

# Package ‘Coralysis’

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**Type** Package

**Title** Coralysis sensitive identification of imbalanced cell types and states in single-cell data via multi-level integration

**Version** 1.3.0

## Description

Coralysis is an R package featuring a multi-level integration algorithm for sensitive integration, reference-mapping, and cell-state identification in single-cell data. The multi-level integration algorithm is inspired by the process of assembling a puzzle - where one begins by grouping pieces based on low-to high-level features, such as color and shading, before looking into shape and patterns. This approach progressively blends the batch effects and separates cell types across multiple rounds of divisive clustering.

**License** GPL-3

**Imports** Matrix, aricode, LiblineaR, SparseM, ggplot2, umap, Rtsne, pheatmap, reshape2, dplyr, SingleCellExperiment, SummarizedExperiment, S4Vectors, methods, stats, utils, RANN, sparseMatrixStats, irlba, flexclust, scran, class, matrixStats, tidyr, cowplot, uwot, scatterpie, RColorBrewer, ggrastr, ggrepel, RSpectra, BiocParallel, withr

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<https://elolab.github.io/Coralysis/>

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---

<code>.randomColors</code>	<i>Random colors</i>
----------------------------	----------------------

---

**Description**

The function returns a group of random colors.

**Usage**

`.randomColors(ncolors)`

**Arguments**

`ncolors`           Integer. Number of colors to generate randomly.

**Value**

Random colors.

---

<code>AggregateClusterExpression</code>	<i>Aggregates cell feature expression by clusters</i>
---	---

---

**Description**

The function aggregates cell feature expression by clusters provided.

**Usage**

`AggregateClusterExpression(mtx, cluster, select.features = NULL, fun = "mean")`

**Arguments**

`mtx`                 Matrix with features vs cells (rows vs cols) with feature expression to aggregate.  
`cluster`            Cluster identities vector corresponding to the cells in `mtx`.  
`select.features`    Should features be selected. By default NULL, all features used.  
`fun`                 Character specifying if feature expression should be aggregated by mean or sum.  
                      By default "mean".

**Value**

Matrix of feature expressed aggregated by clusters.

---

AggregateDataByBatch *Aggregates feature expression by cell clusters, per batch if provided.*

---

### Description

The function aggregates feature expression by cell clusters, per batch if provided.

### Usage

```
AggregateDataByBatch.SingleCellExperiment(object, batch.label, nhvg, p, ...)
```

```
## S4 method for signature 'SingleCellExperiment'
```

```
AggregateDataByBatch(object, batch.label, nhvg = 2000L, p = 30L, ...)
```

### Arguments

object	An object of SingleCellExperiment class.
batch.label	Cluster identities vector corresponding to the cells in mtx.
nhvg	Integer of the number of highly variable features to select. By default 2000.
p	Integer. By default 30.
...	Parameters to be passed to ClusterCells() function.

### Value

A SingleCellExperiment object with feature expression aggregated by clusters.

### Examples

```
# Import package
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Import data from Zenodo
data.url <- "https://zenodo.org/records/14871436/files/pbmc_10Xassays.rds?download=1"
sce <- readRDS(file = url(data.url))

# Run with a batch
set.seed(1204)
sce <- AggregateDataByBatch(object = sce, batch.label = "batch")
logcounts(sce)[1:10, 1:10]
head(metadata(sce)$clusters)

# Run without a batch
set.seed(1204)
sce <- AggregateDataByBatch(object = sce, batch.label = NULL)
logcounts(sce)[1:10, 1:10]
head(metadata(sce)$clusters)
```

---

```
BinCellClusterProbability
    Bin cell cluster probability
```

---

### Description

Bin cell cluster probability by a given cell label.

### Usage

```
BinCellClusterProbability.SingleCellExperiment(
  object,
  label,
  icp.run,
  icp.round,
  funs,
  bins,
  aggregate.bins.by,
  use.assay
)

## S4 method for signature 'SingleCellExperiment'
BinCellClusterProbability(
  object,
  label,
  icp.run = NULL,
  icp.round = NULL,
  funs = "mean",
  bins = 20,
  aggregate.bins.by = "mean",
  use.assay = "logcounts"
)
```

### Arguments

object	An object of SingleCellExperiment class with ICP cell cluster probability tables saved in <code>metadata(object)\$coranalysis\$joint.probability</code> . After running one of <code>RunParallelICP</code> or <code>RunParallelDivisiveICP</code> .
label	Label of interest available in <code>colData(object)</code> to group by the bins of cell cluster probability.
icp.run	ICP run(s) to retrieve from <code>metadata(object)\$coranalysis\$joint.probability</code> . By default NULL, i.e., all are retrieved. Specify a numeric vector to retrieve a specific set of tables.
icp.round	ICP round(s) to retrieve from <code>metadata(object)\$coranalysis\$joint.probability</code> . By default NULL, i.e., all are retrieved. Only relevant if probabilities were obtained with the function <code>RunParallelDivisiveICP</code> , i.e., divisive ICP was performed. Otherwise it is ignored and internally assumed as <code>icp.round = 1</code> , i.e., only one round.
funs	One function to summarise ICP cell cluster probability. One of "mean" or "median". By default "mean".

bins                    Number of bins to bin cell cluster probability by cell label given. By default 20.

aggregate.bins.by        One function to aggregate One of "mean" or "median". By default "mean".

use.assay                Name of the assay that should be used to obtain the average expression of features across cell label probability bins.

### Value

A SingleCellExperiment class object with feature average expression by cell label probability bins.

### Examples

```
# Packages
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Import data from Zenodo
data.url <- "https://zenodo.org/records/14845751/files/pbmc_10Xassays.rds?download=1"
sce <- readRDS(file = url(data.url))

# Prepare data
sce <- PrepareData(object = sce)

# Multi-level integration - 'L = 4' just for highlighting purposes
set.seed(123)
sce <- RunParallelDivisiveICP(
  object = sce, batch.label = "batch", L = 4,
  threads = 2
)

# Cell states SCE object for a given cell type annotation or clustering
cellstate.sce <- BinCellClusterProbability(
  object = sce, label = "cell_type",
  icp.round = 4, bins = 20
)
cellstate.sce
```

---

CellBinsFeatureCorrelation

*Cell bins feature correlation*

---

### Description

Correlation between cell bins for the given labels and features.

### Usage

```
CellBinsFeatureCorrelation.SingleCellExperiment(object, labels, method)

## S4 method for signature 'SingleCellExperiment'
CellBinsFeatureCorrelation(object, labels = NULL, method = "pearson")
```

**Arguments**

object	An object of SingleCellExperiment class obtained with the function BinCellClusterProbabilityDistribution().
labels	Character of label(s) from the label provided to the function BinCellClusterProbabilityDistribution(). By default NULL, i.e., all labels are used.
method	Character specifying the correlation method to use. One of "pearson", "kendall" or "spearman". By default "pearson" is used.

**Value**

A data frame with the correlation coefficient for each feature (rows) across labels (columns).

**Examples**

```
# Packages
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Import data from Zenodo
data.url <- "https://zenodo.org/records/14845751/files/pbmc_10Xassays.rds?download=1"
sce <- readRDS(file = url(data.url))

# Prepare data
sce <- PrepareData(object = sce)

# Multi-level integration - 'L = 4' just for highlighting purposes
set.seed(123)
sce <- RunParallelDivisiveICP(
  object = sce, batch.label = "batch", L = 4,
  threads = 2
)

# Cell states SCE object for a given cell type annotation or clustering
cellstate.sce <- BinCellClusterProbability(
  object = sce, label = "cell_type",
  icp.round = 4, bins = 20
)
cellstate.sce

# Pearson correlated features with "Monocyte"
cor.features.mono <- CellBinsFeatureCorrelation(
  object = cellstate.sce,
  labels = "Monocyte"
)
```

---

CellClusterProbabilityDistribution

*Cell cluster probability distribution*

---

**Description**

Plot cell cluster probability distribution per label by group.

**Usage**

```

CellClusterProbabilityDistribution.SingleCellExperiment(
  object,
  label,
  group,
  probability
)

## S4 method for signature 'SingleCellExperiment'
CellClusterProbabilityDistribution(
  object,
  label,
  group,
  probability = "scaled_mean_probs"
)

```

**Arguments**

<code>object</code>	An object of <code>SingleCellExperiment</code> class with aggregated cell cluster probability available in <code>colData(object)</code> , which can be obtained after running <code>SummariseCellClusterProbability()</code> .
<code>label</code>	Character specifying the <code>colData</code> variable to use as cell type/cluster label.
<code>group</code>	Character specifying the <code>colData</code> variable to use as categorical group variable.
<code>probability</code>	Character specifying the aggregated cell cluster probability variable available in <code>colData</code> , used to plot its distribution. One of "mean_probs", "scaled_mean_probs", "median_probs", "scaled_median_probs". The availability of these variables in <code>colData</code> depends on the parameters given to the function <code>SummariseCellClusterProbability()</code> beforehand. By default assumes that "scaled_mean_probs" is available in <code>colData</code> , which is only true if <code>SummariseCellClusterProbability()</code> function was run with <code>funs = "mean"</code> and <code>scale.funs = TRUE</code> .

**Value**

A plot of class `ggplot`.

