**Description** ASURAT is a software for single-cell data analysis. Using ASURAT, one can simultaneously perform unsupervised clustering and biological interpretation in terms of cell type, disease, biological process, and signaling pathway activity. Inputting a single-cell RNA-seq data and knowledge-based databases, such as Cell Ontology, Gene Ontology, KEGG, etc., ASURAT transforms gene expression tables into original multivariate tables, termed sign-by-sample matrices (SSMs).

**License** GPL-3 + file LICENSE

**biocViews** GeneExpression, SingleCell, Sequencing, Clustering, GeneSignaling

**VignetteBuilder** knitr

**Encoding** UTF-8

**LazyData** TRUE

**Depends** R (>= 4.0.0)

**Imports** SingleCellExperiment, SummarizedExperiment, S4Vectors, Rcpp (>= 1.0.7), cluster, utils, plot3D, ComplexHeatmap, circlize, grid, grDevices, graphics

**Suggests** ggplot2, TENxPBMCDData, dplyr, Rtsne, Seurat, AnnotationDbi, BiocGenerics, stringr, org.Hs.eg.db, knitr, rmarkdown, testthat (>= 3.0.0)

**RoxygenNote** 7.1.2

**LinkingTo** Rcpp

**Config/testthat/edition** 3

**git\_url** <https://git.bioconductor.org/packages/ASURAT>

**git\_branch** devel

**git\_last\_commit** 8cc0b56

**git\_last\_commit\_date** 2024-04-30

**Repository** Bioconductor 3.20

**Date/Publication** 2024-05-08

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---

add_metadata	<i>Add metadata of variables and samples.</i>
--------------	---

---

### Description

This function adds metadata of variables and samples.

### Usage

```
add_metadata(sce = NULL, mitochondria_symbol = NULL)
```

**Arguments**

`sce` A SingleCellExperiment object.

`mitochondria_symbol` A string representing for mitochondrial genes. This function computes percents of reads that map to the mitochondrial genes. Examples are `'^MT-'`, `'^mt-'`, etc.

**Value**

A SingleCellExperiment object.

**Examples**

```
data(pbmc_eg)
pbmc <- add_metadata(sce = pbmc_eg, mitochondria_symbol = "^MT-")
```

---

ASURAT	<i>Functional annotation-driven unsupervised clustering of SingleCell data.</i>
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---

**Description**

ASURAT is a software for single-cell data analysis. Using ASURAT, one can simultaneously perform unsupervised clustering and biological interpretation in terms of cell type, disease, biological process, and signaling pathway activity. Inputting a single-cell RNA-seq data and knowledge-based databases, such as Cell Ontology, Gene Ontology, KEGG, etc., ASURAT transforms gene expression tables into original multivariate tables, termed sign-by-sample matrices (SSMs).

---

<code>bubble_sort</code>	<i>Perform bubble sorting, counting the number of steps.</i>
--------------------------	--

---

**Description**

Perform bubble sorting, counting the number of steps.

**Usage**

```
bubble_sort(listdata)
```

**Arguments**

`listdata` A list of vector and integer. For example, in R code, `listdata = list(vec = c(1, 0, 1, ...), cnt = 0)`. The integer (`cnt = 0`) is the initial number of steps for bubble sorting.

**Value**

A List.

**Examples**

```
bubble_sort(list(vec = c(1, 1, 0), cnt = 0))
```

---

cluster_genesets	<i>Cluster each functional gene set into three groups.</i>
------------------	--

---

**Description**

This function clusters each functional gene set into strongly, variably, and weakly correlated gene sets.

**Usage**

```
cluster_genesets(sce = NULL, cormat = NULL, th_posi = NULL, th_neg = NULL)
```

**Arguments**

sce	A SingleCellExperiment object.
cormat	A correlation matrix of gene expressions.
th_posi	A threshold of positive correlation coefficient.
th_neg	A threshold of negative correlation coefficient.

**Value**

A SingleCellExperiment object.

