Package 'scDblFinder'

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Description The scDblFinder package gathers various methods for the detection and
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     multiple cells captured within the same droplet or reaction volume). It
     includes methods formerly found in the scran package, and the new fast
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```

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 ${\sf addDoublets}$

addDoublets

Description

Adds artificial doublets to an existing dataset

Usage

```
addDoublets(
    x,
    clusters,
    dbr = (0.01 * ncol(x)/1000),
    only.heterotypic = TRUE,
    adjustSize = FALSE,
    prefix = "doublet.",
    ...
)
```

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Arguments

A count matrix of singlets, or a SummarizedExperiment-class

clusters A vector of cluster labels for each column of 'x'

dbr The doublet rate
only.heterotypic
Whether to add only heterotypic doublets.

adjustSize Whether to adjust the library sizes of the doublets.

prefix Prefix for the colnames generated.

Any further arguments to createDoublets.

Value

A 'SingleCellExperiment' with the colData columns 'cluster' and 'type' (indicating whether the cell is a singlet or doublet).

Examples

```
sce <- mockDoubletSCE(dbl.rate=0)
sce <- addDoublets(sce, clusters=sce$cluster)</pre>
```

aggregateFeatures

aggregateFeatures

Description

Aggregates similar features (rows).

Usage

```
aggregateFeatures(
    x,
    dims.use = seq(2L, 12L),
    k = 1000,
    num_init = 2,
    use.mbk = NULL,
    use.subset = 5000,
    use.TFIDF = TRUE,
    ...
)
```

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Arguments

X	A integer/numeric (sparse) matrix, or a 'SingleCellExperiment' including a 'counts' assay.
dims.use	The PCA dimensions to use for clustering rows.
k	The approximate number of meta-features desired
num_init	The number of initializations used for k-means clustering.
use.mbk	Logical; whether to use minibatch k-means (see mbkmeans). If NULL, the minibatch approach will be used if there are more than 30000 features.
use.subset	How many cells (columns) to use to cluster the features.
use.TFIDF	Logical; whether to use TFIDF normalization (instead of standard normalization) to assess the similarity between features. similarity
	Passed to mbkmeans. Can for instance be used to pass the 'BPPARAM' argument for multithreading.

Value

An aggregated version of 'x' (either an array or a 'SingleCellExperiment', depending on the input).

Description

Tests for enrichment of doublets created from each cluster (i.e. cluster's stickiness). Only applicable with >=4 clusters. Note that when applied to an multisample object, this functions assumes that the cluster labels match across samples.

Usage

```
clusterStickiness(
    x,
    type = c("quasibinomial", "nbinom1", "binomial", "poisson"),
    inclDiff = FALSE
)
```

Arguments

X	A table of double statistics, or a SingleCellExperiment on which scDblFinder was run.
type	The type of test to use (quasibinomial recommended).
inclDiff	Logical; whether to include the difficulty in the model. If NULL, will be used only if there is a significant trend with the enrichment.

Value

A table of test results for each cluster.

Examples

```
sce <- mockDoubletSCE(rep(200,5))
sce <- scDblFinder(sce, artificialDoublets=500)
clusterStickiness(sce)</pre>
```

computeDoubletDensity Compute the density of simulated doublets

Description

Identify potential doublet cells based on the local density of simulated doublet expression profiles. This replaces the older doubletCells function from the **scran** package.

Usage

```
computeDoubletDensity(x, ...)
## S4 method for signature 'ANY'
computeDoubletDensity(
  size.factors.norm = NULL,
  size.factors.content = NULL,
  k = 50,
  subset.row = NULL,
  niters = max(10000, ncol(x)),
  block = 10000,
  dims = 25,
  BNPARAM = KmknnParam(),
 BSPARAM = bsparam(),
  BPPARAM = SerialParam()
)
## S4 method for signature 'SummarizedExperiment'
computeDoubletDensity(x, ..., assay.type = "counts")
## S4 method for signature 'SingleCellExperiment'
computeDoubletDensity(x, size.factors.norm = sizeFactors(x), ...)
```

Arguments

Х

A numeric matrix-like object of count values, where each column corresponds to a cell and each row corresponds to an endogenous gene.

Alternatively, a SummarizedExperiment or SingleCellExperiment object containing such a matrix.

For the generic, additional arguments to pass to specific methods. . . .

> For the SummarizedExperiment and SingleCellExperiment methods, additional arguments to pass to the ANY method.

size.factors.norm

A numeric vector of size factors for normalization of x prior to PCA and distance calculations. If NULL, defaults to size factors derived from the library sizes of x.

For the SingleCellExperiment method, the default values are taken from sizeFactors(x), if they are available.

size.factors.content

A numeric vector of size factors for RNA content normalization of x prior to simulating doublets. This is orthogonal to the values in size.factors.norm,

see Details.

k An integer scalar specifying the number of nearest neighbours to use to deter-

mine the bandwidth for density calculations.

subset.row See ?"scran-gene-selection".

niters An integer scalar specifying how many simulated doublets should be generated. block

An integer scalar controlling the rate of doublet generation, to keep memory

usage low.

dims An integer scalar specifying the number of components to retain after the PCA.