**Examples**

```

# Import package
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Create toy SCE data
batches <- c("b1", "b2")
set.seed(239)
batch <- sample(x = batches, size = nrow(iris), replace = TRUE)
sce <- SingleCellExperiment(
  assays = list(logcounts = t(iris[, 1:4])),
  colData = DataFrame(
    "Species" = iris$Species,
    "Batch" = batch
  )
)
colnames(sce) <- paste0("samp", 1:ncol(sce))

```

```

# Prepare SCE object for analysis
sce <- PrepareData(sce)

# Multi-level integration (just for highlighting purposes; use default parameters)
set.seed(123)
sce <- RunParallelDivisiveICP(
  object = sce, batch.label = "Batch",
  k = 4, L = 25, C = 1, d = 0.5,
  train.with.bnn = FALSE,
  use.cluster.seed = FALSE,
  build.train.set = FALSE, ari.cutoff = 0.1,
  threads = 2, RNGseed = 1024
)

# Summarise cell cluster probability
sce <- SummariseCellClusterProbability(object = sce, icp.round = 2) # saved in 'colData'

# Search for differences in probabilities across group(s)
# give an interesting variable to the "group" parameter
prob.dist <- CellClusterProbabilityDistribution(
  object = sce, label = "Species",
  group = "Batch",
  probability = "scaled_mean_probs"
)
prob.dist # print plot

```

---

ClusterCells

*Cluster cells*


---

## Description

The function clusters cells with the K-means++ algorithm

## Usage

```
ClusterCells(object, nclusters = 500, use.emb = TRUE, emb.name = "PCA")
```

## Arguments

object	An object of SingleCellExperiment class.
nclusters	Cluster the cells into n clusters. Ignored if the number of cells in object is lower or equal to nclusters.
use.emb	Should the embedding be used to cluster or the log-transformed data. By default TRUE.
emb.name	Which embedding to use. By default "PCA".

## Value

A SingleCellExperiment object with clusters.

DownOverSampleEvenlyBatches

*Down- and oversample data evenly batches*

---

### **Description**

The function down- and over-samples cluster cells evenly by batch.

### **Usage**

```
DownOverSampleEvenlyBatches(x, batch, n = 50)
```

### **Arguments**

x	A character or numeric vector of data to down-and oversample.
batch	A character vector with batch labels corresponding to x.
n	How many cells to include per cluster.

### **Value**

a list containing the output of the LiblineaR prediction

---

DownOverSampling

*Down- and oversample data*

---

### **Description**

The function implements a script down- and oversamples data to include n cells.

### **Usage**

```
DownOverSampling(x, n = 50)
```

### **Arguments**

x	A character or numeric vector of data to down-and oversample.
n	How many cells to include per cluster.

### **Value**

a list containing the output of the LiblineaR prediction

---

FindAllClusterMarkers *Identification of feature markers for all clusters*

---

## Description

FindAllClusterMarkers enables identifying feature markers for all clusters at once. This is done by differential expression analysis where cells from one cluster are compared against the cells from the rest of the clusters. Feature and cell filters can be applied to accelerate the analysis, but this might lead to missing weak signals.

## Usage

```
FindAllClusterMarkers.SingleCellExperiment(
  object,
  clustering.label,
  test,
  log2fc.threshold,
  min.pct,
  min.diff.pct,
  min.cells.group,
  max.cells.per.cluster,
  return.thresh,
  only.pos
)

## S4 method for signature 'SingleCellExperiment'
FindAllClusterMarkers(
  object,
  clustering.label,
  test = "wilcox",
  log2fc.threshold = 0.25,
  min.pct = 0.1,
  min.diff.pct = NULL,
  min.cells.group = 3,
  max.cells.per.cluster = NULL,
  return.thresh = 0.01,
  only.pos = FALSE
)
```

## Arguments

object	A SingleCellExperiment object.
clustering.label	A variable name (of class character) available in the cell metadata colData(object) with the clustering labels (character or factor) to use.
test	Which test to use. Only "wilcox" (the Wilcoxon rank-sum test, AKA Mann-Whitney U test) is supported at the moment.
log2fc.threshold	Filters out features that have log2 fold-change of the averaged feature expression values below this threshold. Default is 0.25.

<code>min.pct</code>	Filters out features that have dropout rate (fraction of cells expressing a feature) below this threshold in both comparison groups. Default is 0.1.
<code>min.diff.pct</code>	Filters out features that do not have this minimum difference in the dropout rates (fraction of cells expressing a feature) between the two comparison groups. Default is NULL.
<code>min.cells.group</code>	The minimum number of cells in the two comparison groups to perform the DE analysis. If the number of cells is below the threshold, then the DE analysis of this cluster is skipped. Default is 3.
<code>max.cells.per.cluster</code>	The maximum number of cells per cluster if downsampling is performed to speed up the DE analysis. Default is NULL, i.e., no downsampling.
<code>return.thresh</code>	If <code>only.pos=TRUE</code> , then return only features that have the adjusted p-value (adjusted by the Bonferroni method) below or equal to this threshold. Default is 0.01.
<code>only.pos</code>	Whether to return only features that have an adjusted p-value (adjusted by the Bonferroni method) below or equal to the threshold. Default is FALSE.

**Value**

A data frame of the results if positive results were found, else NULL.

**Examples**

```
# Import package
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Create toy SCE data
batches <- c("b1", "b2")
set.seed(239)
batch <- sample(x = batches, size = nrow(iris), replace = TRUE)
sce <- SingleCellExperiment(
  assays = list(logcounts = t(iris[, 1:4])),
  colData = DataFrame(
    "Species" = iris$Species,
    "Batch" = batch
  )
)
colnames(sce) <- paste0("samp", 1:ncol(sce))

# Markers
dge <- FindAllClusterMarkers(sce, clustering.label = "Species")
dge
```

---

FindBatchKNN

*Find batch k nearest neighbors*


---

**Description**

The function finds batch k nearest neighbors for the cell with the highest probability for every batch.

**Usage**

```
FindBatchKNN(idx, group, prob, k = 10)
```

**Arguments**

idx	A numeric vector with the cell ids to retrieve from the data set.
group	A character vector with batch labels corresponding to idx.
prob	A numeric vector with cell probabilities corresponding to idx.
k	The number of nearest neighbors to search for. Default is 10.

**Value**

a list containing the k nearest neighbors for every cell queried

---

FindClusterBatchKNN    *Find batch k nearest neighbors per cluster*

---

**Description**

The function finds batch k nearest neighbors for the cell with the highest probability for every batch per cluster.

**Usage**

```
FindClusterBatchKNN(preds, probs, batch, k = 10, k.prop = NULL)
```

**Arguments**

preds	A numeric vector with the cell cluster predictions.
probs	A numeric matrix with cell cluster probabilities.
batch	A character with batch labels.
k	The number of nearest neighbors to search for. Default is 10.
k.prop	A numeric (higher than 0 and lower than 1) corresponding to the fraction of cells per cluster to use as k nearest neighbors. Default is NULL meaning that the number of k nearest neighbors is equal to k. If given, k parameter is ignored and k is calculated based on k.prop.

**Value**

a list containing the k nearest neighbors for every cluster

---

FindClusterMarkers      *Differential expression between cell clusters*

---

### Description

FindClusterMarkers enables identifying feature markers for one cluster or two arbitrary combinations of clusters, e.g. 1\_2 vs. 3\_4\_5. Feature and cell filters can be applied to accelerate the analysis, but this might lead to missing weak signals.

### Usage

```
FindClusterMarkers.SingleCellExperiment(
  object,
  clustering.label,
  clusters.1,
  clusters.2,
  test,
  log2fc.threshold,
  min.pct,
  min.diff.pct,
  min.cells.group,
  max.cells.per.cluster,
  return.thresh,
  only.pos
)

## S4 method for signature 'SingleCellExperiment'
FindClusterMarkers(
  object,
  clustering.label,
  clusters.1 = NULL,
  clusters.2 = NULL,
  test = "wilcox",
  log2fc.threshold = 0.25,
  min.pct = 0.1,
  min.diff.pct = NULL,
  min.cells.group = 3,
  max.cells.per.cluster = NULL,
  return.thresh = 0.01,
  only.pos = FALSE
)
```

### Arguments

object	A SingleCellExperiment object.
clustering.label	A variable name (of class character) available in the cell metadata colData(object) with the clustering labels (character or factor) to use.
clusters.1	a character or numeric vector denoting which clusters to use in the first group (named group.1 in the results)

<code>clusters.2</code>	a character or numeric vector denoting which clusters to use in the second group (named <code>group.2</code> in the results)
<code>test</code>	Which test to use. Only "wilcoxon" (the Wilcoxon rank-sum test, AKA Mann-Whitney U test) is supported at the moment.
<code>log2fc.threshold</code>	Filters out features that have log2 fold-change of the averaged feature expression values below this threshold. Default is 0.25.
<code>min.pct</code>	Filters out features that have dropout rate (fraction of cells expressing a feature) below this threshold in both comparison groups. Default is 0.1.
<code>min.diff.pct</code>	Filters out features that do not have this minimum difference in the dropout rates (fraction of cells expressing a feature) between the two comparison groups. Default is NULL.
<code>min.cells.group</code>	The minimum number of cells in the two comparison groups to perform the DE analysis. If the number of cells is below the threshold, then the DE analysis is not performed. Default is 3.
<code>max.cells.per.cluster</code>	The maximum number of cells per cluster if downsampling is performed to speed up the DE analysis. Default is NULL, i.e. no downsampling.
<code>return.thresh</code>	If <code>only.pos=TRUE</code> , then return only features that have the adjusted p-value (adjusted by the Bonferroni method) below or equal to this threshold. Default is 0.01.
<code>only.pos</code>	Whether to return only features that have an adjusted p-value (adjusted by the Bonferroni method) below or equal to the threshold. Default is FALSE.

**Value**

a data frame of the results if positive results were found, else NULL

**Examples**

```
# Import package
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Create toy SCE data
batches <- c("b1", "b2")
set.seed(239)
batch <- sample(x = batches, size = nrow(iris), replace = TRUE)
sce <- SingleCellExperiment(
  assays = list(logcounts = t(iris[, 1:4])),
  colData = DataFrame(
    "Species" = iris$Species,
    "Batch" = batch
  )
)
colnames(sce) <- paste0("samp", 1:ncol(sce))

# Markers between versicolor vs virginica
dge <- FindClusterMarkers(sce,
  clustering.label = "Species",
  clusters.1 = "versicolor",
  clusters.2 = "virginica"
)
```

dge

---

GetCellClusterProbability  
*Get ICP cell cluster probability*

---

**Description**

Get ICP cell cluster probability table(s)

**Usage**

```
GetCellClusterProbability.SingleCellExperiment(
  object,
  icp.run,
  icp.round,
  concatenate
)

## S4 method for signature 'SingleCellExperiment'
GetCellClusterProbability(
  object,
  icp.run = NULL,
  icp.round = NULL,
  concatenate = TRUE
)
```

**Arguments**

object	An object of SingleCellExperiment class with ICP cell cluster probability tables saved in <code>metadata(object)\$coranalysis\$joint.probability</code> . After running <code>RunParallelDivisiveICP</code> .
icp.run	ICP run(s) to retrieve from <code>metadata(object)\$coranalysis\$joint.probability</code> . By default NULL, i.e., all are retrieved. Specify a numeric vector to retrieve a specific set of tables.
icp.round	ICP round(s) to retrieve from <code>metadata(object)\$coranalysis\$joint.probability</code> . By default NULL, i.e., all are retrieved.
concatenate	Concatenate list of ICP cell cluster probability tables retrieved. By default TRUE, i.e., the list of ICP cell cluster probability tables is concatenated.

