## Package 'scCB2'

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**Title** CB2 improves power of cell detection in droplet-based single-cell RNA sequencing data

**Depends** R (>= 3.6.0)

**Imports** SingleCellExperiment, SummarizedExperiment, Matrix, methods, utils, stats, edgeR, rhdf5, parallel, DropletUtils, doParallel, iterators, foreach, Seurat

**Suggests** testthat (>= 2.1.0), KernSmooth, beachmat, knitr, BiocStyle, rmarkdown

**biocViews** DataImport, RNASeq, SingleCell, Sequencing, GeneExpression, Transcriptomics, Preprocessing, Clustering

Description scCB2 is an R package implementing CB2 for distinguish-

ing real cells from empty droplets in droplet-based single cell RNA-seq experiments (especially for 10x Chromium).

It is based on clustering similar barcodes and calculating Monte-Carlo p-value for each cluster to test against background distribution.

This cluster-level test outperforms single-barcode-level tests in dealing with low count barcodes and homogeneous sequencing library, while keeping FDR well controlled.

License GPL-3

**NeedsCompilation** yes **VignetteBuilder** knitr

**Encoding** UTF-8

SystemRequirements C++11

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BugReports https://github.com/zijianni/scCB2/issues

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2 CB2FindCell

```
Author Zijian Ni [aut, cre],
      Shuyang Chen [ctb],
      Christina Kendziorski [ctb]
Maintainer Zijian Ni <zni25@wisc.edu>
```

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## **Description**

The main function of scCB2 package. Distinguish real cells from empty droplets using clusteringbased Monte-Carlo test.

## Usage

```
CB2FindCell(
  RawDat,
  FDR_threshold = 0.01,
  lower = 100,
  upper = NULL,
  GeneExpressionOnly = TRUE,
  Ncores = 2,
  TopNGene = 30000,
  verbose = TRUE
)
```

## **Arguments**

RawDat Matrix. Supports standard matrix or sparse matrix. This is the raw feature-by-

barcode count matrix.

Numeric between 0 and 1. Default: 0.01. The False Discovery Rate (FDR) to FDR\_threshold

be controlled for multiple testing.

lower Positive integer. Default: 100. All barcodes whose total count below or equal

to this threshold are defined as background empty droplets. They will be used to estimate the background distribution. The remaining barcodes will be test against background distribution. If sequencing depth is deliberately made higher (lower) than usual, this threshold can be leveled up (down) correspondingly to get reasonable number of cells. Recommended sequencing depth for this default

threshold: 40,000~80,000 reads per cell.

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upper

Positive integer. Default: NULL. This is the upper threshold for large barcodes. All barcodes whose total counts are larger or equal to upper threshold are directly classified as real cells prior to testing. If upper = NULL, the knee point of the log rank curve of barcodes total counts will serve as the upper threshold, which is calculated using package DropletUtils's method. If upper = Inf, no barcodes will be retained prior to testing. If manually specified, it should be greater than pooling threshold.

#### GeneExpressionOnly

Logical. Default: TRUE. For 10x Cell Ranger version >=3, extra features (surface proteins, cell multiplexing oligos, etc) besides genes are measured simultaneously. If GeneExpressionOnly = TRUE, only genes are used for testing. Removing extra features are recommended because the default pooling threshold (100) is chosen only for handling gene expression. Extra features expression level is hugely different from gene expression level. If using the default pooling threshold while keeping extra features, the estimated background distribution will be hugely biased and does not reflect the real background distribution of empty droplets.

Ncores Positive integer. Default: 2. Number of cores for parallel computation.

TopNGene Positive integer. Default: 30000. Number of top highly expressed genes to use.

This threshold avoids high number of false positives in ultra-high dimensional

datasets, e.g. 10x barnyard data.

verbose Logical. Default: TRUE. If verbose = TRUE, progressing messages will be printed.

#### **Details**

Input data is a feature-by-barcode matrix. Background barcodes are defined based on lower. Large barcodes are automatically treated as real cells based on upper. Remaining barcodes will be first clustered into subgroups, then tested against background using Monte-Carlo p-values simulated from Multinomial distribution. The rest barcodes will be further tested using EmptyDrops (Aaron T. L. Lun *et. al.* 2019). FDR is controlled based on FDR\_threshold.