BNPARAM A BiocNeighborParam object specifying the nearest neighbor algorithm. This

should be an algorithm supported by findNeighbors.

BSPARAM A BiocSingularParam object specifying the algorithm to use for PCA, if d is not

NA.

A BiocParallelParam object specifying whether the neighbour searches should **BPPARAM**

be parallelized.

assay.type A string specifying which assay values contain the count matrix.

Details

This function simulates doublets by adding the count vectors for two randomly chosen cells in x. For each original cell, we compute the density of neighboring simulated doublets and compare it to the density of neighboring original cells. Genuine doublets should have a high density of simulated doublets relative to the density of its neighbourhood. Thus, the doublet score for each cell is defined as the ratio of densities of simulated doublets to the density of the original cells.

Densities are calculated in low-dimensional space after a PCA on the log-normalized expression matrix of x. Simulated doublets are projected into the low-dimensional space using the rotation vectors computed from the original cells. For each cell, the density of simulated doublets is computed for a hypersphere with radius set to the median distance to the k nearest neighbour. This is normalized by niters, k and the total number of cells in x to yield the final score.

The two size factor arguments have different roles:

• size.factors.norm contains the size factors to be used for normalization prior to PCA and distance calculations. This defaults to the values returned by librarySizeFactors but can be explicitly set to ensure that the low-dimensional space is consistent with that in the rest of the analysis.

• size.factors.content is much more important, and represents the size factors that preserve RNA content differences. This is usually computed from spike-in RNA and ensures that the simulated doublets have the correct ratio of contributions from the original cells.

It is possible to set both of these arguments as they are orthogonal to each other. Setting size.factors.content will not affect the calculation of log-normalized expression values from x. Conversely, setting size.factors.norm will not affect the ratio in which cells are added together when simulating doublets.

Value

A numeric vector of doublet scores for each cell in x.

Author(s)

Aaron Lun

References

Lun ATL (2018). Detecting doublet cells with *scran*. https://ltla.github.io/SingleCellThoughts/software/doublet_detection/bycell.html

See Also

findDoubletClusters, to detect doublet clusters.

scDblFinder, which uses a hybrid approach involving simulation and overclustering.

More detail on the mathematical background of this function is provided in the corresponding vignette at vignette("computeDoubletDensity",package="scDblFinder").

```
# Mocking up an example.
set.seed(100)
ngenes <- 1000
mu1 <- 2^rnorm(ngenes)</pre>
mu2 <- 2^rnorm(ngenes)</pre>
mu3 <- 2^rnorm(ngenes)</pre>
mu4 <- 2^rnorm(ngenes)</pre>
counts.1 <- matrix(rpois(ngenes*100, mu1), nrow=ngenes) # Pure type 1</pre>
counts.2 <- matrix(rpois(ngenes*100, mu2), nrow=ngenes) # Pure type 2</pre>
counts.3 <- matrix(rpois(ngenes*100, mu3), nrow=ngenes) # Pure type 3</pre>
counts.4 <- matrix(rpois(ngenes*100, mu4), nrow=ngenes) # Pure type 4</pre>
counts.m <- matrix(rpois(ngenes*20, mu1+mu2), nrow=ngenes) # Doublets (1 & 2)</pre>
counts <- cbind(counts.1, counts.2, counts.3, counts.4, counts.m)</pre>
clusters <- rep(1:5, c(rep(100, 4), ncol(counts.m)))</pre>
# Find potential doublets.
scores <- computeDoubletDensity(counts)</pre>
boxplot(split(log10(scores), clusters))
```

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createDoublets

createDoublets

Description

Creates artificial doublet cells by combining given pairs of cells

Usage

```
createDoublets(
   x,
   dbl.idx,
   clusters = NULL,
  resamp = 0.5,
   halfSize = 0.5,
   adjustSize = FALSE,
   prefix = "dbl."
)
```

Arguments

Χ	A count matrix of real cells
dbl.idx	A matrix or data.frame with pairs of cell indexes stored in the first two columns.
clusters	An optional vector of cluster labels (for each column of 'x')
resamp	Logical; whether to resample the doublets using the poisson distribution. Alternatively, if a proportion between 0 and 1, the proportion of doublets to resample.
halfSize	Logical; whether to half the library size of doublets (instead of just summing up the cells). Alternatively, a number between 0 and 1 can be given, determining the proportion of the doublets for which to perform the size adjustment. Ignored if not resampling.
adjustSize	Logical; whether to adjust the size of the doublets using the median sizes per cluster of the originating cells. Requires 'clusters' to be given. Alternatively to a logical value, a number between 0 and 1 can be given, determining the proportion of the doublets for which to perform the size adjustment.
prefix	Prefix for the colnames generated.

Value

A matrix of artificial doublets.

```
sce <- mockDoubletSCE()
idx <- getCellPairs(sce$cluster, n=200)
art.dbls <- createDoublets(sce, idx)</pre>
```

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|--|

Description

Calculates a coexpression-based doublet score using the method developed by Bais and Kostka 2020. This is the original implementation from the scds package, but enabling scores to be calculated for all cells while the gene coexpression is based only on a subset (i.e. excluding known/artificial doublets).

