

# Package ‘scDD’

December 6, 2024

**Version** 1.30.0

**Title** Mixture modeling of single-cell RNA-seq data to identify genes with differential distributions

**Description** This package implements a method to analyze single-cell RNA-seq Data utilizing flexible Dirichlet Process mixture models. Genes with differential distributions of expression are classified into several interesting patterns of differences between two conditions. The package also includes functions for simulating data with these patterns from negative binomial distributions.

**Depends** R (>= 3.4)

**NeedsCompilation** yes

**Imports** fields, mclust, BiocParallel, outliers, ggplot2, EBSeq, arm, SingleCellExperiment, SummarizedExperiment, grDevices, graphics, stats, S4Vectors, scran

**Suggests** BiocStyle, knitr, gridExtra

**License** GPL-2

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**VignetteBuilder** knitr

**biocViews** ImmunoOncology, Bayesian, Clustering, RNASeq, SingleCell, MultipleComparison, Visualization, DifferentialExpression

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**Author** Keegan Korthauer [cre, aut] (<<https://orcid.org/0000-0002-4565-1654>>)

**Maintainer** Keegan Korthauer <keegan@stat.ubc.ca>

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calcMV

*calcMV*

---

### Description

Calculate empirical means and variances of selected genes in a given dataset.

### Usage

```
calcMV(a, FC = 1, FC.thresh = NA, threshold = Inf,
       include.zeroes = FALSE)
```

**Arguments**

a	Numeric vector of values to calculate empirical mean and variance.
FC	Fold change for the mean and standard deviation. Default value is 1.
FC.thresh	Alternate fold change for the mean and standard deviation when the (log nonzero) mean is above the value of threshold. Default value is FC.
threshold	Mean threshold value which dictates which fold change value to use for multiplying mean and standard deviation. Default value is Inf (so FC is always used).
include zeroes	Logical value indicating whether the zero values should be included in the calculations of the empirical means and variances.

**Details**

Calculate empirical means and variances of selected genes in a given dataset. Optionally, multiply the means and standard deviations by a fold change value, which can also vary by mean value. If the mean is below some specified threshold `threshold`, use one fold change value `FC`. If above the threshold, use the alternate fold change value `FC.thresh`. Estimates of mean and variance are robust to outliers.

**Value**

MV Vector of two elements, first contains the empirical mean estimate, second contains the empirical variance estimate (optionally multiplied by a fold change).

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 calcRP

*calcRP*


---

**Description**

Calculate parameter R and P in NB distribution

**Usage**

```
calcRP(Emean, Evar)
```

**Arguments**

Emean	Empirical mean
Evar	Empirical variance

**Value**

RP Vector of two elements, first contains method of moments estimator for r and second contains method of moments estimator for p (parameters of NB distribution)

**References**

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendziorski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

---

 classifyDD

*classifyDD*


---

### Description

Classify significantly DD genes into the four categories (DE, DP, DM or DB) based on posterior distributions of cluster mean parameters

### Usage

```
classifyDD(pe_mat, condition, sig_genes, oa, c1, c2, alpha, m0, s0, a0, b0,
  log.nonzero = TRUE, adjust.perms = FALSE, ref, min.size = 3)
```

### Arguments

pe_mat	Matrix with genes in rows and samples in columns. Column names indicate condition.
condition	Vector of condition indicators (with two possible values).
sig_genes	Vector of the indices of significantly DD genes (indicating the row number of pe_mat)
oa	List item with one item for each gene where the first element contains the cluster membership for each nonzero sample in the overall (pooled) fit.
c1	List item with one item for each gene where the first element contains the cluster membership for each nonzero sample in condition 1 only fit
c2	List item with one item for each gene where the first element contains the cluster membership for each nonzero sample in condition 2 only fit
alpha	Value for the Dirichlet concentration parameter
m0	Prior mean value for generating distribution of cluster means
s0	Prior precision value for generating distribution of cluster means
a0	Prior shape parameter value for the generating distribution of cluster precision
b0	Prior scale parameter value for the generating distribution of cluster precision
log.nonzero	Logical indicating whether to perform log transformation of nonzero values.
adjust.perms	Logical indicating whether or not to adjust the permutation tests for the sample detection rate (proportion of nonzero values). If true, the residuals of a linear model adjusted for detection rate are permuted, and new fitted values are obtained using these residuals.
ref	one of two possible values in condition; represents the referent category.
min.size	a positive integer that specifies the minimum size of a cluster (number of cells) for it to be used during the classification step. Any clusters containing fewer than min.size cells will be considered an outlier cluster and ignored in the classification algorithm. The default value is three.

### Value

cat Character vector of the same length as sig\_genes that indicates which category of DD each significant gene belongs to (DE, DP, DM, DB, or NC (no call))

## References

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendziorski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

feDP

*feDP*

## Description

Function to identify additional DP genes, since clustering process can be consistent within each condition and still have differential proportion within each mode. The Bayes factor score also tends to be small when the correct number of clusters is not correctly detected; in that case differential proportion will manifest as a mean shift.

## Usage

```
feDP(pe_mat, condition, sig_genes, oa, c1, c2, log.nonzero = TRUE,
     testZeroes = FALSE, adjust.perms = FALSE, min.size = 3)
```

## Arguments

pe_mat	Matrix with genes in rows and samples in columns. Column names indicate condition.
condition	Vector of condition indicators (with two possible values).
sig_genes	Vector of the indices of significantly DD genes (indicating the row number of pe_mat)
oa	List item with one item for each gene where the first element contains the cluster membership for each nonzero sample in the overall (pooled) fit.
c1	List item with one item for each gene where the first element contains the cluster membership for each nonzero sample in condition 1 only fit
c2	List item with one item for each gene where the first element contains the cluster membership for each nonzero sample in condition 2 only fit
log.nonzero	Logical indicating whether to perform log transformation of nonzero values.
testZeroes	Logical indicating whether or not to test for a difference in the proportion of zeroes. This will only be done for genes that have at least one zero value (genes where all cells have a nonzero value will have a 'zero.pvalue' of NA).
adjust.perms	Logical indicating whether or not to adjust the permutation tests for the sample detection rate (proportion of nonzero values). If true, the residuals of a linear model adjusted for detection rate are permuted, and new fitted values are obtained using these residuals.
min.size	a positive integer that specifies the minimum size of a cluster (number of cells) for it to be used during the classification step. Any clusters containing fewer than min.size cells will be considered an outlier cluster and ignored in the classification algorithm. The default value is three.

**Details**

The Fisher's Exact test is used to test for independence of condition membership and clustering when the clustering is the same across conditions as it is overall (and is multimodal). When clustering within condition is not multimodal or is different across conditions (most often the case), an FDR-adjusted t-test is performed to detect overall mean shifts.

**Value**

cat Character vector of the same length as sig\_genes that indicates which nonsignificant genes by the permutation test belong to the DP category

---

findFC	<i>findFC</i>
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**Description**

Find the appropriate Fold Change vectors for simulation that will be use in classic differential expression case.