**Examples**

```
data(pbmc_eg)
data(human_GO_eg)
mat <- t(as.matrix(SummarizedExperiment::assay(pbmc_eg, "centered")))
pbmc_cormat <- cor(mat, method = "spearman")
pbmcs <- list(GO = pbmc_eg)
S4Vectors::metadata(pbmcs$GO) <- list(sign = human_GO_eg[["BP"]])
pbmcs$GO <- remove_signs(sce = pbmcs$GO, min_genes = 2, max_genes = 1000)
pbmcs$GO <- cluster_genesets(sce = pbmcs$GO, cormat = pbmc_cormat,
                             th_posi = 0.24, th_neg = -0.20)
# The results are stored in `metadata(pbmcs$GO)$sign`.
```

---

compute\_sepI\_all      *Compute separation indices for each cluster against the others.*

---

### Description

This function computes separation indices for each cluster versus the others.

### Usage

```
compute_sepI_all(sce = NULL, labels = NULL, nrand_samples = NULL)
```

### Arguments

sce                    A SingleCellExperiment object.  
 labels                A vector of labels of all the samples (cells).  
 nrand\_samples        An integer for the number of samples used for random sampling, which samples at least one sample per cluster.

### Value

A SingleCellExperiment object.

### Examples

```
data(pbmcs_eg)
labels <- SummarizedExperiment::colData(pbmcs_eg$GO)$seurat_clusters
pbmcs_eg$GO <- compute_sepI_all(sce = pbmcs_eg$GO, labels = labels,
                               nrand_samples = 10)
# The results are stored in `metadata(pbmcs_eg$GO)$marker_signs`.
```

---

compute\_sepI\_clusters      *Compute separation indices of sign scores for given two clusters.*

---

### Description

This function computes separation indices of sign scores for given two clusters.

### Usage

```
compute_sepI_clusters(
  sce = NULL,
  labels = NULL,
  nrand_samples = NULL,
  ident_1 = NULL,
  ident_2 = NULL
)
```

**Arguments**

sce	A SingleCellExperiment object.
labels	A vector of labels of all the samples.
nrand_samples	An integer for the number of samples used for random sampling, which samples at least one sample per cluster.
ident_1	Label names identifying cluster numbers, e.g., <code>ident_1 = 1</code> , <code>ident_1 = c(1, 3)</code> .
ident_2	Label names identifying cluster numbers, e.g., <code>ident_2 = 2</code> , <code>ident_2 = c(2, 4)</code> .

**Value**

A SingleCellExperiment object.

**Examples**

```
data(pbmcs_eg)
labels <- SummarizedExperiment::colData(pbmcs_eg$G0)$seurat_clusters
pbmcs_eg$G0 <- compute_sepI_clusters(sce = pbmcs_eg$G0, labels = labels,
                                   nrand_samples = 10, ident_1 = 1,
                                   ident_2 = c(0, 2))
# The results are stored in `metadata(pbmcs_eg$G0)$marker_signs`.
```

---

create\_signs

*Define signs for strongly and variably correlated gene sets.*

---

**Description**

This function define signs for strongly and variably correlated gene sets.

**Usage**

```
create_signs(sce = NULL, min_cnt_strg = 2, min_cnt_vari = 2)
```

**Arguments**

sce	A SingleCellExperiment object.
min_cnt_strg	An integer for the cutoff value for strongly correlated gene sets.
min_cnt_vari	An integer for the cutoff value for variably correlated gene sets.

**Value**

A SingleCellExperiment object.

**Examples**

```

data(pbmc_eg)
data(human_GO_eg)
mat <- t(as.matrix(SummarizedExperiment::assay(pbmc_eg, "centered")))
pbmc_cormat <- cor(mat, method = "spearman")
pbmcs <- list(GO = pbmc_eg)
S4Vectors::metadata(pbmcs$GO) <- list(sign = human_GO_eg[["BP"]])
pbmcs$GO <- remove_signs(sce = pbmcs$GO, min_ngenes = 2, max_ngenes = 1000)
pbmcs$GO <- cluster_genesets(sce = pbmcs$GO, cormat = pbmc_cormat,
                             th_posi = 0.24, th_negs = -0.20)
pbmcs$GO <- create_signs(sce = pbmcs$GO, min_cnt_strg = 2, min_cnt_vari = 2)
# The results are stored in `metadata(pbmcs$GO)$sign_all`.