Usage

```
cxds2(x, whichDbls = c(), ntop = 500, binThresh = 0)
```

Arguments

x A matrix of counts, or a 'SingleCellExperiment' containing a 'counts'

whichDbls The columns of 'x' which are known doublets.

ntop The number of top features to keep.

binThresh The count threshold to be considered expressed.

Value

A cxds score or, if 'x' is a 'SingleCellExperiment', 'x' with an added 'cxds_score' colData column.

References

```
https://doi.org/10.1093/bioinformatics/btz698
```

Examples

```
sce <- mockDoubletSCE()
sce <- cxds2(sce)</pre>
```

data

Comparison results

Description

Results of the comparison of doublet detection methods. Each element of the list is a dataset, itself a list with scores for the different methods as well as the demuxlet SNP-based doublet calls ('demuxlet_cls').

Value

a list.

doubletPairwiseEnrichment

doubletPairwiseEnrichment

Description

Calculates enrichment in any type of doublet (i.e. specific combination of clusters) over random expectation. Note that when applied to an multisample object, this functions assumes that the cluster labels match across samples.

Usage

```
doubletPairwiseEnrichment(
    x,
    lower.tail = FALSE,
    sampleWise = FALSE,
    type = c("poisson", "binomial", "chisq", "nbinom1")
)
```

Arguments

Х	A table of double statistics, or a SingleCellExperiment on which scDblFinder was run.
lower.tail	Logical; defaults to FALSE to test enrichment (instead of depletion).
sampleWise	Logical; whether to perform tests sample-wise in multi-sample datasets. If FALSE (default), will aggregate counts before testing.
type	Type of test to use.

Value

A table of significances for each combination.

```
sce <- mockDoubletSCE()
sce <- scDblFinder(sce, artificialDoublets=500)
doubletPairwiseEnrichment(sce)</pre>
```

doubletThresholding 11

doubletThresholding doubletThresholding

Description

Sets the doublet scores threshold; typically called by scDblFinder.

Usage

```
doubletThresholding(
    d,
    dbr = NULL,
    dbr.sd = 0.015,
    stringency = 0.5,
    p = 0.1,
    method = c("auto", "optim", "dbr", "griffiths"),
    perSample = TRUE,
    returnType = c("threshold", "call")
)
```

Arguments

d A data frame of cell properties, with each row representing a cell, as produced

by 'scDblFinder(..., returnType="table")', or minimally containing a 'score' col-

umn.

dbr The expected (mean) doublet rate. If 'd' contains a 'cluster' column, the doublet

rate will be adjusted for homotypic doublets.

dbr.sd The standard deviation of the doublet rate, representing the uncertainty in the

estimate. Ignored if 'method!="optim"'.

stringency A numeric value >0 and <1 which controls the relative weight of false positives

(i.e. real cells) and false negatives (artificial doublets) in setting the threshold. A value of 0.5 gives equal weight to both; a higher value (e.g. 0.7) gives higher weight to the false positives, and a lower to artificial doublets. Ignored if

'method!="optim"'.

p The p-value threshold determining the deviation in doublet score.

method The thresholding method to use, either 'auto' (default, automatic selection de-

pending on the available fields), 'optim' (optimization of misclassification rate and deviation from expected doublet rate), 'dbr' (strictly based on the expected doublet rate), or 'griffiths' (cluster-wise number of median absolute deviation in

doublet score).

perSample Logical; whether to perform thresholding individually for each sample.

returnType The type of value to return, either doublet calls ('call') or thresholds ('thresh-

old').

verbose Logical; output extra information.

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Value

A vector of doublet calls if 'returnType=="call"', or a threshold (or vector of thresholds) if 'return-Type=="threshold"'.

Examples

```
sce <- mockDoubletSCE()
d <- scDblFinder(sce, verbose=FALSE, returnType="table")
th <- doubletThresholding(d, dbr=0.05)
th</pre>
```

fastcluster

fastcluster

Description

Performs a fast two-step clustering: first clusters using k-means with a very large k, then uses louvain clustering of the k cluster averages and reports back the cluster labels.

Usage

```
fastcluster(
    X,
    k = NULL,
    rdname = "PCA",
    nstart = 3,
    iter.max = 20,
    ndims = NULL,
    nfeatures = 1000,
    verbose = TRUE,
    returnType = c("clusters", "preclusters", "metacells", "graph"),
    ...
)
```

Arguments

X	An object of class SCE
k	The number of k-means clusters to use in the primary step (should be much higher than the number of expected clusters). Defaults to 1/10th of the number of cells with a maximum of 3000.
rdname	The name of the dimensionality reduction to use.
nstart	Number of starts for k-means clustering
iter.max	Number of iterations for k-means clustering
ndims	Number of dimensions to use

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Number of features to use (ignored if 'rdname' is given and the corresponding dimensional reduction exists in 'sce')

verbose Logical; whether to output progress messages

returnType See return.

... Arguments passed to 'scater::runPCA' (e.g. BPPARAM or BSPARAM) if 'x' does not have 'rdname'.