**Usage**

```
findFC(SCdat, index, sd.range = c(1, 3), N = 4, overExpressionProb = 0.5,
       plot.FC = FALSE, condition = "condition")
```

**Arguments**

SCdat	An object of class SingleCellExperiment that contains normalized single-cell expression and metadata. The assays slot contains a named list of matrices, where the normalized counts are housed in the one named normcounts. This matrix should have one row for each gene and one sample for each column. The colData slot should contain a data.frame with one row per sample and columns that contain metadata for each sample. This data.frame should contain a variable that represents biological condition, which is in the form of numeric values (either 1 or 2) that indicates which condition each sample belongs to (in the same order as the columns of normcounts). Optional additional metadata about each cell can also be contained in this data.frame, and additional information about the experiment can be contained in the metadata slot as a list.
index	Reasonable set of genes for simulation
sd.range	Numeric vector of length two which describes the interval (lower, upper) of standard deviations of fold changes to randomly select.
N	Integer value for the number of bins to divide range of fold changes for calculating standard deviations
overExpressionProb	Numeric value between 0 and 1 which describes the ratio of over to under expression values to sample.
plot.FC	Logical indicating whether or not to plot the observed and simulated log2 fold changes.

**condition** A character object that contains the name of the column in `colData` that represents the biological group or condition of interest (e.g. treatment versus control). Note that this variable should only contain two possible values since `scDD` can currently only handle two-group comparisons. The default option assumes that there is a column named "condition" that contains this variable.

### Details

This code is a modified version of Sam Younkin's `simulate FC` function. Major things that were changed are (1) standard deviations are calculated only on the nonzeros, (2) the sampling of FCs is uniform on the log scale instead of the raw scale, and (3) the binning is done by quantiles instead of evenly spaced along the average expression values.

### Value

FC.vec Return Fold Change Vectors

### References

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendziorski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

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findIndex

*findIndex*

---

### Description

Find a reasonable set of genes (one mode and at least 25 to use for simulation).

### Usage

```
findIndex(SCdat, condition = "condition")
```

### Arguments

**SCdat** An object of class `SingleCellExperiment` that contains normalized single-cell expression and metadata. The `assays` slot contains a named list of matrices, where the normalized counts are housed in the one named `normcounts`. This matrix should have one row for each gene and one sample for each column. The `colData` slot should contain a `data.frame` with one row per sample and columns that contain metadata for each sample. This `data.frame` should contain a variable that represents biological condition, which is in the form of numeric values (either 1 or 2) that indicates which condition each sample belongs to (in the same order as the columns of `normcounts`). Optional additional metadata about each cell can also be contained in this `data.frame`, and additional information about the experiment can be contained in the `metadata` slot as a list.

**condition** A character object that contains the name of the column in `colData` that represents the biological group or condition of interest (e.g. treatment versus control). Note that this variable should only contain two possible values since `scDD` can currently only handle two-group comparisons. The default option assumes that there is a column named "condition" that contains this variable.

**Value**

Vector of indices for a reasonable set of genes that can be used for simulation.

**References**

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendziorski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

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findOutliers

*findOutliers*

---

**Description**

Find the clusters that are considered outliers

**Usage**

```
findOutliers(clustering, min.size = 3)
```

**Arguments**

clustering	Numeric vector of cluster membership (1st item (named class) in list returned by <code>mclustRestricted</code> )
min.size	Numeric value for the minimum number of samples a cluster must have to be considered in the robust count. Default is 3.

**Details**

Function to obtain a count of the number of clusters that is robust to outliers. Requires at least `min.size` samples to be considered in the robust count.

**Value**

The robust count of the number of unique clusters excluding those with less than `min.size` samples.

**References**

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendziorski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>



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getPosteriorParams      *getPosteriorParams*

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### Description

Given the observations for a single gene and its clustering information, return the calculated posterior parameters

### Usage

```
getPosteriorParams(y, mcobj, alpha, m0, s0, a0, b0)
```

### Arguments

y	Numeric data vector for one gene (log-transformed non-zeroes)
mcobj	Object returned by <code>mclustRestricted</code>
alpha	Value for the Dirichlet concentration parameter
m0	Prior mean value for generating distribution of cluster means
s0	Prior precision value for generating distribution of cluster means
a0	Prior shape parameter value for the generating distribution of cluster precision
b0	Prior scale parameter value for the generating distribution of cluster precision

### Value

A list of posterior parameter values under the DP mixture model framework, given the data and prior parameter values.

### References

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendzioriski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

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jointPosterior      *jointPosterior*

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### Description

Function to obtain the normalized joint posterior of the data and partition.

### Usage

```
jointPosterior(y, mcobj, alpha, m0, s0, a0, b0)
```

**Arguments**

<code>y</code>	Numeric data vector for one gene (log-transformed non-zeros)
<code>mcobj</code>	Object returned by <code>mclustRestricted</code>
<code>alpha</code>	Value for the Dirichlet concentration parameter
<code>m0</code>	Prior mean value for generating distribution of cluster means
<code>s0</code>	Prior precision value for generating distribution of cluster means
<code>a0</code>	Prior shape parameter value for the generating distribution of cluster precision
<code>b0</code>	Prior scale parameter value for the generating distribution of cluster precision

**Details**

Calculates the normalized joint posterior of the data and partition under the Product Partition Model formulation of the Dirichlet Process Mixture model.

**Value**

log joint posterior value

**References**

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendziorski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

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 lu

*lu*


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**Description**

Shortcut for `length(unique())`

**Usage**

`lu(x)`

**Arguments**

<code>x</code>	Numeric vector of cluster membership (1st item (named class) in list returned by <code>mclustRestricted</code> )
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**Details**

Function to obtain a count of the number of clusters

**Value**

The count of the number of unique clusters.

## References

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendzierski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

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luOutlier

*luOutlier*

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## Description

Count the number of clusters with at least `min.size` samples

## Usage

```
luOutlier(x, min.size = 3)
```

## Arguments

<code>x</code>	Numeric vector of cluster membership (1st item (named <code>class</code> ) in list returned by <code>mclustRestricted</code> )
<code>min.size</code>	Numeric value for the minimum number of samples a cluster must have to be considered in the robust count. Default is 3.

## Details

Function to obtain a count of the number of clusters that is robust to outliers. Requires at least `min.size` samples to be considered in the robust count.

## Value

The robust count of the number of unique clusters excluding those with less than `min.size` samples.

## References

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendzierski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

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mclustRestricted	<i>mclustRestricted</i>
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### Description

Function to determine how many normal mixture components are present.

### Usage

```
mclustRestricted(y, restrict = TRUE, min.size)
```

### Arguments

<code>y</code>	Numeric vector of values to fit to a normal mixture model with Mclust.
<code>restrict</code>	Logical indicating whether or not to enforce the restriction on cluster separation based on bimodal index and ratio of largest to smallest variance (see details). If <code>False</code> , then Mclust results as is are returned.
<code>min.size</code>	a positive integer that specifies the minimum size of a cluster (number of cells) for it to be used during the classification step. A clustering with all clusters of size less than <code>min.size</code> is not valid and clusters will be merged if this happens.