**Value**

A list with ICP cell cluster probability tables or a matrix with concatenated tables.

**Examples**

```

# Import package
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Create toy SCE data
batches <- c("b1", "b2")
set.seed(239)
batch <- sample(x = batches, size = nrow(iris), replace = TRUE)
sce <- SingleCellExperiment(
  assays = list(logcounts = t(iris[, 1:4])),
  colData = DataFrame(
    "Species" = iris$Species,
    "Batch" = batch
  )
)
colnames(sce) <- paste0("samp", 1:ncol(sce))

# Prepare SCE object for analysis
sce <- PrepareData(sce)

# Multi-level integration (just for highlighting purposes; use default parameters)
set.seed(123)
sce <- RunParallelDivisiveICP(
  object = sce, batch.label = "Batch",
  k = 2, L = 25, C = 1, train.k.nn = 10,
  train.k.nn.prop = NULL, use.cluster.seed = FALSE,
  build.train.set = FALSE, ari.cutoff = 0.1,
  threads = 2, RNGseed = 1024
)

# Get cluster probability for all ICP runs
probs <- GetCellClusterProbability(object = sce, icp.round = 1, concatenate = TRUE)
probs[1:10, 1:5]

```

---

GetFeatureCoefficients

*Get feature coefficients*

---

**Description**

Get feature coefficients from ICP models.

**Usage**

```

GetFeatureCoefficients.SingleCellExperiment(
  object,
  icp.run = NULL,
  icp.round = NULL
)

## S4 method for signature 'SingleCellExperiment'
GetFeatureCoefficients(object, icp.run = NULL, icp.round = NULL)

```

**Arguments**

object	An object of SingleCellExperiment class with ICP cell cluster probability tables saved in <code>metadata(object)\$coranalysis\$joint.probability</code> . After running <code>RunParallelDivisiveICP</code> .
icp.run	ICP run(s) to retrieve from <code>metadata(object)\$coranalysis\$joint.probability</code> . By default NULL, i.e., all are retrieved. Specify a numeric vector to retrieve a specific set of tables.
icp.round	ICP round(s) to retrieve from <code>metadata(object)\$coranalysis\$joint.probability</code> . By default NULL, i.e., all are retrieved.

**Value**

A list of feature coefficient weights per cluster per ICP run/round.

**Examples**

```
# Import package
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Create toy SCE data
batches <- c("b1", "b2")
set.seed(239)
batch <- sample(x = batches, size = nrow(iris), replace = TRUE)
sce <- SingleCellExperiment(
  assays = list(logcounts = t(iris[, 1:4])),
  colData = DataFrame(
    "Species" = iris$Species,
    "Batch" = batch
  )
)
colnames(sce) <- paste0("samp", 1:ncol(sce))

# Prepare SCE object for analysis
sce <- PrepareData(sce)

# Multi-level integration (just for highlighting purposes; use default parameters)
set.seed(123)
sce <- RunParallelDivisiveICP(
  object = sce, batch.label = "Batch",
  k = 4, L = 25, C = 1, d = 0.5,
  train.with.bnn = FALSE,
  use.cluster.seed = FALSE,
  build.train.set = FALSE, ari.cutoff = 0.1,
  threads = 2, RNGseed = 1024
)

# GetFeatureCoefficients
gene_coefficients_icp_7_1 <- GetFeatureCoefficients(object = sce, icp.run = 7, icp.round = 1)
head(gene_coefficients_icp_7_1$icp_13)
```

HeatmapFeatures

*Heatmap visualization of the expression of features by clusters***Description**

The HeatmapFeatures function draws a heatmap of features by cluster identity.

**Usage**

```
HeatmapFeatures.SingleCellExperiment(
  object,
  clustering.label,
  features,
  use.color,
  seed.color,
  ...
)

## S4 method for signature 'SingleCellExperiment'
HeatmapFeatures(
  object,
  clustering.label,
  features,
  use.color = NULL,
  seed.color = 123,
  ...
)
```

**Arguments**

<code>object</code>	of SingleCellExperiment class
<code>clustering.label</code>	A variable name (of class character) available in the cell metadata <code>colData(object)</code> with the clustering labels (character or factor) to use.
<code>features</code>	Feature names to plot by cluster (character) matching <code>row.names(object)</code> .
<code>use.color</code>	Character specifying the colors for the clusters. By default NULL, i.e., colors are randomly chosen based on the seed given at <code>seed.color</code> . It is overwritten in case the argument <code>annotation_colors</code> is provided.
<code>seed.color</code>	Seed to randomly select colors for the clusters. By default 123. It is overwritten in case the argument <code>annotation_colors</code> is provided.
<code>...</code>	Parameters to pass to <code>pheatmap::pheatmap</code> function.

**Value**

nothing

**Examples**

```

# Import package
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Create toy SCE data
batches <- c("b1", "b2")
set.seed(239)
batch <- sample(x = batches, size = nrow(iris), replace = TRUE)
sce <- SingleCellExperiment(
  assays = list(logcounts = t(iris[, 1:4])),
  colData = DataFrame(
    "Species" = iris$Species,
    "Batch" = batch
  )
)
colnames(sce) <- paste0("samp", 1:ncol(sce))

# Plot features by clustering, i.e., grouping variable
# without scaling rows (using 'logcounts' expression):
HeatmapFeatures(
  object = sce, clustering.label = "Species",
  features = row.names(sce)[1:4]
)

# scaling rows:
HeatmapFeatures(
  object = sce, clustering.label = "Species",
  features = row.names(sce)[1:4], scale = "row"
) # scale

```

---

LogisticRegression      *Clustering projection using logistic regression from the LiblineaR R package*

---

**Description**

The function implements a script that downsamples data a dataset, trains a logistic regression classifier model and then projects its clustering onto itself using a trained L1-regularized logistic regression model.

**Usage**

```

LogisticRegression(
  training.sparse.matrix = NULL,
  training.ident = NULL,
  C = 0.3,
  reg.type = "L1",
  test.sparse.matrix = NULL,
  d = 0.3,
  batch.label = NULL,
  training_ident_subset = NULL
)

```

**Arguments**

<code>training.sparse.matrix</code>	A sparse matrix (dgCMatrix) containing training sample's feature expression data with features in rows and cells in columns. Default is NULL.
<code>training.ident</code>	A named factor containing sample's cluster labels for each cell in <code>training.sparse.matrix</code> . Default is NULL.
<code>C</code>	Cost of constraints violation in L1-regularized logistic regression (C). Default is 0.3.
<code>reg.type</code>	"L1" for LASSO and "L2" for Ridge. Default is "L1".
<code>test.sparse.matrix</code>	A sparse matrix (dgCMatrix) containing test sample's feature expression data with features in rows and cells in columns. Default is NULL.
<code>d</code>	A numeric smaller than 1 and greater than 0 that determines how many cells per cluster should be down- and oversampled ( $d$ in $N/k*d$ ), where $N$ is the total number of cells and $k$ the number of clusters. Default is 0.3.
<code>batch.label</code>	A character vector with batch labels corresponding to the cells given in <code>training.ident</code> . The character batch labels need to be named with the cells names given in <code>training.ident</code> . By default NULL, i.e., cells are sampled evenly regardless their batch.
<code>training_ident_subset</code>	A character or numeric vector with cell ids to use as train set. By default NULL. If given, the down- and oversampled parameters are ignored.

**Value**

a list containing the output of the LiblineR prediction

---

MajorityVotingFeatures

*Majority voting features by label*

---

**Description**

Get ICP feature coefficients for a label of interest by majority voting label across ICP clusters.

**Usage**

```
MajorityVotingFeatures.SingleCellExperiment(object, label)
```

```
## S4 method for signature 'SingleCellExperiment'
MajorityVotingFeatures(object, label)
```

**Arguments**

<code>object</code>	An object of <code>SingleCellExperiment</code> class with ICP cell cluster probability tables saved in <code>metadata(object)\$coranalysis\$joint.probability</code> . After running <code>RunParallelDivisiveICP</code> .
<code>label</code>	Label of interest available in <code>colData(object)</code> .

**Value**

A list of with a list of data frames with feature weights per label and a data frame with a summary by label.

**Examples**

```
## Not run:
# Import package
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Import data from Zenodo
data.url <- "https://zenodo.org/records/14845751/files/pbmc_10Xassays.rds?download=1"
sce <- readRDS(file = url(data.url))

# Multi-level integration (just for highlighting purposes; use default parameters)
set.seed(123)
sce <- RunParallelDivisiveICP(
  object = sce, batch.label = "batch",
  k = 4, L = 10, C = 1, d = 0.5,
  train.with.bnn = FALSE, use.cluster.seed = FALSE,
  build.train.set = FALSE, ari.cutoff = 0.1,
  threads = 2
)

# Get coefficients by majority voting for a given categorical variable
coeff <- MajorityVotingFeatures(object = sce, label = "cell_type")
gene_coeff$summary
order.rows <- order(coeff$feature_coeff$Monocyte$coeff_clt2,
  decreasing = TRUE
)
head(coeff$feature_coeff$Monocyte[order.rows, ], n = 10)

## End(Not run)
```

---

PCAElbowPlot

*Elbow plot of the standard deviations of the principal components*


---

**Description**

Draw an elbow plot of the standard deviations of the principal components to deduce an appropriate value for p.