```

---

human_COMSig_eg	<i>A list of small Cell Ontology and MSigDB databases for human.</i>
-----------------	--

---

**Description**

A list of small Cell Ontology and MSigDB databases for human.

**Usage**

```
human_COMSig_eg
```

**Format**

A list of dataframe.

---

human_GO_eg	<i>A list of small Gene Ontology database for human.</i>
-------------	--

---

**Description**

A list of small Gene Ontology database for human.

**Usage**

```
human_GO_eg
```

**Format**

A list of dataframe.

---

human_KEGG_eg	<i>A list of small KEGG database for human.</i>
---------------	---

---

**Description**

A list of small KEGG database for human.

**Usage**

```
human_KEGG_eg
```

**Format**

A list of dataframe.

---

makeSignMatrix	<i>Create a new SingleCellExperiment object for sign-by-sample matrices.</i>
----------------	--

---

**Description**

This function creates a new `SingleCellExperiment` object for sign-by-sample matrices (SSM) by concatenating SSMs for strongly and variably correlated gene sets.

**Usage**

```
makeSignMatrix(sce = NULL, weight_strg = 0.5, weight_vari = 0.5)
```

**Arguments**

sce	A <code>SingleCellExperiment</code> object.
weight_strg	A weight parameter for strongly correlated gene sets.
weight_vari	A weight parameter for variably correlated gene sets.

**Value**

A `SingleCellExperiment` object.

**Examples**

```

data(pbmc_eg)
data(human_GO_eg)
mat <- t(as.matrix(SummarizedExperiment::assay(pbmc_eg, "centered")))
pbmc_cormat <- cor(mat, method = "spearman")
pbmcs <- list(GO = pbmc_eg)
S4Vectors::metadata(pbmcs$GO) <- list(sign = human_GO_eg[["BP"]])
pbmcs$GO <- remove_signs(sce = pbmcs$GO, min_ngenes = 2, max_ngenes = 1000)
pbmcs$GO <- cluster_genesets(sce = pbmcs$GO, cormat = pbmc_cormat,
                             th_posi = 0.24, th_negs = -0.20)
pbmcs$GO <- create_signs(sce = pbmcs$GO, min_cnt_strg = 2, min_cnt_vari = 2)
pbmcs$GO <- makeSignMatrix(sce = pbmcs$GO, weight_strg = 0.5,
                           weight_vari = 0.5)
# The results can be check by, e.g., assay(pbmcs$GO, "counts").

```

---

pbmcs_eg	<i>A list of SingleCellExperiment objects made from sign-sample matrices.</i>
----------	---

---

**Description**

A list of SingleCellExperiment objects, consisting of small sign-by-sample matrices, pbmcs\_eg\$CM (using Cell Ontology and MSigDB databases), pbmcs\_eg\$GO (using Gene Ontology database), and pbmcs\_eg\$KG (KEGG). Here, pbmcs\_eg\$CM, pbmcs\_eg\$GO, and pbmcs\_eg\$KG include 87, 72, and 64 signs, respectively, and 50 cells.

**Usage**

```
pbmcs_eg
```

**Format**

A list of SingleCellExperiment objects.

---

pbmc_eg	<i>A SingleCellExperiment object made from a gene expression table.</i>
---------	---

---

**Description**

A SingleCellExperiment object, including 50 genes and 50 cells. The original data "4k PBMCs from a Healthy Donor" was downloaded from 10x Genomics database.

**Usage**

```
pbmc_eg
```

**Format**

SingleCellExperiment object.

**Source**

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

---

plot_dataframe3D	<i>Visualize a three-dimensional data with labels and colors.</i>
------------------	---

---

**Description**

This function visualizes a three-dimensional data with labels and colors.

**Usage**

```
plot_dataframe3D(  
  dataframe3D = NULL,  
  labels = NULL,  
  colors = NULL,  
  theta = 30,  
  phi = 30,  
  title = "",  
  xlabel = "",  
  ylabel = "",  
  zlabel = ""  
)
```

**Arguments**

dataframe3D	A dataframe with three columns.
labels	NULL or a vector of labels of all the samples, corresponding to colors.
colors	NULL or a vector of colors of all the samples, corresponding to labels.
theta	Angle of the plot.
phi	Angle of the plot.
title	Title.
xlabel	x-axis label.
ylabel	y-axis label.
zlabel	z-axis label.