### Details

Robust to detecting multiple components that are close together by enforcing that the distance between two clusters of appreciable size (at least 4 samples), have sufficiently high bimodal index (cluster mean difference standardized by average standard deviation and multiplied by a balance factor which is one when clusters are perfectly balanced) and not have variances that differ by more than a ratio of 20. Bimodal index threshold is dependent on sample size to ensure consistent performance in power and type I error of detection of multiple components.

### Value

List object with (1) vector of cluster membership, (2) cluster means, (3) cluster variances, (4) number of model parameters, (5) sample size, (6) BIC of selected model, and (6) loglikelihood of selected model.

### References

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendziorski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

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permMclust	<i>permMclust</i>
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**Description**

Function to obtain bayes factor numerator for permutations of one gene

**Usage**

```
permMclust(y, nperms, condition, remove.zeroes = TRUE, log.transf = TRUE,
  restrict = FALSE, alpha, m0, s0, a0, b0, ref, min.size)
```

**Arguments**

<code>y</code>	Numeric data vector for one gene
<code>nperms</code>	Number of permutations of residuals to evaluate
<code>condition</code>	Vector of condition indicators for each sample
<code>remove.zeroes</code>	Logical indicating whether zeroes need to be removed from <code>y</code>
<code>log.transf</code>	Logical indicating whether the data is in the raw scale (if so, will be log-transformed)
<code>restrict</code>	Logical indicating whether to perform restricted Mclust clustering where close-together clusters are joined.
<code>alpha</code>	Value for the Dirichlet concentration parameter
<code>m0</code>	Prior mean value for generating distribution of cluster means
<code>s0</code>	Prior precision value for generating distribution of cluster means
<code>a0</code>	Prior shape parameter value for the generating distribution of cluster precision
<code>b0</code>	Prior scale parameter value for the generating distribution of cluster precision
<code>ref</code>	one of two possible values in condition; represents the referent category.
<code>min.size</code>	a positive integer that specifies the minimum size of a cluster (number of cells) for it to be used during the classification step. Any clusters containing fewer than <code>min.size</code> cells will be considered an outlier cluster and ignored in the classification algorithm. The default value is three.

**Details**

Obtains bayes factor numerator for data vector `y` representing one gene

**Value**

Bayes factor numerator for the current permutation

**References**

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendzierski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

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permMclustCov      *permMclustCov*

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### Description

Function to obtain bayes factor for permutations of one gene's residuals

### Usage

```
permMclustCov(y, nperms, C, condition, remove.zeroes = TRUE,
  log.transf = TRUE, restrict = FALSE, alpha, m0, s0, a0, b0, ref, min.size)
```

### Arguments

y	Numeric data vector for one gene
nperms	Number of permutations of residuals to evaluate
C	Matrix of confounder variables, where there is one row for each sample and one column for each covariate.
condition	Vector of condition indicators for each sample
remove.zeroes	Logical indicating whether zeroes need to be removed from y
log.transf	Logical indicating whether the data is in the raw scale (if so, will be log-transformed)
restrict	Logical indicating whether to perform restricted Mclust clustering where close-together clusters are joined.
alpha	Value for the Dirichlet concentration parameter
m0	Prior mean value for generating distribution of cluster means
s0	Prior precision value for generating distribution of cluster means
a0	Prior shape parameter value for the generating distribution of cluster precision
b0	Prior scale parameter value for the generating distribution of cluster precision
ref	one of two possible values in condition; represents the referent category.
min.size	a positive integer that specifies the minimum size of a cluster (number of cells) for it to be used during the classification step. Any clusters containing fewer than min.size cells will be considered an outlier cluster and ignored in the classification algorithm. The default value is three.

### Details

Obtains bayes factor numerator for data vector y representing one gene

### Value

Bayes factor numerator for the current permutation

### References

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendziorski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

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permMclustGene      *permMclustGene*

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### Description

Function to obtain bayes factor for all permutations of one gene (not parallelized; to be used when parallelizing over Genes)

### Usage

```
permMclustGene(y, adjust.perms, nperms, condition, remove.zeroes = TRUE,
  log.transf = TRUE, restrict = TRUE, alpha, m0, s0, a0, b0, C, ref,
  min.size)
```

### Arguments

<code>y</code>	Numeric data vector for one gene
<code>adjust.perms</code>	Logical indicating whether or not to adjust the permutation tests for the sample detection rate (proportion of nonzero values). If true, the residuals of a linear model adjusted for detection rate are permuted, and new fitted values are obtained using these residuals.
<code>nperms</code>	Number of permutations of residuals to evaluate
<code>condition</code>	A character object that contains the name of the column in <code>colData</code> that represents the biological group or condition of interest (e.g. treatment versus control). Note that this variable should only contain two possible values since <code>scDD</code> can currently only handle two-group comparisons. The default option assumes that there is a column named "condition" that contains this variable.
<code>remove.zeroes</code>	Logical indicating whether zeroes need to be removed from <code>y</code>
<code>log.transf</code>	Logical indicating whether the data is in the raw scale (if so, will be log-transformed)
<code>restrict</code>	Logical indicating whether to perform restricted Mclust clustering where close-together clusters are joined.
<code>alpha</code>	Value for the Dirichlet concentration parameter
<code>m0</code>	Prior mean value for generating distribution of cluster means
<code>s0</code>	Prior precision value for generating distribution of cluster means
<code>a0</code>	Prior shape parameter value for the generating distribution of cluster precision
<code>b0</code>	Prior scale parameter value for the generating distribution of cluster precision
<code>C</code>	Matrix of confounder variables, where there is one row for each sample and one column for each covariate.
<code>ref</code>	one of two possible values in <code>condition</code> ; represents the referent category.
<code>min.size</code>	a positive integer that specifies the minimum size of a cluster (number of cells) for it to be used during the classification step. Any clusters containing fewer than <code>min.size</code> cells will be considered an outlier cluster and ignored in the classification algorithm. The default value is three.

### Details

Obtains bayes factor for data vector `y` representing one gene

**Value**

Bayes factor numerator for the current permutation

**References**

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendzierski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

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permZero

*permZero*

---

**Description**

Function to generate random permutations of nonzero values.

**Usage**

permZero(m, size, zmat)

**Arguments**

m	Number of permuted sets to generate.
size	Number of samples present in the dataset
zmat	Matrix of indicators of whether the original data value is zero or not. Should contain the same number of rows and columns as original data matrix.

**Details**

Generates random permutations for all genes, where the zeroes are kept fixed (i.e. only permute the nonzero condition labels).

**Value**

a list of length 'm' (nperms) where each item is a 'ngenes' by 'size' matrix

**References**

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendzierski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>



preprocess

*preprocess***Description**

Function to preprocess SingleCellExperiment object (1) to only keep genes with a certain number of nonzero entries, and (2) optionally apply a normalization procedure.