**Usage**

```
PCAElbowPlot.SingleCellExperiment(object, dimred.name, return.plot)

## S4 method for signature 'SingleCellExperiment'
PCAElbowPlot(object, dimred.name = "PCA", return.plot = FALSE)
```

**Arguments**

<code>object</code>	A <code>SingleCellExperiment</code> object obtained after running <code>RunParallelDivisiveICP</code> .
<code>dimred.name</code>	Dimensional reduction name of the PCA to select from <code>reducedDimNames(object)</code> . By default "PCA".
<code>return.plot</code>	logical indicating if the <code>ggplot2</code> object should be returned. By default FALSE.

**Value**

A `ggplot2` object, if `return.plot=TRUE`.

**Examples**

```
# Import package
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Create toy SCE data
batches <- c("b1", "b2")
set.seed(239)
batch <- sample(x = batches, size = nrow(iris), replace = TRUE)
sce <- SingleCellExperiment(
  assays = list(logcounts = t(iris[, 1:4])),
  colData = DataFrame(
    "Species" = iris$Species,
    "Batch" = batch
  )
)
colnames(sce) <- paste0("samp", 1:ncol(sce))

# Prepare SCE object for analysis
sce <- PrepareData(sce)

# Multi-level integration (just for highlighting purposes; use default parameters)
set.seed(123)
sce <- RunParallelDivisiveICP(
  object = sce, batch.label = "Batch",
  k = 2, L = 25, C = 1, train.k.nn = 10,
  train.k.nn.prop = NULL, use.cluster.seed = FALSE,
  build.train.set = FALSE, ari.cutoff = 0.1,
  threads = 2, RNGseed = 1024
)

# Integrated PCA
set.seed(125) # to ensure reproducibility for the default 'irlba' method
sce <- RunPCA(object = sce, assay.name = "joint.probability", p = 10)

# Plot result
cowplot::plot_grid(
  PlotDimRed(
    object = sce, color.by = "Batch",
    legend.nrow = 1
  ),
  PlotDimRed(
    object = sce, color.by = "Species",
    legend.nrow = 1
  ),
)
```

```

    ncol = 2
  )

# Plot Elbow
PCAElbowPlot(sce)

```

---

PlotClusterTree      *Plot cluster tree*

---

### Description

Plot cluster tree by or cluster probability or categorical variable.

### Usage

```

PlotClusterTree.SingleCellExperiment(
  object,
  icp.run,
  color.by,
  use.color,
  seed.color,
  legend.title,
  return.data
)

## S4 method for signature 'SingleCellExperiment'
PlotClusterTree(
  object,
  icp.run,
  color.by = NULL,
  use.color = NULL,
  seed.color = 123,
  legend.title = color.by,
  return.data = FALSE
)

```

### Arguments

object	An object of SingleCellExperiment class.
icp.run	ICP run(s) to retrieve from metadata(object)\$coranalysis\$joint.probability. By default NULL, i.e., all are retrieved. Specify a numeric vector to retrieve a specific set of tables.
color.by	Categorical variable available in colData(object) to plot. If NULL the cluster probability is represented instead. By default NULL.
use.color	Character specifying the colors. By default NULL, i.e., colors are randomly chosen based on the seed given at seed.color.
seed.color	Seed to randomly select colors. By default 123.
legend.title	Legend title. By default the same as given at color.by. Ignored if color.by is NULL.
return.data	Return data frame used to plot. Logical. By default FALSE, i.e., only the plot is returned.

**Value**

A plot of class `ggplot` or a list with a plot of class `ggplot` and a data frame.

**Examples**

```
# Import package
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Create toy SCE data
batches <- c("b1", "b2")
set.seed(239)
batch <- sample(x = batches, size = nrow(iris), replace = TRUE)
sce <- SingleCellExperiment(
  assays = list(logcounts = t(iris[, 1:4])),
  colData = DataFrame(
    "Species" = iris$Species,
    "Batch" = batch
  )
)
colnames(sce) <- paste0("samp", 1:ncol(sce))

# Prepare SCE object for analysis
sce <- PrepareData(sce)

# Multi-level integration (just for highlighting purposes; use default parameters)
set.seed(123)
sce <- RunParallelDivisiveICP(
  object = sce, batch.label = "Batch", k = 4,
  L = 25, C = 1, d = 0.5, train.with.bnn = FALSE,
  use.cluster.seed = FALSE, build.train.set = FALSE,
  ari.cutoff = 0.1, threads = 2, RNGseed = 1024
)

# Plot probability
PlotClusterTree(object = sce, icp.run = 2)

# Plot batch label distribution
PlotClusterTree(object = sce, icp.run = 2, color.by = "Batch")

# Plot species label distribution
PlotClusterTree(object = sce, icp.run = 2, color.by = "Species")
```

---

 PlotDimRed

*Plot dimensional reduction categorical variables*


---

**Description**

Plot categorical variables in dimensional reduction.

**Usage**

```
PlotDimRed.SingleCellExperiment(
  object,
```

```

    color.by,
    dimred,
    dims,
    use.color,
    point.size,
    point.stroke,
    legend.nrow,
    seed.color,
    label,
    plot.theme,
    rasterise,
    rasterise.dpi,
    legend.justification,
    legend.size,
    legend.title
)

## S4 method for signature 'SingleCellExperiment'
PlotDimRed(
  object,
  color.by,
  dimred = tail(reducedDimNames(object), n = 1),
  dims = 1:2,
  use.color = NULL,
  point.size = 1,
  point.stroke = 1,
  legend.nrow = 2,
  seed.color = 123,
  label = FALSE,
  plot.theme = theme_classic(),
  rasterise = (ncol(object) <= 30000),
  rasterise.dpi = 300,
  legend.justification = "center",
  legend.size = 10,
  legend.title = color.by
)

```

### Arguments

<code>object</code>	An object of <code>SingleCellExperiment</code> class.
<code>color.by</code>	Categorical variable available in <code>colData(object)</code> to plot.
<code>dimred</code>	Dimensional reduction available in <code>ReducedDimNames(object)</code> to plot. By default the last dimensional reduction in the object is used.
<code>dims</code>	Dimensions from the dimensional reduction embedding to plot.
<code>use.color</code>	Character specifying the colors. By default <code>NULL</code> , i.e., colors are randomly chosen based on the seed given at <code>seed.color</code> .
<code>point.size</code>	Size of points. By default 1.
<code>point.stroke</code>	Size of stroke. By default 1.
<code>legend.nrow</code>	Display legend items by this number of rows. By default 2.
<code>seed.color</code>	Seed to randomly select colors. By default 123.

label	Logical to add or not categorical labels to the centroid categories. By default FALSE, i.e., labels are not added.
plot.theme	Plot theme available in ggplot2. By default theme_classic().
rasterise	Logical specifying if points should be rasterised or not. By default TRUE, if more than 3e4 cells, otherwise FALSE.
rasterise.dpi	In case rasterise = TRUE, DPI to use. By default 300.
legend.justification	Legend justification. By default "center".
legend.size	Legend size. By default 10
legend.title	Legend title. By default the same as given at color.by.

### Value

A plot of class ggplot.

### Examples

```
# Import package
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Create toy SCE data
batches <- c("b1", "b2")
set.seed(239)
batch <- sample(x = batches, size = nrow(iris), replace = TRUE)
sce <- SingleCellExperiment(
  assays = list(logcounts = t(iris[, 1:4])),
  colData = DataFrame(
    "Species" = iris$Species,
    "Batch" = batch
  )
)
colnames(sce) <- paste0("samp", 1:ncol(sce))

# Compute dimensional reduction
sce <- RunPCA(
  object = sce, assay.name = "logcounts", p = 4,
  pca.method = "stats"
)

# Plot batch
PlotDimRed(object = sce, color.by = "Batch", dimred = "PCA", legend.nrow = 1)

# Plot cell type annotations
PlotDimRed(
  object = sce, color.by = "Species", legend.nrow = 1,
  dimred = "PCA", label = TRUE
)
```

---

PlotExpression	<i>Plot dimensional reduction feature expression</i>
----------------	--

---

### Description

Plot feature expression in dimensional reduction.

### Usage

```
PlotExpression.SingleCellExperiment(
  object,
  color.by,
  dimred,
  scale.values,
  color.scale,
  plot.theme,
  legend.title,
  point.size,
  point.stroke
)

## S4 method for signature 'SingleCellExperiment'
PlotExpression(
  object,
  color.by,
  dimred = tail(reducedDimNames(object), n = 1),
  scale.values = FALSE,
  color.scale = "inferno",
  plot.theme = theme_classic(),
  legend.title = color.by,
  point.size = 1,
  point.stroke = 1
)
```

### Arguments

<code>object</code>	An object of <code>SingleCellExperiment</code> class.
<code>color.by</code>	Categorical variable available in <code>colData(object)</code> to plot.
<code>dimred</code>	Dimensional reduction available in <code>ReducedDimNames(object)</code> to plot. By default the last dimensional reduction in the object is used.
<code>scale.values</code>	Logical specifying if values should be scaled. By default <code>FALSE</code> , i.e., values are not scaled.
<code>color.scale</code>	Character of color scale palette to be passed to <code>ggplot2::scale_color_viridis_c</code> . By default <code>inferno</code> . Other palettes are also available such as <code>viridis</code> .
<code>plot.theme</code>	Plot theme available in <code>ggplot2</code> . By default <code>theme_classic()</code> .
<code>legend.title</code>	Legend title. By default the same as given at <code>color.by</code> .
<code>point.size</code>	Size of points. By default 1.
<code>point.stroke</code>	Size of stroke. By default 1.

**Value**

A plot of class `ggplot`.

**Examples**

```
# Import package
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Create toy SCE data
batches <- c("b1", "b2")
set.seed(239)
batch <- sample(x = batches, size = nrow(iris), replace = TRUE)
sce <- SingleCellExperiment(
  assays = list(logcounts = t(iris[, 1:4])),
  colData = DataFrame(
    "Species" = iris$Species,
    "Batch" = batch
  )
)
colnames(sce) <- paste0("samp", 1:ncol(sce))

# Compute dimensional reduction
sce <- RunPCA(
  object = sce, assay.name = "logcounts", p = 4,
  pca.method = "stats"
)

# Plot expression level of one or more features
## one
PlotExpression(object = sce, color.by = "Petal.Width")

## more than one
features <- row.names(sce)[1:4]
exp.plots <- lapply(X = features, FUN = function(x) {
  PlotExpression(object = sce, color.by = x, scale.values = TRUE)
})
cowplot::plot_grid(plotlist = exp.plots, ncol = 2, align = "vh")
```

---

 PrepareData

*Prepare SingleCellExperiment object for analysis*


---

**Description**

This function prepares the `SingleCellExperiment` object for analysis. The only required input is an object of class `SingleCellExperiment` with at least data in the `logcounts` slot.