This function supports parallel computation. Ncores is used to specify number of cores.

Under CellRanger version >=3, extra features other than genes are simultaneously measured (e.g. surface protein, cell multiplexing oligo). We recommend filtering them out using GeneExpressionOnly = TRUE because the expression of extra features is not in the same scale as gene expression counts. If using the default pooling threshold while keeping extra features, the estimated background distribution will be hugely biased and does not reflect the real background distribution of empty droplets. The resulting matrix will contain lots of barcodes who have almost zero gene expression and relatively high extra features expression, which are usually not useful for RNA-Seq study.

## Value

An object of class SummarizedExperiment. The slot assays contains the real cell barcode matrix distinguished during cluster-level test, single-barcode-level test plus large cells who exceed the upper threshold. The slot metadata contains (1) testing statistics (Pearson correlation to the background) for all candidate barcode clusters, (2) barcode IDs for all candidate barcode clusters, the name of each cluster is its median barcode size, (3) testing statistics (log likelihood under background distribution) for remaining single barcodes not clustered, (4) background distribution count vector without Good-Turing correction.

#### **Examples**

# raw data, all barcodes

```
data(mbrainSub)
str(mbrainSub)

# run CB2 on the first 10000 barcodes
CBOut <- CB2FindCell(mbrainSub[,1:10000], FDR_threshold = 0.01,
    lower = 100, Ncores = 2)
RealCell <- GetCellMat(CBOut, MTfilter = 0.05)

# real cells
str(RealCell)</pre>
```

 ${\it Check Background Cutoffs\ and\ recommend\ an\ appropriate}$  one

## Description

The key parameter of CB2 as well as other similar methods is the background cutoff, which divides barcodes into two groups: (1) small barcodes that are most likely to be background; (2) the rest barcodes that can be either background or cell, and remain to be tested. Those small barcodes will be used to estimate a background distribution, which guides the identification of cells from background. It is crucial to have an unbiased estimation of the background distribution.

### Usage

CheckBackgroundCutoff(RawDat)

#### **Arguments**

RawDat

Matrix. Supports standard matrix or sparse matrix. This is the raw feature-by-barcode count matrix.

#### **Details**

An appropriate background cutoff should be reasonably large to contain enough background information, but shouldn't be too large to mistakenly include real cells. We recommend a background cutoff which (1) puts more than 90 (2) puts more than 10 The smallest cutoff satisfying either condition is the recommended cutoff.

## Value

A list containing a data frame summarizing background information under different background cutoffs, and the recommended background cutoff for the input data. For the data frame, 'n\_bg\_bcs' is the number of barcodes less or equal to the cutoff, 'n\_bg\_counts' is the number of UMI counts within the barcodes less or equal to the cutoff, 'prop\_bg\_bcs' and 'prop\_bg\_counts' are the corresponding proportions.

#### **Examples**

```
data(mbrainSub)
CheckBackgroundCutoff(mbrainSub)
```

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FilterGB	Filter out low count genes and barcodes from count matrix	

#### **Description**

This function is used for filtering out low count genes and barcodes from count matrix based on total gene expression count (row sums) and barcode expression count (column sums). CB2FindCell has already integrated this function into it with g\_threshold = 0 and b\_threshold = 0. If users plan to customize their filtering threshold, this function can be applied to the raw expression count matrix prior to running CB2FindCell.

## Usage

```
FilterGB(dat, g_threshold = 0, b_threshold = 0)
```

## **Arguments**

dat Input count matrix to be filtered.

g\_threshold Nonnegative integer. Default: 0. Filtering threshold for genes. Any gene whose

total expression count is less or equal to g\_threshold will be filtered out.

b\_threshold Nonnegative integer. Default: 0. Filtering threshold for barcodes. Any barcode

whose total count is less or equal to b\_threshold will be filtered out.

## Value

A filtered matrix with the same format as input matrix.