**Usage**

```
preprocess(SCdat, condition = "condition", zero.thresh = 0.9,
           scran_norm = FALSE, median_norm = FALSE)
```

**Arguments**

SCdat	An object of class SingleCellExperiment that contains single-cell expression and metadata. The assays slot contains a named list of matrices, where the normalized counts are housed in the one named normcounts, and unnormalized counts are stored in the one named counts. If either scran_norm or median_norm is set to TRUE, the normcounts slot will be created from the counts slot. The counts and normalized counts matrices should have one row for each gene and one sample for each column. The colData slot should contain a data.frame with one row per sample and columns that contain metadata for each sample. This data.frame should contain a variable that represents biological condition, which is in the form of numeric values (either 1 or 2) that indicates which condition each sample belongs to (in the same order as the columns of normcounts). Optional additional metadata about each cell can also be contained in this data.frame, and additional information about the experiment can be contained in the metadata slot as a list.
condition	A character object that contains the name of the column in colData that represents the biological group or condition of interest (e.g. treatment versus control). Note that this variable should only contain two possible values since scDD can currently only handle two-group comparisons. The default option assumes that there is a column named "condition" that contains this variable.
zero.thresh	A numeric value between 0 and 1 that represents the maximum proportion of zeroes per gene allowable in the processed dataset
scran_norm	Logical indicating whether or not to normalize the data using scran Normalization from scran
median_norm	Logical indicating whether or not to normalize the data using Median Normalization from EBSeq

**Value**

An object of class SingleCellExperiment with genes removed if they have more than zero.thresh zeroes, and the normcounts assay added if either scran\_norm or median\_norm is set to TRUE and only counts is provided. If normcounts already exists and either scran\_norm or median\_norm is set to TRUE, then the new normalized counts are placed in the normcounts assay slot, and the original values are moved to a new slot called normcounts-orig.

## References

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendziorski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

## Examples

```
# load toy example SingleCellExperiment object

data(scDatEx)

# apply the preprocess function to filter out genes if they have more than
# 75% zero

scDatEx <- preprocess(scDatEx, zero.thresh=0.75)

# apply the preprocess function again, but this time threshold on the
# proportion of zeroes and apply scran normalization
# set the zero.thresh argument to 0.75 so that genes with more than 75%
# zeroes are filtered out
# set the scran_norm argument to TRUE to return scran normalized counts

scDatEx.scran <- preprocess(scDatEx, zero.thresh=0.75, scran_norm=TRUE)

# set the median_norm argument to TRUE to return Median normalized counts

scDatEx.median <- preprocess(scDatEx, zero.thresh=0.75, median_norm=TRUE)
```

---

results

*results*

---

## Description

extract results objects after running scDD analysis

## Usage

```
results(SCdat, type = c("Genes", "Zhat.c1", "Zhat.c2", "Zhat.combined"))
```

## Arguments

SCdat	An object of class <code>SingleCellExperiment</code> that contains normalized single-cell expression and metadata, and the output of the <code>scDD</code> function.
type	A character variable specifying which output is desired, with possible values "Genes", "Zhat.c1", "Zhat.c2", and "Zhat.overall". The default value is "Genes", which contains a data frame with nine columns: gene name (matches row-names of SCdat), permutation p-value for testing of independence of condition membership with clustering, Benjamini-Hochberg adjusted version of the previous column, p-value for test of difference in dropout rate (only for non-DD genes), Benjamini-Hochberg adjusted version of the previous column, name of the DD (DE, DP, DM, DB) pattern or DZ (otherwise NS = not significant), the

number of clusters identified overall, the number of clusters identified in condition 1 alone, and the number of clusters identified in condition 2 alone.

If type is "Zhat.c1", then a matrix is returned that contains the fitted cluster memberships (partition estimates  $Z$ ) for each sample (cluster number given by 1,2,3,...) in columns and gene in rows only for condition 1. The same information is returned only for condition 2, and for the overall clustering, when type is set to "Zhat.c2" or "Zhat.overall", respectively. Zeroes, which are not involved in the clustering, are labeled as zero.

### Details

Convenient helper function to extract the results (gene classifications, pvalues, and clustering information). Results data.frames/matrices are stored in the metadata slot and can also be accessed without the help of this convenience function by calling `metadata(SCdat)`.

### Value

A data.frame which contains either the gene classification and p-value results, or cluster membership information, as detailed in the description of the type input parameter.

### Examples

```
# load toy simulated example SingleCellExperiment object to find DD genes
data(scDatExSim)

# set arguments to pass to scDD function

prior_param=list(alpha=0.01, mu0=0, s0=0.01, a0=0.01, b0=0.01)

# call the scDD function to perform permutations and classify DD genes

scDatExSim <- scDD(scDatExSim, prior_param=prior_param, testZeroes=FALSE)

# extract main results object

RES <- results(scDatExSim)
```

---

scDatEx

*Data: Toy example data*

---

### Description

Toy example data in SingleCellExperiment format for 500 genes to illustrate how to generate simulated data from example data using [simulateSet](#).

### Usage

```
data(scDatEx)
```

### Format

An object of class SingleCellExperiment containing data for 500 genes for 142 samples (78 from condition 1 and 64 from condition 2). Condition labels (1 or 2) are stored in the colData slot. The assays slot contains both normcounts and counts for illustration, but these objects are identical.

**Value**

An RData object, see Format section for details

**References**

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendziorski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

**Examples**

```
# load toy example data
data(scDatEx)
```

---

scDatExList

*Data: Toy example data list*

---

**Description**

Toy example data list (one item for each of two conditions) for 100 genes to illustrate how to use the function [preprocess](#).

**Usage**

```
data(scDatExList)
```

**Format**

A list of two matrices (one for each of two conditions) labeled "C1" and "C2". Each matrix contains data for 100 genes and a variable number of samples (78 in C1 and 64 in C2).

**Value**

An RData object, see Format section for details

**References**

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendziorski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

**Examples**

```
# load toy example data list
data(scDatExList)
```

---

 scDatExSim

*Data: Toy example of simulated data*


---

### Description

Toy example data in SingleCellExperiment format for 500 genes to illustrate how to generate simulated data from example data using `simulateSet`. Contains 5 genes from each category (DE, DP, DM, DB, EE, and EP).

### Usage

```
data(scDatExSim)
```

### Format

An object of class `SingleCellExperiment` containing data for 30 genes for 200 samples (100 from condition 1 and 100 from condition 2). Condition labels (1 or 2) are stored in the `colData` slot. Row names of the `assayData` slot contain the two letter category label that the gene was simulated from (e.g. 'EE', 'DB', ...) along with the row number (1-30).