**Value**

A scatter3D object in plot3D package.

**Examples**

```

data(pbmcs_eg)
mat <- SingleCellExperiment::reducedDim(pbmcs_eg$CM, "UMAP")[, 1:3]
dataframe3D <- as.data.frame(mat)
labels <- SummarizedExperiment::colData(pbmcs_eg$CM)$seurat_clusters
plot_dataframe3D(dataframe3D = dataframe3D, labels = labels, colors = NULL,
                 theta = 45, phi = 20, title = "PBMC (CO & MSigDB)",
                 xlabel = "UMAP_1", ylabel = "UMAP_2", zlabel = "UMAP_3")

```

---

plot\_multiheatmaps      *Visualize multivariate data by heatmaps.*

---

**Description**

This function visualizes multivariate data by heatmaps.

**Usage**

```

plot_multiheatmaps(
  ssm_list = NULL,
  gem_list = NULL,
  ssmlabel_list = NULL,
  gemlabel_list = NULL,
  nrand_samples = NULL,
  show_row_names = FALSE,
  title = NULL
)

```

**Arguments**

ssm_list	A list of sign-by-sample matrices.
gem_list	A list of gene-by-sample matrices.
ssmlabel_list	NULL or a list of dataframes of sample (cell) labels and colors. The length of the list must be as same as that of ssm_list, and the order of labels in each list must be as same as those in ssm_list.
gemlabel_list	NULL or a list of dataframes of sample (cell) annotations and colors. The length of the list must be as same as that of gem_list, and the order of labels in each list must be as same as those in gem_list.
nrand_samples	Number of samples (cells) used for random sampling.
show_row_names	TRUE or FALSE: if TRUE, row names are shown.
title	Title.

**Value**

A ComplexHeatmap object.

## Examples

```

data(pbmcs_eg)
mat_CM <- SummarizedExperiment::assay(pbmcs_eg$CM, "counts")
mat_GO <- SummarizedExperiment::assay(pbmcs_eg$GO, "counts")
mat_KG <- SummarizedExperiment::assay(pbmcs_eg$KG, "counts")
ssm_list <- list(SSM_COMSig = mat_CM, SSM_GO = mat_GO, SSM_KEGG = mat_KG)
se <- SingleCellExperiment::altExp(pbmcs_eg$CM, "logcounts")
mat <- SummarizedExperiment::assay(se, "counts")
se <- SingleCellExperiment::altExp(pbmcs_eg$CM, "logcounts")
gem_list <- list(GeneExpr = SummarizedExperiment::assay(se, "counts"))
labels <- list() ; ssmlabel_list <- list()
for(i in seq_along(pbmcs_eg)){
  fa <- SummarizedExperiment::colData(pbmcs_eg[[i]])$seurat_clusters
  labels[[i]] <- data.frame(label = fa)
  colors <- rainbow(length(unique(labels[[i]]$label)))[labels[[i]]$label]
  labels[[i]]$color <- colors
  ssmlabel_list[[i]] <- labels[[i]]
}
names(ssmlabel_list) <- c("Label_COMSig", "Label_GO", "Label_KEGG")
phases <- SummarizedExperiment::colData(pbmcs_eg$CM)$Phase
label_CC <- data.frame(label = phases, color = NA)
gemlabel_list <- list(CellCycle = label_CC)
plot_multiheatmaps(ssm_list = ssm_list, gem_list = gem_list,
  ssmlabel_list = ssmlabel_list,
  gemlabel_list = gemlabel_list, nrand_samples = 50,
  show_row_names = FALSE, title = "PBMC")

```

---

remove\_samples

*Remove samples based on expression profiles across variables.*

---

## Description

This function removes sample data by setting minimum and maximum threshold values for the metadata.

## Usage

```

remove_samples(
  sce = NULL,
  min_nReads = NULL,
  max_nReads = NULL,
  min_nGenes = NULL,
  max_nGenes = NULL,
  min_percMT = NULL,
  max_percMT = NULL
)

```

**Arguments**

sce	A SingleCellExperiment object.
min_nReads	A minimum threshold value of the number of reads.
max_nReads	A maximum threshold value of the number of reads.
min_nGenes	A minimum threshold value of the number of non-zero expressed genes.
max_nGenes	A maximum threshold value of the number of non-zero expressed genes.
min_percMT	A minimum threshold value of the percent of reads that map to mitochondrial genes.
max_percMT	A maximum threshold value of the percent of reads that map to mitochondrial genes.