#### **Examples**

```
data(mbrainSub)
dim(mbrainSub)
mbrainSub_f <- FilterGB(mbrainSub)
dim(mbrainSub_f)</pre>
```

GetCellMat Extract real cell matrix from CB2FindCell output and optionally filter out low-quality cells

## **Description**

Handy function to extract real cell matrix from CB2FindCell output. It provides the option to filter out broken cells based on proportion of mitochondrial gene expressions. The input can also be a sparse matrix only for cell filtering.

#### Usage

```
GetCellMat(CBout, MTfilter = 0.25, MTgene = NULL)
```

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## **Arguments**

CBout Output object from CB2FindCell, or a sparse matrix (for example, from QuickCB2).

MTfilter Numeric value between 0 and 1. Default: 0.25. For each barcode, if the pro-

portion of mitochondrial gene expression exceeds MTfilter, this barcode will be filtered out. By default, cell barcodes with more than 25 filtered out. Set

MTfilter = 1 for no filtering.

The proportion of mitochondrial gene expressions is usually a criterion for evaluating cell quality, and is calculated using the scaled sum of all genes starting with "MT-" (human) or "mt-" (mouse) if row names are gene symbols, or cus-

tomized mitochondrial genes specified by MTgene.

MTgene Character vector. User may specify customized mitochondrial gene names to

perform the filtering. This should correspond to a subset of row names in raw

data.

#### Value

A dgCMatrix count matrix of real cells.

#### **Examples**

mbrainSub

Subset of 1k Brain Cells from an E18 Mouse

#### **Description**

1k Brain Cells from an E18 Mouse is a public dataset from 10X Genomics. This subset is the first 50,000 barcodes of original data.

## Usage

data(mbrainSub)

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#### **Format**

An object of class "dgCMatrix".

#### **Source**

1k Brain Cells from an E18 Mouse

#### **Examples**

```
data(mbrainSub)
str(mbrainSub)
```

QuickCB2

All-in-one function from raw data to filtered cell matrix

## Description

All-in-one function for scCB2. Take 10x output raw data as input and return either a matrix of real cells identified by CB2 or a Seurat object containing this matrix, which can be incorporated with downstream analysis using Seurat pipeline.

## Usage

```
QuickCB2(
   dir = NULL,
   h5file = NULL,
   FDR_threshold = 0.01,
   MTfilter = 1,
   MTgene = NULL,
   AsSeurat = FALSE,
   Ncores = 2,
   ...
)
```

## **Arguments**

dir

The directory of 10x output data. For Cell Ranger version <3, directory should include three files: barcodes.tsv, genes.tsv, matrix.mtx. For Cell Ranger version >=3, directory should include three files: barcodes.tsv.gz, features.tsv.gz, matrix.mtx.gz.

h5file

The path of 10x output HDF5 file (ended with .h5).

FDR\_threshold

Numeric between 0 and 1. Default: 0.01. The False Discovery Rate (FDR) to be controlled for multiple testing.

MTfilter

Numeric value between 0 and 1. Default: 1 (No filtering). For each barcode, if the proportion of mitochondrial gene expression exceeds MTfilter, this barcode will be filtered out. No barcode exceeds 100% mitochondrial gene expression, thus the default (100%) corresponds to no filtering. The proportion of mitochondrial gene expressions is usually a criterion for evaluating cell quality, and is calculated using the scaled sum of all genes starting with "MT-" (human) or "mt-" (mouse) if row names are gene symbols, or customized mitochondrial genes specified by MTgene.

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MIgene	Character vector. User may specify customized mitochondrial gene IDs to perform the filtering. This should correspond to a subset of row names in raw data.
AsSeurat	Logical. Default: FALSE. Decides whether a Seurat object is returned instead of cell matrix. Set to TRUE so that users can directly apply Seurat pipeline for downstream analyses.
Ncores	Positive integer. Default: detectCores() - 2. Number of cores for parallel computation.
	Additional arguments to be passed to CB2FindCell.