### Value

An RData object, see Format section for details

### References

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendzierski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

### Examples

```
# load toy example of simulated data
data(scDatExSim)
```

---

 scDD

*scDD*


---

### Description

Find genes with differential distributions (DD) across two conditions

### Usage

```
scDD(SCdat, prior_param = list(alpha = 0.1, mu0 = 0, s0 = 0.01, a0 = 0.01, b0 = 0.01),
      permutations = 0, testZeroes = TRUE, adjust.perms = FALSE,
      param = bpparam(), parallelBy = c("Genes", "Permutations"),
      condition = "condition", min.size = 3, min.nonzero = NULL,
      level = 0.05, categorize = TRUE)
```

## Arguments

SCdat	An object of class <code>SingleCellExperiment</code> that contains normalized single-cell expression and metadata. The <code>assays</code> slot contains a named list of matrices, where the normalized counts are housed in the one named <code>normcounts</code> . This matrix should have one row for each gene and one sample for each column. The <code>colData</code> slot should contain a <code>data.frame</code> with one row per sample and columns that contain metadata for each sample. This <code>data.frame</code> should contain a variable that represents biological condition, which is in the form of numeric values (either 1 or 2) that indicates which condition each sample belongs to (in the same order as the columns of <code>normcounts</code> ). Optional additional metadata about each cell can also be contained in this <code>data.frame</code> , and additional information about the experiment can be contained in the <code>metadata</code> slot as a list.
prior_param	A list of prior parameter values to be used when modeling each gene as a mixture of DP normals. Default values are given that specify a vague prior distribution on the cluster-specific means and variances.
permutations	The number of permutations to be used in calculating empirical p-values. If set to zero (default), the full Bayes Factor permutation test will not be performed. Instead, a fast procedure to identify the genes with significantly different expression distributions will be performed using the nonparametric Kolmogorov-Smirnov test, which tests the null hypothesis that the samples are generated from the same continuous distribution. This test will yield slightly lower power than the full permutation testing framework (this effect is more pronounced at smaller sample sizes, and is more pronounced in the DB category), but is orders of magnitude faster. This option is recommended when compute resources are limited. The remaining steps of the scDD framework will remain unchanged (namely, categorizing the significant DD genes into patterns that represent the major distributional changes, as well as the ability to visualize the results with violin plots using the <code>sideViolin</code> function).
testZeroes	Logical indicating whether or not to test for a difference in the proportion of zeroes. This will only be done for genes that have at least one zero value (genes where all cells have a nonzero value will have a 'zero.pvalue' of NA).
adjust.perms	Logical indicating whether or not to adjust the permutation tests for the sample detection rate (proportion of nonzero values). If true, the residuals of a linear model adjusted for detection rate are permuted, and new fitted values are obtained using these residuals.
param	a <code>MulticoreParam</code> or <code>SnowParam</code> object of the <code>BiocParallel</code> package that defines a parallel backend. The default option is <code>BiocParallel::bpparam()</code> which will automatically create a cluster appropriate for the operating system. Alternatively, the user can specify the number of cores they wish to use by first creating the corresponding <code>MulticoreParam</code> (for Linux-like OS) or <code>SnowParam</code> (for Windows) object, and then passing it into the <code>scDD</code> function. This could be done to specify a parallel backend on a Linux-like OS with, say 12 cores by setting <code>param=BiocParallel::MulticoreParam(workers=12)</code>
parallelBy	For the permutation test (if invoked), the manner in which to parallelize. The default option is "Genes" which will spawn processes that divide up the genes across all cores defined in <code>param</code> cores, and then loop through the permutations. The alternate option is "Permutations" which loop through each gene and spawn processes that divide up the permutations across all cores defined in <code>param</code> . The default option is recommended when analyzing more genes than the number of permutations.

condition	A character object that contains the name of the column in colData that represents the biological group or condition of interest (e.g. treatment versus control). Note that this variable should only contain two possible values since scDD can currently only handle two-group comparisons. The default option assumes that there is a column named "condition" that contains this variable.
min.size	a positive integer that specifies the minimum size of a cluster (number of cells) for it to be used during the classification step. Any clusters containing fewer than min.size cells will be considered an outlier cluster and ignored in the classification algorithm. The default value is three.
min.nonzero	a positive integer that specifies the minimum number of nonzero cells in each condition required for the test of differential distributions. If a gene has fewer nonzero cells per condition, it will still be tested for DZ (if testZeroes is TRUE). Default value is NULL (no minimum value is enforced).
level	numeric value between 0 and 1 that specifies the alpha level for significance of a differential gene test (default value 0.05). This is used to decide whether to classify a gene into one of the differential patterns. If 'testZeroes' is FALSE and the adjusted p-value for a given gene is below 'level', then the gene is categorized. Alternatively, if 'testZeroes' is TRUE, then the adjusted p-value must be below 'level/2' in order to be considered significant and categorized. This is done to control for multiple testing since 'testZeroes=TRUE' means that each gene is tested for a difference in nonzeros and zeroes separately.
categorize	a logical indicating whether to determine which categories (DE, DP, DM, DB) each gene belongs to (default = TRUE). This can only be set to FALSE if 'permutations' is set to zero, since the full model fitting will automatically be carried out if permutations are run.

## Details

Find genes with differential distributions (DD) across two conditions. Models each log-transformed gene as a Dirichlet Process Mixture of normals and uses a permutation test to determine whether condition membership is independent of sample clustering. The FDR adjusted (Benjamini-Hochberg) permutation p-value is returned along with the classification of each significant gene (with p-value less than 0.05 (or 0.025 if also testing for a difference in the proportion of zeroes)) into one of four categories (DE, DP, DM, DB). For genes that do not show significant influence, of condition on clustering, an optional test of whether the proportion of zeroes (dropout rate) is different across conditions is performed (DZ).

## Value

A SingleCellExperiment object that contains the data and sample information from the input object, but where the results objects are now added to the metadata slot. The metadata slot is now a list with four items: the first (main results object) is a data.frame with the following columns:

- 'gene': gene name (matches rownames of SCdat)
- 'DDcategory': name of the DD (DE, DP, DM, DB, DZ) pattern (or NS = not significant)
- 'Clusters.combined': the number of clusters identified overall
- 'Clusters.C1': the number of clusters identified in condition 1 alone
- 'Clusters.C2': the number of clusters identified in condition 2 alone
- 'nonzero.pvalue': permutation (or KS) p-value for testing difference in nonzero expression values

- ‘nonzero.pvalue.adj’: Benjamini-Hochberg adjusted version of the ‘nonzero.pvalue’ column
- ‘zero.pvalue’: p-value for test of difference in dropout rate (only if ‘testZeroes’ is TRUE)
- ‘zero.pvalue’: Benjamini-Hochberg adjusted version of the previous column (only if ‘testZeroes’ is TRUE)
- ‘combined.pvalue’: Fisher’s combined p-value for a difference in nonzero or zero values (only if ‘testZeroes’ is TRUE).
- ‘combined.pvalue.adj’: Benjamini-Hochberg adjusted version of the previous column (only if ‘testZeroes’ is TRUE)

The remaining three elements are matrices (first for condition 1 and 2 combined, then condition 1 alone, then condition 2 alone) that contains the cluster memberships for each sample (cluster 1,2,3,...) in columns and genes in rows. Zeroes, which are not involved in the clustering, are labeled as zero. See the results function for a convenient way to extract these results objects.