Value

By default, a vector of cluster labels. If 'returnType='preclusters', returns the k-means pre-clusters. If 'returnType='metacells', returns the metacells aggretated by pre-clusters and the corresponding cell indexes. If 'returnType='graph', returns the graph of (meta-)cells and the corresponding cell indexes.

Examples

```
sce <- mockDoubletSCE()
sce$cluster <- fastcluster(sce)</pre>
```

Description

Identify potential clusters of doublet cells based on whether they have intermediate expression profiles, i.e., their profiles lie between two other "source" clusters.

Usage

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Arguments

A numeric matrix-like object of count values, where each column corresponds х to a cell and each row corresponds to an endogenous gene. Alternatively, a SummarizedExperiment or SingleCellExperiment object containing such a matrix. For the generic, additional arguments to pass to specific methods. For the ANY method, additional arguments to pass to findMarkers. For the SummarizedExperiment method, additional arguments to pass to the ANY method. For the SingleCellExperiment method, additional arguments to pass to the SummarizedExperiment method. A vector of length equal to ncol(x), containing cluster identities for all cells. If clusters x is a SingleCellExperiment, this is taken from colLabels(x) by default. See ?"scran-gene-selection". subset.row threshold A numeric scalar specifying the FDR threshold with which to identify significant genes. get.all.pairs Logical scalar indicating whether statistics for all possible source pairings should be returned. A string specifying which assay values to use, e.g., "counts" or "logcounts". assay.type

Details

This function detects clusters of doublet cells in a manner similar to the method used by Bach et al. (2017). For each "query" cluster, we examine all possible pairs of "source" clusters, hypothesizing that the query consists of doublets formed from the two sources. If so, gene expression in the query cluster should be strictly intermediate between the two sources after library size normalization.

We apply pairwise t-tests to the normalized log-expression profiles to reject this null hypothesis. This is done by identifying genes that are consistently up- or down-regulated in the query compared to *both* sources. We count the number of genes that reject the null hypothesis at the specified FDR threshold. For each query cluster, the most likely pair of source clusters is that which minimizes the number of significant genes.

Potential doublet clusters are identified using the following characteristics, in order of importance:

- Low number of significant genes (i.e., num.de). Ideally, median.de is also high to indicate that the absence of strong DE is not due to a lack of power.
- A reasonable proportion of cells in the cluster, i.e., prop. This requires some expectation of the doublet rate in the experimental protocol.
- Library sizes of the source clusters that are below that of the query cluster, i.e., lib.size* values below unity. This assumes that the doublet cluster will contain more RNA and have more counts than either of the two source clusters.

For each query cluster, the function will only report the pair of source clusters with the lowest num.de. Setting get.all.pairs=TRUE will retrieve statistics for all pairs of potential source clusters. This can be helpful for diagnostics to identify relationships between specific clusters.

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The reported p.value is of little use in a statistical sense, and is only provided for inspection. Technically, it could be treated as the Simes combined p-value against the doublet hypothesis for the query cluster. However, this does not account for the multiple testing across all pairs of clusters for each chosen cluster, especially as we are chosing the pair that is most concordant with the doublet null hypothesis.

We use library size normalization (via librarySizeFactors) even if existing size factors are present. This is because intermediate expression of the doublet cluster is not guaranteed for arbitrary size factors. For example, expression in the doublet cluster will be higher than that in the source clusters if normalization was performed with spike-in size factors.

Value

A DataFrame containing one row per query cluster with the following fields:

source1: String specifying the identity of the first source cluster.

source2: String specifying the identity of the second source cluster.

num.de: Integer, number of genes that are significantly non-intermediate in the query cluster compared to the two putative source clusters.

median.de: Integer, median number of genes that are significantly non-intermediate in the query cluster across all possible source cluster pairings.

best: String specifying the identify of the top gene with the lowest p-value against the doublet hypothesis for this combination of query and source clusters.

p.value: Numeric, containing the adjusted p-value for the best gene.

lib.size1: Numeric, ratio of the median library sizes for the first source cluster to the query cluster.

lib.size2: Numeric, ratio of the median library sizes for the second source cluster to the query cluster.

prop: Numeric, proportion of cells in the query cluster.

all.pairs: A SimpleList object containing the above statistics for every pair of potential source clusters, if get.all.pairs=TRUE.

Each row is named according to its query cluster.

Author(s)

Aaron Lun

References

Bach K, Pensa S, Grzelak M, Hadfield J, Adams DJ, Marioni JC and Khaled WT (2017). Differentiation dynamics of mammary epithelial cells revealed by single-cell RNA sequencing. *Nat Commun.* 8, 1:2128.

See Also

findMarkers, to detect DE genes between clusters.

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Examples

```
# Mocking up an example.
library(SingleCellExperiment)
sce <- mockDoubletSCE()</pre>
# Compute doublet-ness of each cluster:
dbl <- findDoubletClusters(counts(sce), sce$cluster)</pre>
dbl
# Narrow this down to clusters with very low 'N':
library(scuttle)
isOutlier(dbl$num.de, log=TRUE, type="lower")
# Get help from "lib.size" below 1.
dbl$lib.size1 < 1 & dbl$lib.size2 < 1
```

 ${\tt getArtificialDoublets} \quad {\tt getArtificialDoublets}$

Description

Create expression profiles of random artificial doublets.