#### **Details**

QuickCB2 is a quick function to apply CB2 on 10x Cell Ranger raw data by combining Read10xRaw, Read10xRawH5, CB2FindCell and GetCellMat into one simple function under default parameters.

#### Value

Either a sparse matrix of real cells identified by CB2 or a Seurat object containing real cell matrix.

#### **Examples**

```
# simulate 10x output files
data(mbrainSub)
mbrainSub <- mbrainSub[,1:10000]</pre>
data_dir <- file.path(tempdir(),"CB2example")</pre>
dir.create(data_dir)
gene_name <- rownames(mbrainSub)</pre>
# For simplicity, use gene names to generate gene IDs to fit the format.
gene_id <- paste0("ENSG_fake_",gene_name)</pre>
barcode_id <- colnames(mbrainSub)</pre>
Matrix::writeMM(mbrainSub,file = file.path(data_dir,"matrix.mtx"))
write.table(barcode_id,file = file.path(data_dir, "barcodes.tsv"),
    sep = "\t", quote = FALSE, col.names = FALSE, row.names = FALSE)
write.table(cbind(gene_id,gene_name),file = file.path(data_dir,"genes.tsv"),
    sep = "\t", quote = FALSE, col.names = FALSE, row.names = FALSE)
# Run QuickCB2 on 10x raw data and get cell matrix.
# Control FDR at 1%. Use 2-core parallel computation.
RealCell <- QuickCB2(dir = data_dir,</pre>
                      FDR_threshold = 0.01,
                      Ncores = 2)
str(RealCell)
```

Read10xRaw

Read 10x output data

## **Description**

Read10xRaw is a one-line handy function for reading 10x Cell Ranger output data, producing a count matrix for input to CB2FindCell. Read10xRawH5 is for reading 10x Cell Ranger output HDF5 file (ended with .h5). Works under both old (<3) and new (>=3) Cell Ranger version.

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#### Usage

```
Read10xRaw(dir = NULL, row.name = "symbol", meta = FALSE)
Read10xRawH5(h5file, row.name = "symbol", meta = FALSE)
```

#### **Arguments**

dir The directory of 10x output data. For Cell Ranger version <3, directory should

include three files: barcodes.tsv, genes.tsv, matrix.mtx. For Cell Ranger version >=3, directory should include three files: barcodes.tsv.gz, features.tsv.gz,

matrix.mtx.gz.

row.name Specify either using gene symbols (row.name = "symbol") or gene Ensembl

IDs (row.name = "id") as row names of the count matrix. Default is row.name

= "symbol".

meta Logical. If TRUE, returns a list containing both the count matrix and metadata

of genes (features). Metadata includes feature names, IDs and other additional information depending on Cell Ranger output. If FALSE (default), only returns

the count matrix.

h5file The path of 10x output HDF5 file (ended with .h5).

#### Value

If meta = TRUE, returns a list of two elements: a "dgCMatrix" sparse matrix containing expression counts and a data frame containing metadata of genes (features). For the count matrix, each row is a gene (feature) and each column is a barcode. If meta = FALSE, only returns the count matrix.

## **Examples**

```
# simulate 10x output files
data(mbrainSub)
data_dir <- file.path(tempdir(), "CB2example")</pre>
dir.create(data_dir)
gene_name <- rownames(mbrainSub)</pre>
# For simplicity, use gene names to generate gene IDs to fit the format.
gene_id <- paste0("ENSG_fake_",gene_name)</pre>
barcode_id <- colnames(mbrainSub)</pre>
Matrix::writeMM(mbrainSub,file = file.path(data_dir,"matrix.mtx"))
write.table(barcode_id,file = file.path(data_dir,"barcodes.tsv"),
    sep = "\t", quote = FALSE, col.names = FALSE, row.names = FALSE)
write.table(cbind(gene_id,gene_name),file = file.path(data_dir,"genes.tsv"),
    sep = "\t", quote = FALSE, col.names = FALSE, row.names = FALSE)
# read files
list.files(data_dir)
mbrainSub_new <- Read10xRaw(data_dir)</pre>
str(mbrainSub_new)
identical(mbrainSub, mbrainSub_new)
```

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