## References

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendzioriski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

## Examples

```
# load toy simulated example SingleCellExperiment object to find DD genes
data(scDatExSim)

# check that this object is a member of the SingleCellExperiment class
# and that it contains 200 samples and 30 genes

class(scDatExSim)
show(scDatExSim)

# set arguments to pass to scDD function
# we will perform 100 permutations on each of the 30 genes

prior_param=list(alpha=0.01, mu0=0, s0=0.01, a0=0.01, b0=0.01)
nperms <- 100

# call the scDD function to perform permutations, classify DD genes,
# and return results
# we won't perform the test for a difference in the proportion of zeroes
# since none exists in this simulated toy example data
# this step will take significantly longer with more genes and/or
# more permutations

scDatExSim <- scDD(scDatExSim, prior_param=prior_param, permutations=nperms,
                  testZeroes=FALSE)
```



---

sideHist	<i>sideHist</i>
----------	-----------------

---

**Description**

Plots two histograms side by side with smoothed density overlay

**Usage**

```
sideHist(x, y, logT = TRUE, title.gene = "")
```

**Arguments**

x	First numeric vector of data to plot.
y	Second numeric vector of data to plot.
logT	Logical that indicates whether to take the log(x+1) transformation.
title.gene	Character vector that contains the gene name that you are plotting

**Value**

NULL (creates a baseR plot)

**References**

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendzierski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

---

sideViolin	<i>sideViolin</i>
------------	-------------------

---

**Description**

Plots two histograms side by side with smoothed density overlay

**Usage**

```
sideViolin(y, cond, MAP = NULL, logT = TRUE, title.gene = "",  
           conditionLabels = unique(cond), axes.titles = TRUE)
```

**Arguments**

<code>y</code>	Numeric vector of data to plot.
<code>cond</code>	Vector of condition labels corresponding to elements of <code>x</code> .
<code>MAP</code>	List of MAP partition estimates with conditions as list items and samples as elements (integer indicating which cluster each observation belongs to; zeroes belong to cluster 1)
<code>logT</code>	Logical that indicates whether to take the $\log(x+1)$ transformation.
<code>title.gene</code>	Character vector that contains the gene name that you are plotting.
<code>conditionLabels</code>	Character vector containing the names of the two conditions.
<code>axes.titles</code>	Logical indicating whether or not to include axes labels on plots.

**Value**

ggplot object

**References**

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendzierski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

**Examples**

```
# load toy simulated example ExpressionSet to find DD genes
data(scDatExSim)

# load SingleCellExperiment package to facilitate subset operations
library(SingleCellExperiment)

# plot side by side violin plots for Gene 1 (DE)
sideViolin(normcounts(scDatExSim)[1,], scDatExSim$condition,
            title.gene=rownames(scDatExSim)[1])

# plot side by side violin plots for Gene 6 (DP)
sideViolin(normcounts(scDatExSim)[6,], scDatExSim$condition,
            title.gene=rownames(scDatExSim)[6])

# plot side by side violin plots for Gene 11 (DM)
sideViolin(normcounts(scDatExSim)[11,], scDatExSim$condition,
            title.gene=rownames(scDatExSim)[11])
```

```
# plot side by side violin plots for Gene 16 (DB)

sideViolin(normcounts(scDatExSim)[16,], scDatExSim$condition,
            title.gene=rownames(scDatExSim)[16])

# plot side by side violin plots for Gene 21 (EP)

sideViolin(normcounts(scDatExSim)[21,], scDatExSim$condition,
            title.gene=rownames(scDatExSim)[21])

# plot side by side violin plots for Gene 26 (EE)

sideViolin(normcounts(scDatExSim)[26,], scDatExSim$condition,
            title.gene=rownames(scDatExSim)[26])
```

---

simuDB

*simuDB*


---

## Description

Simulation for Differential "Both" Case - both Differential Modality and Differential Mean

## Usage

```
simuDB(Dataset1, Simulated_Data, DEIndex, samplename, Zeropercent_Base, f, FC,
       coeff, RP, modeFC, DP, generateZero, constantZero, varInflation)
```

## Arguments

Dataset1	Numeric matrix of expression values with genes in rows and samples in columns.
Simulated_Data	Required input empty matrix to provide structure information of output matrix with simulated data
DEIndex	Index for DE genes
samplename	The name for genes that chosen for simulation
Zeropercent_Base	Zero percentage for corresponding gene expression values
f	Fold change values (number of SDs) for each gene
FC	Fold Change values for DE Simulation
coeff	Relationship coefficients for Mean and Variance
RP	matrix for NB parameters for genes in samplename
modeFC	Vector of values to use for fold changes between modes for DP, DM, and DB.
DP	Differetial Proportion vector
generateZero	Specification of how to generate the zero values. If "empirical" (default), the observed proportion of zeroes in each gene is used for the simulated data, and the nonzeroes are simulated from a truncated negative binomial distribution. If "simulated", all values are simulated out of a negative binomial distribution, including the zeroes. If "constant", then each gene has a fixed proportion of zeroes equal to constantZero.

constantZero	Numeric value between 0 and 1 that indicates the fixed proportion of zeroes for every gene. Ignored if generateZero method is not equal to "constant".
varInflation	Optional numeric vector with one element for each condition that corresponds to the multiplicative variance inflation factor to use when simulating data. Useful for sensitivity studies to assess the impact of confounding effects on differential variance across conditions. Currently assumes all samples within a condition are subject to the same variance inflation factor.

### Value

Simulated\_Data Simulated dataset for DB

### References

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendziorski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

---

simuDE

*simuDE*

---

### Description

Simulation for Classic Differentially Expressed Case.

### Usage

```
simuDE(Dataset1, Simulated_Data, DEIndex, samplename, Zeropercent_Base, f, FC,
      coeff, RP, modeFC, generateZero, constantZero, varInflation)
```

### Arguments

Dataset1	Numeric matrix of expression values with genes in rows and samples in columns.
Simulated_Data	Required input empty matrix to provide structure information of output matrix with simulated data
DEIndex	Index for DE genes
samplename	The name for genes that chosen for simulation
Zeropercent_Base	Zero percentage for corresponding gene expression values
f	Fold change values (number of SDs) for each gene
FC	Fold Change values for DE Simulation
coeff	Relationship coefficients for Mean and Variance
RP	matrix for NB parameters for genes in samplename
modeFC	Vector of values to use for fold changes between modes for DP, DM, and DB.

generateZero	Specification of how to generate the zero values. If "empirical" (default), the observed proportion of zeroes in each gene is used for the simulated data, and the nonzeros are simulated from a truncated negative binomial distribution. If "simulated", all values are simulated out of a negative binomial distribution, including the zeroes. If "constant", then each gene has a fixed proportion of zeroes equal to constantZero.
constantZero	Numeric value between 0 and 1 that indicates the fixed proportion of zeroes for every gene. Ignored if generateZero method is not equal to "constant".
varInflation	Optional numeric vector with one element for each condition that corresponds to the multiplicative variance inflation factor to use when simulating data. Useful for sensitivity studies to assess the impact of confounding effects on differential variance across conditions. Currently assumes all samples within a condition are subject to the same variance inflation factor.