**Usage**

```
PrepareData.SingleCellExperiment(object)

## S4 method for signature 'SingleCellExperiment'
PrepareData(object)
```

**Arguments**

`object` An object of `SingleCellExperiment` class.

**Value**

An object of `SingleCellExperiment` class.

**Examples**

```
# Import package
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Create toy SCE data
batches <- c("b1", "b2")
set.seed(239)
batch <- sample(x = batches, size = nrow(iris), replace = TRUE)
sce <- SingleCellExperiment(
  assays = list(logcounts = t(iris[, 1:4])),
  colData = DataFrame(
    "Species" = iris$Species,
    "Batch" = batch
  )
)
colnames(sce) <- paste0("samp", 1:ncol(sce))
sce <- PrepareData(sce)
```

---

RandomlyDivisiveClustering

*Split randomly every cluster into k clusters*

---

**Description**

Splits randomly every cluster given into k clusters

**Usage**

```
RandomlyDivisiveClustering(cluster, k, cluster.names = NULL)
```

**Arguments**

`cluster` A clustering result in which each cluster will be divided randomly into k clusters.

`k` The number of clusters that each cluster given in `cluster` should be randomly divided into.

`cluster.names` Names to name the cluster result given. By default `NULL`.

**Value**

A clustering result where every cluster given was split randomly into k clusters.

---

ReferenceMapping	<i>Reference mapping</i>
------------------	--------------------------

---

### Description

This function allows to project new query data sets onto a reference built with Coralysis as well as transfer cell labels from the reference to queries.

### Usage

```
ReferenceMapping.SingleCellExperiment(
  ref,
  query,
  ref.label,
  label.prune.cutoff,
  scale.query.by,
  project.umap,
  select.icp.models,
  k.nn,
  dimred.name.prefix
)

## S4 method for signature 'SingleCellExperiment,SingleCellExperiment'
ReferenceMapping(
  ref,
  query,
  ref.label,
  label.prune.cutoff = 0.5,
  scale.query.by = NULL,
  project.umap = FALSE,
  select.icp.models = metadata(ref)$coralysis$pca.params$select.icp.tables,
  k.nn = 10,
  dimred.name.prefix = ""
)
```

### Arguments

<code>ref</code>	An object of <code>SingleCellExperiment</code> class trained with Coralysis and after running <code>RunPCA(..., return.model = TRUE)</code> function.
<code>query</code>	An object of <code>SingleCellExperiment</code> class to project onto <code>ref</code> .
<code>ref.label</code>	A character cell metadata column name from the <code>ref</code> object to transfer to the queries.
<code>label.prune.cutoff</code>	A numeric cutoff value used to prune low-confidence predicted cell labels, based on the confidence probability scores stored in the <code>coral_probability</code> column of <code>colData</code> . By default is 0.5, i.e., cell labels with confidence scores less than or equal to 0.5 are considered unclassified and set to NA. The resulting pruned cell labels are stored in <code>pruned_coral_labels</code> . Set to 0 to ignore it.
<code>scale.query.by</code>	Should the query data be scaled by cell or by feature. By default is NULL, i.e., is not scaled. Scale it if reference was scaled.

<code>project.umap</code>	Project query data onto reference UMAP (logical). By default FALSE. If TRUE, the ref object needs to have a UMAP embedding obtained with <code>RunUMAP(..., return.model = TRUE)</code> function.
<code>select.icp.models</code>	Select the reference ICP models to use for query cluster probability prediction. By default <code>metadata(ref)\$coranalysis\$pca.params\$select.icp.tables</code> , i.e., the models selected to compute the reference PCA are selected. If NULL all are used. Otherwise a numeric vector should be given to select the ICP models of interest.
<code>k.nn</code>	The number of k nearest neighbors to use in the classification KNN algorithm used to transfer labels from the reference to queries (integer). By default 10.
<code>dimred.name.prefix</code>	Dimensional reduction name prefix to add to the computed PCA and UMAP. By default nothing is added, i.e., <code>dimred.name.prefix = ""</code> .

### Value

An object of `SingleCellExperiment` class.

### Examples

```
# Import package
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Create toy SCE data
batches <- c("b1", "b2")
set.seed(239)
batch <- sample(x = batches, size = nrow(iris), replace = TRUE)
sce <- SingleCellExperiment(
  assays = list(logcounts = t(iris[, 1:4])),
  colData = DataFrame(
    "Species" = iris$Species,
    "Batch" = batch
  )
)
colnames(sce) <- paste0("samp", 1:ncol(sce))

# Create reference & query SCE objects
ref <- sce[, sce$Batch == "b1"]
query <- sce[, sce$Batch == "b2"]

# 1) Train the reference
set.seed(123)
ref <- RunParallelDivisiveICP(
  object = ref, k = 2, L = 25, C = 1,
  train.k.nn = 10, train.k.nn.prop = NULL,
  use.cluster.seed = FALSE,
  build.train.set = FALSE, ari.cutoff = 0.1,
  threads = 2, RNGseed = 1024
)

# 2) Compute reference PCA & UMAP
ref <- RunPCA(ref, p = 5, return.model = TRUE, pca.method = "stats")
set.seed(123)
ref <- RunUMAP(ref, return.model = TRUE)
```

```

# Plot
PlotDimRed(object = ref, color.by = "Species", legend.nrow = 1)

# 3) Project & predict query cell labels
map <- ReferenceMapping(
  ref = ref, query = query, ref.label = "Species",
  project.umap = TRUE
)

# Confusion matrix: predictions (rows) x ground-truth (cols)
preds_x_truth <- table(map$coral_labels, map$Species)
print(preds_x_truth)

# Accuracy score
acc <- sum(diag(preds_x_truth)) / sum(preds_x_truth) * 100
print(paste0("Coralysis accuracy score: ", round(acc), "%"))

# Visualize: ground-truth, prediction, confidence scores
cowplot::plot_grid(
  PlotDimRed(
    object = map, color.by = "Species",
    legend.nrow = 1
  ),
  PlotDimRed(
    object = map, color.by = "coral_labels",
    legend.nrow = 1
  ),
  PlotExpression(
    object = map, color.by = "coral_probability",
    color.scale = "viridis"
  ),
  ncol = 2, align = "vh"
)

```

---

RunDivisiveICP

*Divisive Iterative Clustering Projection (ICP) clustering*


---

## Description

The function implements divisive Iterative Clustering Projection (ICP) clustering: a supervised learning-based clustering, which maximizes clustering similarity between the clustering and its projection by logistic regression, doing it in a divisive clustering manner.

## Usage

```

RunDivisiveICP(
  normalized.data = NULL,
  batch.label = NULL,
  k = 16,
  d = 0.3,
  r = 5,
  C = 5,
  reg.type = "L1",

```

```

max.iter = 200,
icp.batch.size = Inf,
train.with.bnn = TRUE,
train.k.nn = 10,
train.k.nn.prop = NULL,
cluster.seed = NULL,
divisive.method = "random",
allow.free.k = FALSE,
ari.cutoff = 0.5,
verbose = TRUE
)

```

## Arguments

normalized.data	A sparse matrix (dgCMatrix) containing normalized feature expression data with cells in rows and features in columns. Default is NULL.
batch.label	A character vector with batch labels corresponding to the cells given in normalized.data. The character batch labels need to be named with the cells names given in the rows of normalized.data. By default NULL, i.e., cells are sampled evenly regardless their batch.
k	A positive integer power of two, i.e., $2^{*n}$ , where $n > 0$ , specifying the number of clusters in the last Iterative Clustering Projection (ICP) round. Decreasing k leads to smaller cell populations diversity and vice versa. Default is 16, i.e., the divisive clustering 2 -> 4 -> 8 -> 16 is performed.
d	A numeric that defines how many cells per cluster should be down- and oversampled (d in ceiling( $N/k*d$ )), when stratified.downsampling=FALSE, or what fraction should be downsampled in the stratified approach ,stratified.downsampling=TRUE. Default is 0.3.
r	A positive integer that denotes the number of reiterations performed until the algorithm stops. Default is 5.
C	Cost of constraints violation (C) for L1-regulatization. Default is 0.3.
reg.type	"L1" for LASSO and "L2" for Ridge. Default is "L1".
max.iter	A positive integer that denotes the maximum number of iterations performed until the algorithm ends. Default is 200.
icp.batch.size	A positive integer that specifies how many cells to randomly select for each ICP run from the complete data set. This is a new feature intended to speed up the process with larger data sets. Default is Inf, which means using all cells.
train.with.bnn	Train data with batch nearest neighbors. Default is TRUE. Only used if batch.label is given.
train.k.nn	Train data with batch nearest neighbors using k nearest neighbors. Default is 10. Only used if train.with.bnn is TRUE.
train.k.nn.prop	A numeric (higher than 0 and lower than 1) corresponding to the fraction of cells per cluster to use as train.k.nn nearest neighbors. Default is NULL meaning that the number of train.k.nn nearest neighbors is equal to train.k.nn. If given, train.k.nn parameter is ignored and train.k.nn is calculated based on train.k.nn.prop. A vector with different proportions for the different divisive clustering rounds can be given, otherwise the same value is given for all.

<code>cluster.seed</code>	A cluster seed to start and guide the clustering to more reproducible clusterings across runs (factor). Default is NULL. Otherwise, a random clustering takes place to start divisive clustering with ICP.
<code>divisive.method</code>	Divisive method (character). One of "random" (randomly sample two clusters out of every cluster previously found), "cluster" or "cluster.batch" (sample two clusters out of every cluster previously found based on the cluster probability distribution across batches or per batch). By default "random".
<code>allow.free.k</code>	Allow free k (logical). Allow ICP algorithm to decrease the k given in case it does not find k target clusters. By default FALSE.
<code>ari.cutoff</code>	Include ICP models and probability tables with an Adjusted Rand Index higher than <code>ari.cutoff</code> (numeric). By default 0.5. A value that can range between 0 (include all) and lower than 1.
<code>verbose</code>	A logical value to print verbose during the ICP run in case. Default is TRUE. Verbose might help debugging errors by printing intermediate ICP projection results.