Usage

```
getArtificialDoublets(
  х,
  n = 3000,
  clusters = NULL,
  resamp = 0.25,
  halfSize = 0.25,
  adjustSize = 0.25,
  propRandom = 0.1,
  selMode = c("proportional", "uniform", "sqrt"),
  n.meta.cells = 2,
  meta.triplets = TRUE,
  trim.q = c(0.05, 0.95)
)
```

Arguments

A count matrix, with features as rows and cells as columns. Х

The approximate number of doublet to generate (default 3000).

The optional clusters labels to use to build cross-cluster doublets. clusters

Logical; whether to resample the doublets using the poisson distribution. Alterresamp

natively, if a proportion between 0 and 1, the proportion of doublets to resample.

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halfSize	Logical; whether to half the library size of doublets (instead of just summing up the cells). Alternatively, a number between 0 and 1 can be given, determining the proportion of the doublets for which to perform the size adjustment.
adjustSize	Logical; whether to adjust the size of the doublets using the ratio between each cluster's median library size. Alternatively, a number between 0 and 1 can be given, determining the proportion of the doublets for which to perform the size adjustment.
propRandom	The proportion of the created doublets that are fully random (default 0.1); the rest will be doublets created across clusters. Ignored if 'clusters' is NULL.
selMode	The cell pair selection mode for inter-cluster doublet generation, either 'uniform' (same number of doublets for each combination), 'proportional' (proportion expected from the clusters' prevalences), or 'sqrt' (roughly the square root of the expected proportion).
n.meta.cells	The number of meta-cell per cluster to create. If given, additional doublets will be created from cluster meta-cells. Ignored if 'clusters' is missing.
meta.triplets	Logical; whether to create triplets from meta cells. Ignored if 'clusters' is missing.
trim.q	A vector of two values between 0 and 1

Value

A list with two elements: 'counts' (the count matrix of the artificial doublets) and 'origins' the clusters from which each artificial doublets originated (NULL if 'clusters' is not given).

Examples

getCellPairs getCellPairs

Description

Given a vector of cluster labels, returns pairs of cross-cluster cells

Usage

```
getCellPairs(x, n = 1000, ...)
```

Arguments

X	A vector of cluster labels for each cell, or a list containing metacells and graph
n	The number of cell pairs to obtain
	Further arguments, for instance the 'k' vector of precluster labels if 'x' is a metacell graph.

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Value

A data frame with the columns

Examples

```
# create random labels
x <- sample(head(LETTERS), 100, replace=TRUE)
getCellPairs(x, n=6)</pre>
```

getExpectedDoublets getExpectedDoublets

Description

getExpectedDoublets

Usage

```
getExpectedDoublets(x, dbr = NULL, only.heterotypic = TRUE)
```

Arguments

x A vector of cluster labels for each cell

dbr The expected doublet rate.

only.heterotypic

Logical; whether to return expectations only for heterotypic doublets

Value

The expected number of doublets of each combination of clusters

```
# random cluster labels
cl <- sample(head(LETTERS,4), size=2000, prob=c(.4,.2,.2), replace=TRUE)
getExpectedDoublets(cl)</pre>
```

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mockDoubletSCE

mockDoubletSCE

Description

Creates a mock random single-cell experiment object with doublets

Usage

```
mockDoubletSCE(
  ncells = c(200, 300),
  ngenes = 200,
  mus = NULL,
  dbl.rate = 0.1,
  only.heterotypic = TRUE
)
```

Arguments

ncells A positive integer vector indicating the number of cells per cluster (min 2 clus-

ters)

ngenes The number of genes to simulate. Ignored if 'mus' is given.

mus A list of cluster averages.

dbl.rate The doublet rate

only.heterotypic

Whether to create only heterotypic doublets

Value

A SingleCellExperiment object, with the colData columns 'type' indicating whether the cell is a singlet or doublet, and 'cluster' indicating from which cluster (or cluster combination) it was simulated.

```
sce <- mockDoubletSCE()</pre>
```

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plotDoubletMap	plotDoubletMap
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Description

Plots a heatmap of observed versus expected doublets. Requires the 'ComplexHeatmap' package.

Usage

```
plotDoubletMap(
    sce,
    colorBy = "enrichment",
    labelBy = "observed",
    addSizes = TRUE,
    col = NULL,
    column_title = "Clusters",
    row_title = "Clusters",
    column_title_side = "bottom",
    na_col = "white",
    ...
)
```

Arguments

sce	A SingleCellExperiment object on which 'scDblFinder' has been run.	
colorBy	Determines the color mapping. Either "enrichment" (for log2-enrichment over expectation) or any column of 'metadata(sce)\$scDblFinder.stats'	
labelBy	Determines the cell labels. Either "enrichment" (for log2-enrichment over expectation) or any column of 'metadata(sce)\$scDblFinder.stats'	
addSizes	Logical; whether to add the sizes of clusters to labels	
col	The colors scale to use (passed to 'ComplexHeatmap::Heatmap')	
column_title	passed to 'ComplexHeatmap::Heatmap'	
row_title	passed to 'ComplexHeatmap::Heatmap'	
column_title_side		
	passed to 'ComplexHeatmap::Heatmap'	
na_col	color for NA cells	
	passed to 'ComplexHeatmap::Heatmap'	

Value

```
a Heatmap object
```

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 ${\tt plotThresholds}$

plotThresholds

Description

Plots scores used for thresholding.