### Details

Method called by main function `singleCellSimu` to simulate genes that have different means in each condition. Not intended to be called directly by user.

### Value

Simulated\_Data Simulated dataset for DE

### References

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendziorski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

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simuDM	<i>simuDM</i>
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---

### Description

Simulation for Differential Modalities Case

### Usage

```
simuDM(Dataset1, Simulated_Data, DEIndex, samplename, Zeropercent_Base, f, FC,
       coeff, RP, modeFC, generateZero, constantZero, varInflation)
```

### Arguments

Dataset1	Numeric matrix of expression values with genes in rows and samples in columns.
Simulated_Data	Required input empty matrix to provide structure information of output matrix with simulated data
DEIndex	Index for DE genes
samplename	The name for genes that chosen for simulation

Zeropercent_Base	Zero percentage for corresponding gene expression values
f	Fold change values (number of SDs) for each gene
FC	Fold Change values for DE Simulation
coeff	Relationship coefficients for Mean and Variance
RP	matrix for NB parameters for genes in samplename
modeFC	Vector of values to use for fold changes between modes for DP, DM, and DB.
generateZero	Specification of how to generate the zero values. If "empirical" (default), the observed proportion of zeroes in each gene is used for the simulated data, and the nonzeros are simulated from a truncated negative binomial distribution. If "simulated", all values are simulated out of a negative binomial distribution, including the zeroes. If "constant", then each gene has a fixed proportion of zeroes equal to constantZero.
constantZero	Numeric value between 0 and 1 that indicates the fixed proportion of zeroes for every gene. Ignored if generateZero method is not equal to "constant".
varInflation	Optional numeric vector with one element for each condition that corresponds to the multiplicative variance inflation factor to use when simulating data. Useful for sensitivity studies to assess the impact of confounding effects on differential variance across conditions. Currently assumes all samples within a condition are subject to the same variance inflation factor.

## Value

Simulated\_Data Simulated dataset for DM

## References

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendziorski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

---

simuDP

*simuDP*

---

## Description

Simulation for Differential Proportion Case

## Usage

```
simuDP(Dataset1, Simulated_Data, DEIndex, samplename, Zeropercent_Base, f, FC,
coeff, RP, modeFC, DP, generateZero, constantZero, varInflation)
```

**Arguments**

Dataset1	Numeric matrix of expression values with genes in rows and samples in columns.
Simulated_Data	Required input empty matrix to provide structure information of output matrix with simulated data
DEIndex	Index for DE genes
samplename	The name for genes that chosen for simulation
Zeropercent_Base	Zero percentage for corresponding gene expression values
f	Fold change values (number of SDs) for each gene
FC	Fold Change values for DE Simulation
coeff	Relationship coefficients for Mean and Variance
RP	matrix for NB parameters for genes in samplename
modeFC	Vector of values to use for fold changes between modes for DP, DM, and DB.
DP	Differetial Proportion vector
generateZero	Specification of how to generate the zero values. If "empirical" (default), the observed proportion of zeroes in each gene is used for the simuated data, and the nonzeroes are simulated from a truncated negative binomial distribution. If "simulated", all values are simulated out of a negative binomial distribution, including the zeroes. If "constant", then each gene has a fixed proportion of zeroes equal to constantZero.
constantZero	Numeric value between 0 and 1 that indicates the fixed proportion of zeroes for every gene. Ignored if generateZero method is not equal to "constant".
varInflation	Optional numeric vector with one element for each condition that corresponds to the multiplicative variance inflation factor to use when simulating data. Useful for sensitivity studies to assess the impact of confounding effects on differential variance across conditions. Currently assumes all samples within a condition are subject to the same variance inflation factor.

**Value**

Simulated\_Data Simulated dataset for DP

**References**

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendziorski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

---

simulateSet

*simulateSet*

---

**Description**

Simulation of a complete dataset, where the number of each type of differential distributions and equivalent distributions is specified.

**Usage**

```
simulateSet(SCdat, numSamples = 100, nDE = 250, nDP = 250, nDM = 250,
  nDB = 250, nEE = 5000, nEP = 4000, sd.range = c(1, 3), modeFC = c(2,
  3, 4), plots = TRUE, plot.file = NULL, random.seed = 284,
  varInflation = NULL, condition = "condition", param = bpparam())
```

**Arguments**

SCdat	An object of class <code>SingleCellExperiment</code> that contains normalized single-cell expression and metadata. The <code>assays</code> slot contains a named list of matrices, where the normalized counts are housed in the one named <code>normcounts</code> . This matrix should have one row for each gene and one sample for each column. The <code>colData</code> slot should contain a <code>data.frame</code> with one row per sample and columns that contain metadata for each sample. This <code>data.frame</code> should contain a variable that represents biological condition, which is in the form of numeric values (either 1 or 2) that indicates which condition each sample belongs to (in the same order as the columns of <code>normcounts</code> ). Optional additional metadata about each cell can also be contained in this <code>data.frame</code> , and additional information about the experiment can be contained in the <code>metadata</code> slot as a list.
numSamples	numeric value for the number of samples in each condition to simulate
nDE	Number of DE genes to simulate
nDP	Number of DP genes to simulate
nDM	Number of DM genes to simulate
nDB	Number of DB genes to simulate
nEE	Number of EE genes to simulate
nEP	Number of EP genes to simulate
sd.range	Numeric vector of length two which describes the interval (lower, upper) of standard deviations of fold changes to randomly select.
modeFC	Vector of values to use for fold changes between modes for DP, DM, and DB.
plots	Logical indicating whether or not to generate fold change and validation plots
plot.file	Character containing the file string if the plots are to be sent to a pdf instead of to the standard output.
random.seed	Numeric value for a call to <code>set.seed</code> for reproducibility.
varInflation	Optional numeric vector with one element for each condition that corresponds to the multiplicative variance inflation factor to use when simulating data. Useful for sensitivity studies to assess the impact of confounding effects on differential variance across conditions. Currently assumes all samples within a condition are subject to the same variance inflation factor.
condition	A character object that contains the name of the column in <code>colData</code> that represents the biological group or condition of interest (e.g. treatment versus control). Note that this variable should only contain two possible values since <code>scDD</code> can currently only handle two-group comparisons. The default option assumes that there is a column named "condition" that contains this variable.
param	a <code>MulticoreParam</code> or <code>SnowParam</code> object of the <code>BiocParallel</code> package that defines a parallel backend. The default option is <code>BiocParallel::bpparam()</code> which will automatically creates a cluster appropriate for the operating system. Alternatively, the user can specify the number of cores they wish to use by first



creating the corresponding MulticoreParam (for Linux-like OS) or SnowParam (for Windows) object, and then passing it into the scDD function. This could be done to specify a parallel backend on a Linux-like OS with, say 12 cores by setting `param=BiocParallel::MulticoreParam(workers=12)`