**Value**

A list that includes the probability matrix and the clustering similarity measures: ARI, NMI, etc.

---

RunParallelDivisiveICP

*Multi-level integration*

---

**Description**

Run divisive ICP clustering in parallel in order to perform multi-level integration.

**Usage**

```
RunParallelDivisiveICP.SingleCellExperiment(
  object,
  batch.label,
  k,
  d,
  L,
  r,
  C,
  reg.type,
  max.iter,
  threads,
  icp.batch.size,
  train.with.bnn,
  train.k.nn,
  train.k.nn.prop,
  build.train.set,
  build.train.params,
  scale.by,
  use.cluster.seed,
```

```

    divisive.method,
    allow.free.k,
    ari.cutoff,
    verbose,
    RNGseed,
    BPPARAM
)

## S4 method for signature 'SingleCellExperiment'
RunParallelDivisiveICP(
  object,
  batch.label = NULL,
  k = 16,
  d = 0.3,
  L = 50,
  r = 5,
  C = 0.3,
  reg.type = "L1",
  max.iter = 200,
  threads = 0,
  icp.batch.size = Inf,
  train.with.bnn = TRUE,
  train.k.nn = 10,
  train.k.nn.prop = 0.3,
  build.train.set = TRUE,
  build.train.params = list(),
  scale.by = NULL,
  use.cluster.seed = TRUE,
  divisive.method = "cluster.batch",
  allow.free.k = TRUE,
  ari.cutoff = 0.3,
  verbose = FALSE,
  RNGseed = 123,
  BPPARAM = NULL
)

```

### Arguments

<code>object</code>	An object of <code>SingleCellExperiment</code> class.
<code>batch.label</code>	A variable name (of class character) available in the cell metadata <code>colData(object)</code> with the batch labels (character or factor) to use. The variable provided must not contain NAs. By default NULL, i.e., cells are sampled evenly regardless their batch.
<code>k</code>	A positive integer power of two, i.e., $2^{*n}$ , where $n > 0$ , specifying the number of clusters in the last Iterative Clustering Projection (ICP) round. Decreasing $k$ leads to smaller cell populations diversity and vice versa. Default is 16, i.e., the divisive clustering 2 -> 4 -> 8 -> 16 is performed.
<code>d</code>	A numeric greater than 0 and smaller than 1 that determines how many cells $n$ are down- or oversampled from each cluster into the training data ( $n = N/k*d$ ), where $N$ is the total number of cells, $k$ is the number of clusters in ICP. Increasing above 0.3 leads gradually to smaller cell populations diversity. Default is 0.3.

L	A positive integer greater than 1 denoting the number of the ICP runs to run. Default is 50.
r	A positive integer that denotes the number of reiterations performed until the ICP algorithm stops. Increasing recommended with a significantly larger sample size (tens of thousands of cells). Default is 5.
C	A positive real number denoting the cost of constraints violation in the L1-regularized logistic regression model from the LIBLINEAR library. Decreasing leads to more stringent feature selection, i.e. less features are selected that are used to build the projection classifier. Decreasing to a very low value ( $\sim 0.01$ ) can lead to failure to identify central cell populations. Default 0.3.
reg.type	"L1" or "L2". L2-regularization was not investigated in the manuscript, but it leads to a more conventional outcome (less subpopulations). Default is "L1".
max.iter	A positive integer that denotes the maximum number of iterations performed until ICP stops. This parameter is only useful in situations where ICP converges extremely slowly, preventing the algorithm to run too long. In most cases, reaching the number of reiterations ( $r=5$ ) terminates the algorithm. Default is 200.
threads	A positive integer that specifies how many logical processors (threads) to use in parallel computation. Set 1 to disable parallelism altogether or 0 to use all available threads except one. Default is 0. This argument is ignored if BPPARAM is provided as threads should be given directly to the BiocParallelParam object.
icp.batch.size	A positive integer that specifies how many cells to randomly select. It behaves differently depending on build.train.set. If build.train.set=FALSE, it randomly samples cells for each ICP run from the complete dataset. If build.train.set=TRUE, it randomly samples cells once, before building the training set with the sampled cells (per batch if batch.label different than NULL). Default is Inf, which means using all cells.
train.with.bnn	Train data with batch nearest neighbors. Default is TRUE. Only used if batch.label is given.
train.k.nn	Train data with batch nearest neighbors using k nearest neighbors. Default is 10. Only used if train.with.bnn is TRUE and train.k.nn.prop is NULL.
train.k.nn.prop	A numeric (higher than 0 and lower than 1) corresponding to the fraction of cells per cluster to use as train.k.nn nearest neighbors. If NULL the number of train.k.nn nearest neighbors is equal to train.k.nn. If given, train.k.nn parameter is ignored and train.k.nn is calculated based on train.k.nn.prop. By default 0.3 meaning that 30 proportions for the different divisive clustering rounds can be given, otherwise the same value is given for all.
build.train.set	Logical specifying if a training set should be built from the data or the whole data should be used for training. By default TRUE.
build.train.params	A list of parameters to be passed to the function AggregateDataByBatch(). Only provided if build.train.set is TRUE.
scale.by	A character specifying if the data should be scaled by cell or by feature before training. Default is NULL, i.e., the data is not scaled before training.
use.cluster.seed	Should the same starting clustering result be provided to ensure more reproducible results (logical). If FALSE, each ICP run starts with a total random clustering and, thus, independent clustering. By default TRUE, i.e., the same clustering result is provided based on PCA density sampling. If batch.label different than NULL, the PCA density sampling is performed in a batch wise manner.

<code>divisive.method</code>	Divisive method (character). One of "random" (randomly sample two clusters out of every cluster previously found), "cluster" or "cluster.batch" (sample two clusters out of every cluster previously found based on the cluster probability distribution across batches or per batch). By default "cluster.batch". If <code>batch.label</code> is NULL, it is automatically set to <code>cluster</code> . It can be set to <code>random</code> if explicitly provided.
<code>allow.free.k</code>	Allow free k (logical). Allow ICP algorithm to decrease the k given in case it does not find k target clusters. By default TRUE.
<code>ari.cutoff</code>	Include ICP models and probability tables with an Adjusted Rand Index higher than <code>ari.cutoff</code> (numeric). By default 0.3. A value that can range between 0 (include all) and lower than 1.
<code>verbose</code>	A logical value to print verbose during the ICP run in case. Default is FALSE. Verbose might help debugging errors by printing intermediate ICP projection results.
<code>RNGseed</code>	Seed number passed to the parallel backend via <code>BiocParallel</code> to ensure reproducibility. Defaults to 123. If the <code>BPPARAM</code> parameter is provided, <code>RNGseed</code> is ignored and should be set within <code>BPPARAM</code> .
<code>BPPARAM</code>	A <code>BiocParallelParam</code> object specifying the parallel backend to use. This controls how tasks are distributed across workers. Use <code>MulticoreParam</code> (for Unix-like systems) and <code>SnowParam</code> (for Windows or cross-platform). If not specified, i.e., NULL, the default backend uses <code>MulticoreParam</code> for Unix-like systems and <code>SnowParam</code> for Windows.

### Value

A `SingleCellExperiment` object.

### Examples

```
# Import package
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Create toy SCE data
batches <- c("b1", "b2")
set.seed(239)
batch <- sample(x = batches, size = nrow(iris), replace = TRUE)
sce <- SingleCellExperiment(
  assays = list(logcounts = t(iris[, 1:4])),
  colData = DataFrame(
    "Species" = iris$Species,
    "Batch" = batch
  )
)
colnames(sce) <- paste0("samp", 1:ncol(sce))

# Prepare SCE object for analysis
sce <- PrepareData(sce)

# Multi-level integration (just for highlighting purposes; use default parameters)
set.seed(123)
sce <- RunParallelDivisiveICP(
  object = sce, batch.label = "Batch",
  k = 2, L = 25, C = 1, train.k.nn = 10,
```

```

    train.k.nn.prop = NULL, use.cluster.seed = FALSE,
    build.train.set = FALSE, ari.cutoff = 0.1,
    threads = 2, RNGseed = 1024
  )

  # Integrated PCA
  set.seed(125) # to ensure reproducibility for the default 'irlba' method
  sce <- RunPCA(object = sce, assay.name = "joint.probability", p = 10)

  # Plot result
  cowplot::plot_grid(
    PlotDimRed(
      object = sce, color.by = "Batch",
      legend.nrow = 1
    ),
    PlotDimRed(
      object = sce, color.by = "Species",
      legend.nrow = 1
    ),
    ncol = 2
  )

```

---

RunPCA

*Principal Component Analysis*


---

## Description

Perform principal component analysis using assays or the joint probability matrix as input.