Usage

```
plotThresholds(
    d,
    ths = (0:100)/100,
    dbr = NULL,
    dbr.sd = 0.015,
    do.plot = TRUE
)
```

Arguments

d	A data.frame of cell properties, with each row representing a cell, as produced by 'scDblFinder(, returnType="table")'.
ths	A vector of thresholds between 0 and 1 at which to plot values.
dbr	The expected (mean) doublet rate.
dbr.sd	The standard deviation of the doublet rate, representing the uncertainty in the estimate.
do.plot	Logical; whether to plot the data.

Value

A ggplot, or a data.frame if 'do.plot==FALSE'.

propHomotypic propHomotypic

Description

Computes the proportion of pairs expected to be made of elements from the same cluster.

Usage

```
propHomotypic(clusters)
```

Arguments

clusters

A vector of cluster labels

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Value

A numeric value between 0 and 1.

Examples

```
clusters <- sample(LETTERS[1:5], 100, replace=TRUE)
propHomotypic(clusters)</pre>
```

recoverDoublets

Recover intra-sample doublets

Description

Recover intra-sample doublets that are neighbors to known inter-sample doublets in a multiplexed experiment.

Usage

```
recoverDoublets(x, ...)
## S4 method for signature 'ANY'
recoverDoublets(
  х,
 doublets,
  samples,
 k = 50,
  transposed = FALSE,
  subset.row = NULL,
 BNPARAM = KmknnParam(),
 BPPARAM = SerialParam()
)
## S4 method for signature 'SummarizedExperiment'
recoverDoublets(x, ..., assay.type = "logcounts")
## S4 method for signature 'SingleCellExperiment'
recoverDoublets(x, ..., use.dimred = NULL)
```

Arguments

Χ

A log-expression matrix for all cells (including doublets) in columns and genes in rows. If transposed=TRUE, this should be a matrix of low-dimensional coordinates where each row corresponds to a cell.

Alternatively, a SummarizedExperiment or SingleCellExperiment containing (i) a log-expression matrix in the assays as specified by assay.type, or (ii) a matrix of reduced dimensions in the reducedDims as specified by use.dimred.

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	For the generic, additional arguments to pass to specific methods.
	For the SummarizedExperiment method, additional arguments to pass to the ANY method.
	For the SingleCellExperiment method, additional arguments to pass to the SummarizedExperiment method.
doublets	A logical, integer or character vector specifying which cells in x are known (inter-sample) doublets.
samples	A numeric vector containing the relative proportions of cells from each sample, used to determine how many cells are to be considered as intra-sample doublets.
k	Integer scalar specifying the number of nearest neighbors to use for computing the local doublet proportions.
transposed	Logical scalar indicating whether x is transposed, i.e., cells in the rows.
subset.row	A logical, integer or character vector specifying the genes to use for the neighbor search. Only used when transposed=FALSE.
BNPARAM	A BiocNeighborParam object specifying the algorithm to use for the nearest neighbor search.
BPPARAM	A BiocParallelParam object specifying the parallelization to use for the nearest neighbor search.
assay.type	A string specifying which assay values contain the log-expression matrix.
use.dimred	A string specifying whether existing values in reducedDims(x) should be used.

Details

In multiplexed single-cell experiments, we can detect doublets as libraries with labels for multiple samples. However, this approach fails to identify doublets consisting of two cells with the same label. Such cells may be problematic if they are still sufficiently abundant to drive formation of spurious clusters.

This function identifies intra-sample doublets based on the similarity in expression profiles to known inter-sample doublets. For each cell, we compute the proportion of the k neighbors that are known doublets. Of the "unmarked" cells that are not known doublets, those with top X largest proportions are considered to be intra-sample doublets. We use samples to obtain a reasonable estimate for X, see the vignette for details.

A larger value of k provides more stable estimates of the doublet proportion in each cell. However, this comes at the cost of assuming that each cell actually has k neighboring cells of the same state. For example, if a doublet cluster has fewer than k members, its doublet proportions will be "diluted" by inclusion of unmarked cells in the next-closest cluster.

Value

A DataFrame containing one row per cell and the following fields:

- proportion, a numeric field containing the proportion of neighbors that are doublets.
- known, a logical field indicating whether this cell is a known inter-sample doublet.
- predicted, a logical field indicating whether this cell is a predicted intra-sample doublet.

The metadata contains intra, a numeric scalar containing the expected number of intra-sample doublets.

Author(s)

Aaron Lun

See Also

doubletCells and doubletCluster, for alternative methods of doublet detection when no prior doublet information is available.

hashedDrops from the DropletUtils package, to identify doublets from cell hashing experiments.

More detail on the mathematical background of this function is provided in the corresponding vignette at vignette("recoverDoublets",package="scDblFinder").