**Value**

A SingleCellExperiment object.

**Examples**

```
data(pbmc_eg)
pbmc <- add_metadata(sce = pbmc_eg, mitochondria_symbol = "^MT-")
pbmc <- remove_samples(sce = pbmc, min_nReads = 0, max_nReads = 1e+10,
                      min_nGenes = 0, max_nGenes = 1e+10,
                      min_percMT = NULL, max_percMT = NULL)
```

---

remove\_signs

*Remove signs including too few or too many genes.*

---

**Description**

This function removes signs including too few or too many genes.

**Usage**

```
remove_signs(sce = NULL, min_ngenes = 2, max_ngenes = 1000)
```

**Arguments**

sce	A SingleCellExperiment object.
min_ngenes	Minimum number of genes, which must be greater than one.
max_ngenes	Maximum number of genes, which must be greater than one.

**Value**

A SingleCellExperiment object.

**Examples**

```

data(pbmc_eg)
data(human_GO_eg)
pbmcs <- list(GO = pbmc_eg)
S4Vectors::metadata(pbmcs$GO) <- list(sign = human_GO_eg[["BP"]])
pbmcs$GO <- remove_signs(sce = pbmcs$GO, min_ngenes = 2, max_ngenes = 1000)
# The results are stored in `metadata(pbmcs$GO)$sign`.

```

---

remove\_signs\_manually *Remove signs by specifying keywords.*

---

**Description**

This function removes signs by specifying keywords.

**Usage**

```
remove_signs_manually(sce = NULL, keywords = NULL)
```

**Arguments**

sce	A SingleCellExperiment object.
keywords	keywords separated by pipes ' '. 

**Value**

A SingleCellExperiment object.

**Examples**

```

data(pbmc_eg)
data(human_GO_eg)
mat <- t(as.matrix(SummarizedExperiment::assay(pbmc_eg, "centered")))
pbmc_cormat <- cor(mat, method = "spearman")
pbmcs <- list(GO = pbmc_eg)
S4Vectors::metadata(pbmcs$GO) <- list(sign = human_GO_eg[["BP"]])
pbmcs$GO <- remove_signs(sce = pbmcs$GO, min_ngenes = 2, max_ngenes = 1000)
pbmcs$GO <- cluster_genesets(sce = pbmcs$GO, cormat = pbmc_cormat,
                             th_posi = 0.24, th_neg = -0.20)
pbmcs$GO <- create_signs(sce = pbmcs$GO, min_cnt_strg = 2, min_cnt_vari = 2)
keywords <- "Covid19|foofoo|hogegege"
pbmcs$GO <- remove_signs_manually(sce = pbmcs$GO, keywords = keywords)
# The results are stored in `metadata(pbmcs$GO)$sign_SCG`,
# `metadata(pbmcs$GO)$sign_VCG`, and `metadata(pbmcs$GO)$sign_all`.

```

---

 remove\_signs\_redundant

*Remove redundant signs using semantic similarity matrices.*


---

## Description

This function removes redundant signs using semantic similarity matrices.

## Usage

```
remove_signs_redundant(
  sce = NULL,
  similarity_matrix = NULL,
  threshold = NULL,
  keep_rareID = NULL
)
```

## Arguments

sce	A SingleCellExperiment object.
similarity_matrix	A semantic similarity matrix.
threshold	A threshold value of semantic similarity, used for regarding biological terms as similar ones
keep_rareID	If TRUE, biological terms with the larger ICs are kept.

## Value

A SingleCellExperiment object.