## Usage

```

RunPCA.SingleCellExperiment(
  object,
  assay.name,
  p,
  scale,
  center,
  threshold,
  pca.method,
  return.model,
  select.icp.tables,
  features,
  dimred.name
)

## S4 method for signature 'SingleCellExperiment'
RunPCA(
  object,
  assay.name = "joint.probability",
  p = 50,
  scale = TRUE,
  center = TRUE,

```

```

threshold = 0,
pca.method = "irlba",
return.model = FALSE,
select.icp.tables = NULL,
features = NULL,
dimred.name = "PCA"
)

```

### Arguments

<code>object</code>	A <code>SingleCellExperiment</code> object.
<code>assay.name</code>	Name of the assay to compute PCA. One of <code>assayNames(object)</code> or <code>joint.probability</code> . By default <code>joint.probability</code> is used. Use <code>joint.probability</code> to obtain an integrated embedding after running <code>RunParallelDivisiveICP</code> . One of the assays in <code>assayNames(object)</code> can be provided before performing integration to assess if data requires integration.
<code>p</code>	A positive integer denoting the number of principal components to calculate and select. Default is 50.
<code>scale</code>	A logical specifying whether the probabilities should be standardized to unit-variance before running PCA. Default is <code>TRUE</code> .
<code>center</code>	A logical specifying whether the probabilities should be centered before running PCA. Default is <code>TRUE</code> .
<code>threshold</code>	A threshold for filtering out ICP runs before PCA with the lower terminal projection accuracy below the threshold. Default is 0.
<code>pca.method</code>	A character specifying the PCA method. One of <code>"irlba"</code> (default), <code>"RSpectra"</code> or <code>"stats"</code> . Set seed before, if the method is <code>"irlba"</code> to ensure reproducibility.
<code>return.model</code>	A logical specifying if the PCA model should or not be retrieved. By default <code>FALSE</code> . Only implemented for <code>pca.method = "stats"</code> . If <code>TRUE</code> , the <code>pca.method</code> is coerced to <code>"stats"</code> .
<code>select.icp.tables</code>	Select the ICP cluster probability tables to perform PCA. By default <code>NULL</code> , i.e., all are used, except if the ICP tables were obtained with the function <code>RunParallelDivisiveICP</code> , in which the ICP tables correspond to the last round of divisive clustering for every epoch. A vector of integers should be given otherwise.
<code>features</code>	A character of feature names matching <code>row.names(object)</code> to select from before computing PCA. Only used if <code>assay.name</code> is one of the assays in <code>assayNames(object)</code> , otherwise it is ignored.
<code>dimred.name</code>	Dimensional reduction name given to the returned PCA. By default <code>"PCA"</code> .

### Value

object of `SingleCellExperiment` class

### Examples

```

# Import package
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Create toy SCE data
batches <- c("b1", "b2")
set.seed(239)

```

```

batch <- sample(x = batches, size = nrow(iris), replace = TRUE)
sce <- SingleCellExperiment(
  assays = list(logcounts = t(iris[, 1:4])),
  colData = DataFrame(
    "Species" = iris$Species,
    "Batch" = batch
  )
)
colnames(sce) <- paste0("samp", 1:ncol(sce))

# Prepare SCE object for analysis
sce <- PrepareData(sce)

# Multi-level integration (just for highlighting purposes; use default parameters)
set.seed(123)
sce <- RunParallelDivisiveICP(
  object = sce, batch.label = "Batch",
  k = 2, L = 25, C = 1, train.k.nn = 10,
  train.k.nn.prop = NULL, use.cluster.seed = FALSE,
  build.train.set = FALSE, ari.cutoff = 0.1,
  threads = 2, RNGseed = 1024
)

# Integrated PCA
set.seed(125) # to ensure reproducibility for the default 'irlba' method
sce <- RunPCA(object = sce, assay.name = "joint.probability", p = 10)

# Plot result
cowplot::plot_grid(
  PlotDimRed(
    object = sce, color.by = "Batch",
    legend.nrow = 1
  ),
  PlotDimRed(
    object = sce, color.by = "Species",
    legend.nrow = 1
  ),
  ncol = 2
)

```

RunTSNE

*Barnes-Hut implementation of t-Distributed Stochastic Neighbor Embedding (t-SNE)*

## Description

Run nonlinear dimensionality reduction using t-SNE with the PCA-transformed consensus matrix as input.

## Usage

```
RunTSNE.SingleCellExperiment(
  object,
```

```

    dims,
    dimred.type,
    perplexity,
    dimred.name,
    ...
)

## S4 method for signature 'SingleCellExperiment'
RunTSNE(
  object,
  dims = NULL,
  dimred.type = "PCA",
  perplexity = 30,
  dimred.name = "TSNE",
  ...
)

```

### Arguments

<code>object</code>	Object of <code>SingleCellExperiment</code> class.
<code>dims</code>	Dimensions to select from <code>dimred.type</code> . By default <code>NULL</code> , i.e., all the dimensions are selected. Provide a numeric vector to select a specific range, e.g., <code>dims = 1:10</code> to select the first 10 dimensions.
<code>dimred.type</code>	Dimensional reduction type to use. By default <code>"PCA"</code> .
<code>perplexity</code>	Perplexity of t-SNE.
<code>dimred.name</code>	Dimensional reduction name given to the returned t-SNE. By default <code>"TSNE"</code> .
<code>...</code>	Parameters to be passed to the <code>Rtsne</code> function. The parameters given should match the parameters accepted by the <code>Rtsne</code> function. Check possible parameters with <code>?Rtsne::Rtsne</code> .

### Value

A `SingleCellExperiment` object.

### Examples

```

# Import package
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Create toy SCE data
batches <- c("b1", "b2")
set.seed(239)
batch <- sample(x = batches, size = nrow(iris), replace = TRUE)
sce <- SingleCellExperiment(
  assays = list(logcounts = t(iris[, 1:4])),
  colData = DataFrame(
    "Species" = iris$Species,
    "Batch" = batch
  )
)
colnames(sce) <- paste0("samp", 1:ncol(sce))

# Run PCA

```

```

set.seed(125) # to ensure reproducibility for the default 'irlba' method
sce <- RunPCA(
  object = sce, assay.name = "logcounts",
  pca.method = "stats", p = nrow(sce)
)

# Run t-SNE
set.seed(125) # to ensure reproducibility for the default 'irlba' method
sce <- RunTSNE(object = sce, dimred.type = "PCA", check_duplicates = FALSE)

# Plot result
cowplot::plot_grid(
  PlotDimRed(
    object = sce, color.by = "Batch",
    legend.nrow = 1
  ),
  PlotDimRed(
    object = sce, color.by = "Species",
    legend.nrow = 1
  ),
  ncol = 2
)

```

---

RunUMAP

*Uniform Manifold Approximation and Projection (UMAP)*


---

## Description

Run nonlinear dimensionality reduction using UMAP with a dimensional reduction as input.

## Usage

```

RunUMAP.SingleCellExperiment(
  object,
  dims,
  dimred.type,
  return.model,
  umap.method,
  dimred.name,
  ...
)

## S4 method for signature 'SingleCellExperiment'
RunUMAP(
  object,
  dims = NULL,
  dimred.type = "PCA",
  return.model = FALSE,
  umap.method = "umap",
  dimred.name = "UMAP",
  ...
)

```

**Arguments**

<code>object</code>	An object of <code>SingleCellExperiment</code> class.
<code>dims</code>	Dimensions to select from <code>dimred.type</code> . By default <code>NULL</code> , i.e., all the dimensions are selected. Provide a numeric vector to select a specific range, e.g., <code>dims = 1:10</code> to select the first 10 dimensions.
<code>dimred.type</code>	Dimensional reduction type to use. By default <code>"PCA"</code> .
<code>return.model</code>	Return UMAP model. By default <code>FALSE</code> .
<code>umap.method</code>	UMAP method to use: <code>"umap"</code> or <code>"uwot"</code> . By default <code>"umap"</code> .
<code>dimred.name</code>	Dimensional reduction name given to the returned UMAP. By default <code>"UMAP"</code> .
<code>...</code>	Parameters to be passed to the <code>umap</code> function. The parameters given should match the parameters accepted by the <code>umap</code> function depending on the <code>umap.method</code> given. Check possible parameters with <code>?umap::umap</code> or <code>?uwot::umap</code> depending if <code>umap.method</code> is <code>"umap"</code> or <code>"uwot"</code> .

**Value**

A `SingleCellExperiment` object.

**Examples**

```
# Import package
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Create toy SCE data
batches <- c("b1", "b2")
set.seed(239)
batch <- sample(x = batches, size = nrow(iris), replace = TRUE)
sce <- SingleCellExperiment(
  assays = list(logcounts = t(iris[, 1:4])),
  colData = DataFrame(
    "Species" = iris$Species,
    "Batch" = batch
  )
)
colnames(sce) <- paste0("samp", 1:ncol(sce))

# Prepare SCE object for analysis
sce <- PrepareData(sce)

# Multi-level integration (just for highlighting purposes; use default parameters)
set.seed(123)
sce <- RunParallelDivisiveICP(
  object = sce, batch.label = "Batch",
  k = 2, L = 25, C = 1, train.k.nn = 10,
  train.k.nn.prop = NULL, use.cluster.seed = FALSE,
  build.train.set = FALSE, ari.cutoff = 0.1,
  threads = 2, RNGseed = 1024
)

# Integrated PCA
set.seed(125) # to ensure reproducibility for the default 'irlba' method
sce <- RunPCA(object = sce, assay.name = "joint.probability", p = 10)
```

```

# Plot result
cowplot::plot_grid(
  PlotDimRed(
    object = sce, color.by = "Batch",
    legend.nrow = 1
  ),
  PlotDimRed(
    object = sce, color.by = "Species",
    legend.nrow = 1
  ),
  ncol = 2
)

# Run UMAP
set.seed(123)
sce <- RunUMAP(sce, dimred.type = "PCA")

# Plot results
# Plot result
cowplot::plot_grid(
  PlotDimRed(
    object = sce, color.by = "Batch",
    legend.nrow = 1
  ),
  PlotDimRed(
    object = sce, color.by = "Species",
    legend.nrow = 1
  ),
  ncol = 2
)

```

---

SampleClusterBatchProbs

*Sample cells based on cluster probabilities distribution batch wise*

---

## Description

Samples cells based on cluster probabilities distribution batch wise

## Usage

```
SampleClusterBatchProbs(cluster, probs, batch, q.split = 0.5)
```

## Arguments

cluster	Clustering cell labels predicted by ICP (factor).
probs	Clustering probabilities predicted by ICP (matrix).
batch	Batch labels for the corresponding clusters (character or factor).
q.split	Split (cell) batch principal component distribution by this quantile (numeric). By default 0.5, i.e., median.

**Value**

A factor with cell cluster identities.

---

SampleClusterProbs	<i>Sample cells based on cluster probabilities distribution</i>
--------------------	---

---

**Description**

Samples cells based on cluster probabilities distribution

**Usage**

```
SampleClusterProbs(cluster, probs, q.split = 0.5)
```

**Arguments**

cluster	Clustering cell labels predicted by ICP (factor).
probs	Clustering probabilities predicted by ICP (matrix).
q.split	Split (cell) batch principal component distribution by this quantile (numeric). By default 0.5, i.e., median.