Examples

```
# Mocking up an example.
set.seed(100)
ngenes <- 1000
mu1 <- 2^rnorm(ngenes, sd=2)
mu2 <- 2^rnorm(ngenes, sd=2)

counts.1 <- matrix(rpois(ngenes*100, mu1), nrow=ngenes) # Pure type 1
counts.2 <- matrix(rpois(ngenes*100, mu2), nrow=ngenes) # Pure type 2
counts.m <- matrix(rpois(ngenes*20, mu1+mu2), nrow=ngenes) # Doublets (1 & 2)
all.counts <- cbind(counts.1, counts.2, counts.m)
lcounts <- scuttle::normalizeCounts(all.counts)

# Pretending that half of the doublets are known. Also pretending that
# the experiment involved two samples of equal size.
known <- 200 + seq_len(10)
out <- recoverDoublets(lcounts, doublets=known, k=10, samples=c(1, 1))
out</pre>
```

scDblFinder

scDblFinder

Description

Identification of heterotypic (or neotypic) doublets in single-cell RNAseq using cluster-based generation of artificial doublets.

Usage

```
scDblFinder(
   sce,
   clusters = NULL,
   samples = NULL,
   trajectoryMode = FALSE,
   artificialDoublets = NULL,
```

```
knownDoublets = NULL,
  dbr = NULL,
  clustCor = NULL,
  dbr.sd = NULL,
  nfeatures = 1000,
  dims = 20,
  k = NULL
  removeUnidentifiable = TRUE,
  includePCs = 1:5,
  propRandom = 0,
  propMarkers = 0,
  aggregateFeatures = FALSE,
  returnType = c("sce", "table", "full"),
  score = c("xgb", "weighted", "ratio"),
  processing = "default",
 metric = "logloss",
  nrounds = 0.25,
 max_depth = 5,
  iter = 1,
 multiSampleMode = c("split", "singleModel", "singleModelSplitThres"),
  threshold = TRUE,
  verbose = is.null(samples),
 BPPARAM = SerialParam(),
)
```

Arguments

sce

 $\label{lem:class} A \ {\tt SummarizedExperiment-class}, \ {\tt SingleCellExperiment-class}, \ {\tt or} \ {\tt array} \ {\tt of} \ {\tt counts}.$

clusters

The optional cluster assignments (if omitted, will run clustering). This is used to make doublets more efficiently. clusters should either be a vector of labels for each cell, or the name of a colData column of sce. Alternatively, if it is a single integer, will determine how many clusters to create (using k-means clustering). This options should be used when distinct subpopulations are not expected in the data (e.g. trajectories).

samples

A vector of the same length as cells (or the name of a column of colData(x)), indicating to which sample each cell belongs. Here, a sample is understood as being processed independently. If omitted, doublets will be searched for with all cells together. If given, doublets will be searched for independently for each sample, which is preferable if they represent different captures. If your samples were multiplexed using cell hashes, want you want to give here are the different batches/wells (i.e. independent captures, since doublets cannot arise across them) rather than biological samples.

trajectoryMode

Logical; whether to generate fewer doublets from cells that are closer to each other, for datasets with gradients rather than separated subpopulations. This disrupts the proportionality and is not anymore the recommended way of handling such datasets. See vignette("scDblFinder") for more details.

artificialDoublets

The approximate number of artificial doublets to create. If NULL, will be the maximum of the number of cells or 5*nbClusters^2.

knownDoublets

An optional logical vector of known doublets (e.g. through cell barcodes), or the name of a colData column of 'sce' containing that information. Including known doublets tends to increase the sensitivity of doublet identification, but decrease the specificity (since some of the known doublets are homotypic).

dbr

The expected doublet rate. By default this is assumed to be 1% per thousand cells captured (so 4% among 4000 thousand cells), which is appropriate for 10x datasets. Corrections for homeotypic doublets will be performed on the given rate.

clustCor

Include Spearman correlations to cell type averages in the predictors. If 'clust-Cor' is a matrix of cell type marker expressions (with features as rows and cell types as columns), the subset of these which are present in the selected features will be correlated to each cell to produce additional predictors (i.e. one per cell type). Alternatively, if 'clustCor' is a positive integer, this number of inter-cluster markers will be selected and used for correlation (se 'clustCor=Inf' to use all available genes).

dbr.sd

The uncertainty range in the doublet rate, interpreted as a +/- around 'dbr'. During thresholding, deviation from the expected doublet rate will be calculated from these boundaries, and will be considered null within these boundaries. If NULL, will be 40% of 'dbr'. Set to 'dbr.sd=0' to disable.

nfeatures

The number of top features to use (default 1000)

dims

The number of dimensions used.

k

Number of nearest neighbors (for KNN graph). If more than one value is given, the doublet density will be calculated at each k (and other values at the highest k), and all the information will be used by the classifier. If omitted, a reasonable set of values is used.

removeUnidentifiable

Logical; whether to remove artificial doublets of a combination that is generally found to be unidentifiable.

includePCs

The index of principal components to include in the predictors (e.g. 'include-PCs=1:2').

propRandom

The proportion of the artificial doublets which should be made of random cells (as opposed to inter-cluster combinations).