## Examples

```
data(pbmc_eg)
data(human_GO_eg)
mat <- t(as.matrix(SummarizedExperiment::assay(pbmc_eg, "centered")))
pbmc_cormat <- cor(mat, method = "spearman")
pbmcs <- list(GO = pbmc_eg)
S4Vectors::metadata(pbmcs$GO) <- list(sign = human_GO_eg[["BP"]])
pbmcs$GO <- remove_signs(sce = pbmcs$GO, min_ngenes = 2, max_ngenes = 1000)
pbmcs$GO <- cluster_genesets(sce = pbmcs$GO, cormat = pbmc_cormat,
  th_posi = 0.24, th_neg = -0.20)
pbmcs$GO <- create_signs(sce = pbmcs$GO, min_cnt_strg = 2, min_cnt_vari = 2)
pbmcs$GO <- remove_signs_redundant(
  sce = pbmcs$GO, similarity_matrix = human_GO_eg$similarity_matrix$BP,
  threshold = 0.80, keep_rareID = TRUE)
# The results are stored in `metadata(pbmcs$GO)$sign_SCG`,
# `metadata(pbmcs$GO)$sign_VCG`, `metadata(pbmcs$GO)$sign_all`,
# and if there exist, `metadata(pbmcs$GO)$sign_SCG_redundant` and
```

```
# `metadata(pbmc$GO)$sign_VCG_redundant`.
```

---

remove\_variables      *Remove variables based on expression profiles across samples.*

---

### Description

This function removes low expressed variable data.

### Usage

```
remove_variables(sce = NULL, min_nsamples = 0)
```

### Arguments

sce                    A SingleCellExperiment object.

min\_nsamples        An integer. This function removes variables for which the numbers of non-zero expressing samples are less than this value.

### Value

A SingleCellExperiment object.

### Examples

```
data(pbmc_eg)
pbmc <- add_metadata(sce = pbmc_eg, mitochondria_symbol = "^MT-")
pbmc <- remove_variables(sce = pbmc, min_nsamples = 10)
```

---

remove\_variables\_second      *Remove variables based on the mean expression levels across samples.*

---

### Description

This function removes variable data such that the mean expression levels across samples are less than 'min\_meannReads'.

### Usage

```
remove_variables_second(sce = NULL, min_meannReads = 0)
```

**Arguments**

sce                    A SingleCellExperiment object.

min\_meannReads    An integer. This function removes variables for which the mean read counts are less than this value.

**Value**

A SingleCellExperiment object.

**Examples**

```
data(pbmc_eg)
pbmc <- remove_variables_second(sce = pbmc_eg, min_meannReads = 0.01)
```

---

select\_signs\_manually    *Select signs by specifying keywords.*

---

**Description**

This function selects signs by specifying keywords.

**Usage**

```
select_signs_manually(sce = NULL, keywords = NULL)
```

**Arguments**

sce                    An ASURAT object.

keywords              Keywords separated by a pipe.

**Value**

An ASURAT object.

**Examples**

```
data(pbmc_eg)
data(human_GO_eg)
mat <- t(as.matrix(SummarizedExperiment::assay(pbmc_eg, "centered")))
pbmc_cormat <- cor(mat, method = "spearman")
pbmcs <- list(GO = pbmc_eg)
S4Vectors::metadata(pbmcs$GO) <- list(sign = human_GO_eg[["BP"]])
pbmcs$GO <- remove_signs(sce = pbmcs$GO, min_ngenes = 2, max_ngenes = 1000)
pbmcs$GO <- cluster_genesets(sce = pbmcs$GO, cormat = pbmc_cormat,
                             th_posi = 0.24, th_negs = -0.20)
pbmcs$GO <- create_signs(sce = pbmcs$GO, min_cnt_strg = 2, min_cnt_vari = 2)
keywords <- "cell|process"
```

```
pbmcs$GO <- select_signs_manually(sce = pbmcs$GO, keywords = keywords)
# The results are stored in `metadata(pbmcs$GO)$sign_SCG`,
# `metadata(pbmcs$GO)$sign_VCG`, and `metadata(pbmcs$GO)$sign_all`.
```

---

swap\_pass

*Perform one-shot adjacent swapping for each element.*

---

### **Description**

Perform one-shot adjacent swapping for each element.

### **Usage**

```
swap_pass(listdata)
```

### **Arguments**

listdata      A list of vector and integer.

### **Value**

A List.

### **Examples**

```
swap_pass(list(vec = c(1, 1, 0), cnt = 0))
```

# Index

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