**Value**

A factor with cell cluster identities.

---

SamplePCACells	<i>Sample cells based on principal components distribution</i>
----------------	--

---

**Description**

Samples cells based on their distributions along one principal component

**Usage**

```
SamplePCACells(
  data,
  batch = NULL,
  q.split = 0.5,
  p = 30,
  use.pc = "PC1",
  center = TRUE,
  scale. = TRUE
)
```

**Arguments**

<code>data</code>	Data to compute PCA and sample cells from. Rows and columns should represent cells and features, respectively.
<code>batch</code>	Batch cell label identity (character) matching cells giving in data. Use NULL in the absence of batches. If the batch is given the cells are sampled in a batch wise manner, otherwise the cells are sampled without any grouping factor. By default is NULL.
<code>q.split</code>	Split (cell) batch principal component distribution by this quantile (numeric). By default 0.5, i.e., median.
<code>p</code>	Number of principal components to compute (integer). By default 30.
<code>use.pc</code>	Which principal component should be used for sampling cells per batch. By default "PC1", i.e., first principal component is used.
<code>center</code>	Should the features given in data centered before performing the PCA (logical). By default TRUE.
<code>scale.</code>	Should the features given in data scaled before performing the PCA (logical). By default TRUE.

**Value**

A factor with cell cluster identities (two clusters).

---

Scale	<i>Scale a sparse matrix by row or column</i>
-------	---

---

**Description**

Faster implementation of scale function. It diverges from the scale function by not performing the root-mean-square when `scale=TRUE` and `center=FALSE`. In this case it divides the values by the standard deviation of the column or row used (depending on `scale.by`).

**Usage**

```
Scale(x, center = TRUE, scale = TRUE, scale.by = "col")
```

**Arguments**

<code>x</code>	A matrix of class 'dgCMatrix'.
<code>center</code>	A logical. By default TRUE. Subtract the values by the row or column mean depending on the 'scale.by' parameter.
<code>scale</code>	A logical. By default TRUE. Divide the values by the row or column standard deviation depending on the 'scale.by' parameter
<code>scale.by</code>	Scale by 'row' or 'col' (=column), i.e., use the row or column mean and/or standard deviations to center and /or scale the data. Default is col.

**Value**

A matrix of class 'dgCMatrix'.

---

ScaleByBatch                      *Scale sparse matrix by features (column) by batch*

---

**Description**

Scales features by batch

**Usage**

```
ScaleByBatch(x, batch)
```

**Arguments**

x	A matrix of class 'dgCMatrix'. Cells by features (rows x columns).
batch	A character vector with batch labels corresponding to the cells given in x. The character batch labels need to be named with the cells names given in the rows of x.

**Value**

A scaled matrix of class 'dgCMatrix'.

---

SummariseCellClusterProbability  
*Summarise ICP cell cluster probability*

---

**Description**

Summarise ICP cell cluster probability table(s)

**Usage**

```
SummariseCellClusterProbability.SingleCellExperiment(
  object,
  icp.run,
  icp.round,
  funs,
  scale.funs,
  save.in.sce
)

## S4 method for signature 'SingleCellExperiment'
SummariseCellClusterProbability(
  object,
  icp.run = NULL,
  icp.round = NULL,
  funs = c("mean", "median"),
  scale.funs = TRUE,
  save.in.sce = TRUE
)
```

**Arguments**

object	An object of SingleCellExperiment class with ICP cell cluster probability tables saved in <code>metadata(object)\$coranalysis\$joint.probability</code> . After running <code>RunParallelDivisiveICP</code> .
icp.run	ICP run(s) to retrieve from <code>metadata(object)\$coranalysis\$joint.probability</code> . By default NULL, i.e., all are retrieved. Specify a numeric vector to retrieve a specific set of tables.
icp.round	ICP round(s) to retrieve from <code>metadata(object)\$coranalysis\$joint.probability</code> . By default NULL, i.e., all are retrieved.
funs	Functions to summarise ICP cell cluster probability: "mean" and/or "median". By default <code>c("mean", "median")</code> , i.e, both mean and median are calculated. Set to NULL to not estimate any.
scale.funs	Scale in the range 0-1 the summarised probability obtained with funs. By default TRUE, i.e., summarised probability will be scaled in the 0-1 range.
save.in.sce	Save the data frame into the cell metadata from the SingleCellExperiment object or return the data frame. By default TRUE, i.e., the summary of probabilities retrieved is save in the SCE object in <code>colData(object)</code> .

**Value**

A data frame or a SingleCellExperiment object with ICP cell cluster probability summarised.

**Examples**

```
# Import package
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Create toy SCE data
batches <- c("b1", "b2")
set.seed(239)
batch <- sample(x = batches, size = nrow(iris), replace = TRUE)
sce <- SingleCellExperiment(
  assays = list(logcounts = t(iris[, 1:4])),
  colData = DataFrame(
    "Species" = iris$Species,
    "Batch" = batch
  )
)
colnames(sce) <- paste0("samp", 1:ncol(sce))

# Prepare SCE object for analysis
sce <- PrepareData(sce)

# Multi-level integration (just for highlighting purposes; use default parameters)
set.seed(123)
sce <- RunParallelDivisiveICP(
  object = sce, batch.label = "Batch",
  k = 2, L = 25, C = 1, train.k.nn = 10,
  train.k.nn.prop = NULL, use.cluster.seed = FALSE,
  build.train.set = FALSE, ari.cutoff = 0.1,
  threads = 2, RNGseed = 1024
)
```

```

# Integrated PCA
set.seed(125) # to ensure reproducibility for the default 'irlba' method
sce <- RunPCA(object = sce, assay.name = "joint.probability", p = 10)

# Summarise cluster probability
sce <- SummariseCellClusterProbability(
  object = sce, icp.round = 1,
  save.in.sce = TRUE
) # saved in 'colData'

# Plot the clustering result for ICP run no. 3
PlotDimRed(object = sce, color.by = "icp_run_round_3_1_clusters")

# Plot Coralysis mean cell cluster probabilities
PlotExpression(
  object = sce, color.by = "mean_probs",
  color.scale = "viridis"
)

```

---

TabulateCellBinsByGroup

*Tabulate cell bins by group*

---

## Description

Frequency of cells per cell cluster probability bin by group for each label. The label has to be specified beforehand to the function `BinCellClusterProbability()`.

## Usage

```
TabulateCellBinsByGroup.SingleCellExperiment(object, group, relative, margin)
```

```
## S4 method for signature 'SingleCellExperiment'
```

```
TabulateCellBinsByGroup(object, group, relative = FALSE, margin = 1)
```

## Arguments

<code>object</code>	An object of <code>SingleCellExperiment</code> class obtained with the function <code>BinCellClusterProbability()</code> .
<code>group</code>	Character specifying the <code>colData</code> variable from the <code>SingleCellExperiment</code> object provided to the function <code>BinCellClusterProbability()</code> to use as categorical group variable.
<code>relative</code>	Logical specifying if relative proportions of cell bins per group should be returned. By default <code>FALSE</code> , i.e., absolute values are returned.
<code>margin</code>	If <code>relative</code> is <code>TRUE</code> , proportions should be calculated by: rows (1, the default); columns (2); or overall ( <code>NULL</code> ).

## Value

A list of tables with the frequency of cells per bin of cell cluster probability by group for each label.

**Examples**

```

# Packages
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Import data from Zenodo
data.url <- "https://zenodo.org/records/14845751/files/pbmc_10Xassays.rds?download=1"
sce <- readRDS(file = url(data.url))

# Prepare data
sce <- PrepareData(object = sce)

# Multi-level integration - 'L = 4' just for highlighting purposes
set.seed(123)
sce <- RunParallelDivisiveICP(
  object = sce, batch.label = "batch", L = 4,
  threads = 2
)

# Cell states SCE object for a given cell type annotation or clustering
cellstate.sce <- BinCellClusterProbability(
  object = sce, label = "cell_type",
  icp.round = 4, bins = 20
)
cellstate.sce

# Tabulate cell bins by group
# give an interesting variable to the "group" parameter
cellbins.tables <- TabulateCellBinsByGroup(
  object = cellstate.sce,
  group = "batch",
  relative = TRUE,
  margin = 1
)

```

---

VlnPlot

*Visualization of feature expression using violin plots*


---

**Description**

The VlnPlot function enables visualizing expression levels of feature(s), across clusters using violin plots.

**Usage**

```

VlnPlot.SingleCellExperiment(
  object,
  clustering.label,
  features,
  return.plot,
  rotate.x.axis.labels
)

```

```
## S4 method for signature 'SingleCellExperiment'
VlnPlot(
  object,
  clustering.label,
  features,
  return.plot = FALSE,
  rotate.x.axis.labels = FALSE
)
```

### Arguments

`object` of `SingleCellExperiment` class

`clustering.label` A variable name (of class character) available in the cell metadata `colData(object)` with the clustering labels (character or factor) to use.

`features` Feature names to plot by cluster (character) matching `row.names(object)`.

`return.plot` `return.plot` whether to return the `ggplot2` object. Default is `FALSE`.

`rotate.x.axis.labels` a logical denoting whether the x-axis labels should be rotated 90 degrees or just draw it. Default is `FALSE`.

### Value

A `ggplot2` object if `return.plot=TRUE`.

### Examples

```
# Import package
suppressPackageStartupMessages(library("SingleCellExperiment"))

# Create toy SCE data
batches <- c("b1", "b2")
set.seed(239)
batch <- sample(x = batches, size = nrow(iris), replace = TRUE)
sce <- SingleCellExperiment(
  assays = list(logcounts = t(iris[, 1:4])),
  colData = DataFrame(
    "Species" = iris$Species,
    "Batch" = batch
  )
)
colnames(sce) <- paste0("samp", 1:ncol(sce))

# Plot features by clustering/grouping variable
VlnPlot(sce,
  clustering.label = "Species",
  features = row.names(sce)[1:4],
  rotate.x.axis.labels = TRUE
)
```

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