propMarkers

The proportion of features to select based on marker identification.

aggregateFeatures

Whether to perform feature aggregation (recommended for ATAC). Can also be a positive integer, in which case this will indicate the number of components to use for feature aggregation (if TRUE, 'dims' will be used.)

returnType

Either "sce" (default), "table" (to return the table of cell attributes including artificial doublets), or "full" (returns an SCE object containing both the real and artificial cells.

score

Score to use for final classification.

processing Counts (real and artificial) processing before KNN. Either 'default' (normal

scater-based normalization and PCA), "rawPCA" (PCA without normalization), "rawFeatures" (no normalization/dimensional reduction) or a custom function with (at least) arguments 'e' (the matrix of counts) and 'dims' (the desired number of dimensions), returning a named matrix with cells as rows and com-

ponents as columns.

metric Error metric to optimize during training (e.g. 'merror', 'logloss', 'auc', 'aucpr').

nrounds Maximum rounds of boosting. If NULL, will be determined through cross-

validation.

max_depth Maximum depths of each tree.

iter A positive integer indicating the number of scoring iterations (ignored if 'score'

isn't based on classifiers). At each iteration, real cells that would be called as doublets are excluding from the training, and new scores are calculated. Rec-

ommended values are 1 or 2.

multiSampleMode

Either "split" (recommended if there is a lot of heterogeneity across samples), "singleModel" (recommended _only_ if the samples are very similar), or "singleModelSplitThres" (use a single classifier, but sample-specific thresholds).

threshold Logical; whether to threshold scores into binary doublet calls verbose Logical; whether to print messages and the thresholding plot.

BPPARAM Used for multithreading when splitting by samples (i.e. when 'samples!=NULL');

otherwise passed to eventual PCA and K/SNN calculations.

... further arguments passed to getArtificialDoublets.

Details

This function generates artificial doublets from clusters of real cells, evaluates their prevalence in the neighborhood of each cells, and uses this along with additional features to classify doublets. The approach is complementary to doublets identified via cell hashes and SNPs in multiplexed samples: the latter can identify doublets formed by cells of the same type from two samples, which are nearly undistinguishable from real cells transcriptionally, but cannot identify doublets made by cells of the same sample. See vignette("scDblFinder") for more details on the method.

When multiple samples/captures are present, they should be specified using the samples argument. In this case, we recommend the use of BPPARAM to perform several of the steps in parallel. Artificial doublets and kNN networks will be computed separately; then the behavior will then depend on the 'multiSampleMode' argument. If 'split', the whole process is split by sample (this is recommended when there is heterogeneity between samples, for instance in the number of cells); if 'singleModel', the classifier and thresholding will be trained globally (this is not recommended unless the samples are extremely comparable); if 'singleModelSplitThres', the classifierwill be trained globally, but the thresholding be performed separately for each samples.

When inter-sample doublets are available, they can be provided to 'scDblFinder' through the knownDoublets argument to improve the identification of further doublets. However, because such 'true' doublets can include a lot of homotypic doublets, in practice this often lead to a slight decrease in the accuracy of detecting neotypic doublets.

Finally, for some types of data, such as single-cell ATAC-seq, selecting a number of top features is ineffective due to the high sparsity of the signal. In such contexts, rather than _selecting_ features

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we recommend to use the alternative approach of _aggregating_ similar features (with 'aggregate-Features=TRUE'), which strongly improves accuracy.

Value

The sce object with several additional colData columns, in particular 'scDblFinder.score' (the final score used) and 'scDblFinder.class' (whether the cell is called as 'doublet' or 'singlet'). See vignette("scDblFinder") for more details; for alternative return values, see the 'returnType' argument.

Examples

```
library(SingleCellExperiment)
sce <- mockDoubletSCE()
sce <- scDblFinder(sce, dbr=0.1)
table(truth=sce$type, call=sce$scDblFinder.class)</pre>
```

selFeatures

selFeatures

Description

Selects features based on cluster-wise expression or marker detection, or a combination.

Usage

```
selFeatures(
   sce,
   clusters = NULL,
   nfeatures = 1000,
   propMarkers = 0,
   FDR.max = 0.05
)
```

Arguments

A SummarizedExperiment-class, SingleCellExperiment-class with a 'counts' assay.

Clusters Optional cluster assignments. Should either be a vector of labels for each cell.

The number of features to select.

propMarkers The proportion of features to select from markers (rather than on the basis of high expression). Ignored if 'clusters' isn't given.

FDR.max The maximum marker binom FDR to be included in the selection. (see findMarkers).

Value

A vector of feature (i.e. row) names.

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Examples

```
sce <- mockDoubletSCE()
selFeatures(sce, clusters=sce$cluster, nfeatures=5)</pre>
```

TFIDF

TFIDF

Description

The Term Frequency - Inverse Document Frequency (TF-IDF) normalization, as implemented in Stuart & Butler et al. 2019.

Usage

```
TFIDF(x, sf = 10000)
```

Arguments

x The matrix of occurrences

sf Scaling factor

Value

An array of same dimensions as 'x'

```
m <- matrix(rpois(500,1),nrow=50)
m <- TFIDF(m)</pre>
```

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