### Value

An object of class `SingleCellExperiment` that contains simulated single-cell expression and meta-data. The `assays` slot contains a named list of matrices, where the simulated counts are housed in the one named `normcounts`. This matrix should have one row for each gene ( $n_{DE} + n_{DP} + n_{DM} + n_{DB} + n_{EE} + n_{EP}$  rows) and one sample for each column (`numSamples` columns). The `colData` slot contains a `data.frame` with one row per sample and a column that represents biological condition, which is in the form of numeric values (either 1 or 2) that indicates which condition each sample belongs to (in the same order as the columns of `normcounts`). The `rowData` slot contains information about the category of the gene (EE, EP, DE, DM, DP, or DB), as well as the simulated foldchange value.

### References

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendziorski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

### Examples

```
# Load toy example ExpressionSet to simulate from

data(scDatEx)

# check that this object is a member of the ExpressionSet class
# and that it contains 142 samples and 500 genes

class(scDatEx)
show(scDatEx)

# set arguments to pass to simulateSet function
# we will simulate 30 genes total; 5 genes of each type;
# and 100 samples in each of two conditions

nDE <- 5
nDP <- 5
nDM <- 5
nDB <- 5
nEE <- 5
nEP <- 5
numSamples <- 100
seed <- 816

# create simulated set with specified numbers of DE, DP, DM, DM, EE, and
# EP genes,
# specified number of samples, DE genes are 2 standard deviations apart, and
# multimodal genes have modal distance of 4 standard deviations
```

```
SD <- simulateSet(scDatEx, numSamples=numSamples, nDE=nDE, nDP=nDP, nDM=nDM,
                 nDB=nDB, nEE=nEE, nEP=nEP, sd.range=c(2,2), modeFC=4,
                 plots=FALSE,
                 random.seed=seed)
```

---

singleCellSimu      *singleCellSimu*

---

## Description

Called by `simulateSet` to simulate a specified number of genes from one DD category at a time.

## Usage

```
singleCellSimu(Dataset1, Method, index, FC, modeFC, DP, Validation = FALSE,
              numGenes = 1000, numDE = 100, numSamples = 100,
              generateZero = c("empirical", "simulated", "constant"),
              constantZero = NULL, varInflation = NULL)
```

## Arguments

Dataset1	Numeric matrix of expression values with genes in rows and samples in columns.
Method	Type of simulation should choose from "DE" "DP" "DM" "DB"
index	Reasonable set of genes for simulation
FC	Fold Change values for DE Simulation
modeFC	Vector of values to use for fold changes between modes for DP, DM, and DB.
DP	Differential Proportion vector
Validation	Show Validation plots or not
numGenes	numeric value for the number of genes to simulate
numDE	numeric value for the number of genes that will differ between two conditions
numSamples	numeric value for the number of samples in each condition to simulate
generateZero	Specification of how to generate the zero values. If "empirical" (default), the observed proportion of zeroes in each gene is used for the simulated data, and the nonzeros are simulated from a truncated negative binomial distribution. If "simulated", all values are simulated out of a negative binomial distribution, including the zeroes. If "constant", then each gene has a fixed proportion of zeroes equal to constantZero.
constantZero	Numeric value between 0 and 1 that indicates the fixed proportion of zeroes for every gene. Ignored if generateZero method is not equal to "constant".
varInflation	Optional numeric vector with one element for each condition that corresponds to the multiplicative variance inflation factor to use when simulating data. Useful for sensitivity studies to assess the impact of confounding effects on differential variance across conditions. Currently assumes all samples within a condition are subject to the same variance inflation factor.

## Value

Simulated\_Data A list object where the first element contains a matrix of the simulated dataset, the second element contains the DEIndex, and the third element contains the fold change (between two conditions for DE, between two modes for DP, DM, and DB).

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testKS	<i>testKS</i>
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**Description**

Function to perform KS test

**Usage**

```
testKS(dat, condition, inclZero = TRUE, numDE = NULL, DEIndex)
```

**Arguments**

dat	Matrix of single-cell RNA-seq data with genes in rows and samples in columns.
condition	Vector containing the indicator of which condition each sample (in the columns of dat) belongs to.
inclZero	Logical indicating whether to include zero in the test of different distributions
numDE	numeric value for the number of genes that will differ between two conditions
DEIndex	Vector containing the row numbers of the DE genes

**Value**

List object containing the significant gene indices, their adjusted p-values, and (if DE genes are supplied) the power and fdr.

**References**

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendziorski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

**Examples**

```
# load toy simulated example ExpressionSet to find KS genes
data(scDatExSim)

# load SingleCellExperiment package to facilitate subset operations
library(SingleCellExperiment)

# check that this object is a member of the ExpressionSet class
# and that it contains 200 samples and 30 genes

class(scDatExSim)
show(scDatExSim)

# perform KS test and obtain adjusted p-values
RES_KS <- testKS(normcounts(scDatExSim), scDatExSim$condition, inclZero=FALSE,
                 numDE=20, DEIndex=1:20)
```

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testZeroes	<i>testZeroes</i>
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**Description**

Test for a difference in the proportion of zeroes between conditions for a specified set of genes

**Usage**

```
testZeroes(dat, cond, these = 1:nrow(dat))
```

**Arguments**

dat	Matrix of single cell expression data with genes in rows and samples in columns.
cond	Vector of condition labels
these	vector of row numbers (gene numbers) to test for a difference in the proportion of zeroes.

**Details**

Test for a difference in the proportion of zeroes between conditions that is not explained by the detection rate. Utilizes Bayesian logistic regression.

**Value**

Vector of FDR adjusted p-values

---

validation	<i>validation</i>
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---

**Description**

Draw validation plots to show that the simulated dataset emulates characteristics of observed dataset.

**Usage**

```
validation(MV, DEIndex, Zeropercent_Base, Simulated_Data, numGenes)
```

**Arguments**

MV	Mean and Variance matrix for observed data
DEIndex	Index for genes chosen to be DE (can be NULL)
Zeropercent_Base	Zero percentage for corresponding gene expression values
Simulated_Data	Simulated dataset
numGenes	numeric value for the number of genes to simulate

**Value**

Validation plots

**References**

Korthauer KD, Chu LF, Newton MA, Li Y, Thomson J, Stewart R, Kendzierski C. A statistical approach for identifying differential distributions in single-cell RNA-seq experiments. *Genome Biology*. 2016 Oct 25;17(1):222. <https://genomebiology.biomedcentral.com/articles/10.1186/s13059-016-1077-y>

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