

Package ‘Cepo’

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Title Cepo for the identification of differentially stable genes

Version 1.15.0

Description

Defining the identity of a cell is fundamental to understand the heterogeneity of cells to various environmental signals and perturbations. We present Cepo, a new method to explore cell identities from single-cell RNA-sequencing data using differential stability as a new metric to define cell identity genes. Cepo computes cell-type specific gene statistics pertaining to differential stable gene expression.

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methods, SingleCellExperiment, SummarizedExperiment, ggplot2,
rlang, grDevices, patchwork, reshape2, BiocParallel, stats,
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VignetteBuilder knitr

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cellbench	<i>cellbench</i>
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Description

A single-cell RNA-seq dataset adapted from [sc_mixology](#)

Usage

`data(cellbench)`

Format

An object of SingleCellExperiment class with 895 cells and 2001 genes.

Source

https://github.com/LuyiTian/sc_mixology

Cepo	<i>Computing Cepo cell identity genes</i>
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Description

ExprsMat accepts various matrix objects, including DelayedArray and HDF5Array for out-of-memory computations. See vignette.

Usage

```
Cepo(  
  exprsMat,  
  cellTypes,  
  minCells = 20,  
  minCelltype = 3,  
  exprsPct = 0.05,  
  prefilter_sd = NULL,  
  prefilter_pzero = NULL,  
  logfc = NULL,  
  computePvalue = NULL,  
  computeFastPvalue = TRUE,
```

```

    variability = "CV",
    method = "weightedMean",
    weight = c(0.5, 0.5),
    workers = 1L,
    block = NULL,
    ...
)

```

Arguments

<code>exprsMat</code>	Expression matrix where columns denote cells and rows denote genes
<code>cellTypes</code>	Vector of cell type labels
<code>minCells</code>	Integer indicating the minimum number of cells required within a cell type
<code>minCelltype</code>	Integer indicating the minimum number of cell types required in each batch
<code>exprsPct</code>	Percentage of lowly expressed genes to remove. Default to NULL to not remove any genes.
<code>prefilter_sd</code>	Numeric value indicating threshold relating to standard deviation of genes. Used with <code>prefilter_zeros</code> .
<code>prefilter_pzero</code>	Numeric value indicating threshold relating to the percentage of zero expression of genes. Used with <code>prefilter_sd</code> .
<code>logfc</code>	Numeric value indicating the threshold of log fold-change to use to filter genes.
<code>computePvalue</code>	Whether to compute p-values using bootstrap test. Default to NULL to not make computations. Set this to an integer to set the number of bootstraps needed (recommend to be at least 100).
<code>computeFastPvalue</code>	Logical vector indicating whether to perform a faster version of p-value calculation. Set to TRUE by default.
<code>variability</code>	A character indicating the stability measure (CV, IQR, MAD, SD). Default is set to CV.
<code>method</code>	Character indicating the method for integration the two stability measures. By default this is set to 'weightedMean' with equal weights.
<code>weight</code>	Vector of two values indicating the weights for each stability measure. By default this value is <code>c(0.5, 0.5)</code> .
<code>workers</code>	Number of cores to use. Default to 1, which invokes <code>BiocParallel::SerialParam</code> . For workers greater than 1, see the <code>workers</code> argument in <code>BiocParallel::MulticoreParam</code> and <code>BiocParallel::SnowParam</code> .
<code>block</code>	Vector of batch labels
<code>...</code>	Additional arguments passed to <code>BiocParallel::MulticoreParam</code> and <code>BiocParallel::SnowParam</code> .

Value

Returns a list of key genes.

Examples

```

library(SingleCellExperiment)
data('cellbench', package = 'Cepo')
cellbench

```

```
cepoOutput <- Cepo(logcounts(cellbench), cellbench$celltype)
cepoOutput
```

plotDensities	<i>Plot densities</i>
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Description

Plot densities

Usage

```
plotDensities(
  x,
  cepoOutput,
  nGenes = 2,
  assay = "logcounts",
  celltypeColumn,
  celltype = NULL,
  genes = NULL,
  plotType = c("histogram", "density"),
  color = NULL
)
```

Arguments

x	a SummarizedExperiment or a SingleCellExperiment object.
cepoOutput	an output from Cepo or doLimma/doVoom/doTtest/doWilcoxon functions
nGenes	number of top genes from each celltype to plot. Default to 2.
assay	a character ('logcounts' by default), indicating the name of the assays(x) element which stores the expression data (i.e., assays(x)\$name_assays_expression). We strongly encourage using normalized data, such as counts per million (CPM) or log-CPM.
celltypeColumn	a character, indicating the name of the name of the cell type column in the col-Data(x).
celltype	a character, indicating the name of the cell type to plot. Default is NULL which selects all celltypes in the cepoOutput.
genes	a character vector, indicating the name of the genes to plot. Default to NULL, so that 2 top genes from each celltype will be plotted.
plotType	Either 'histogram' or 'density'
color	a named color vector. The names should correspond to the celltype argument above

Value

A [ggplot](#) object with cell-type specific densities for a gene.

A [ggplot](#) object.

Examples

```
library(SingleCellExperiment)
data('cellbench', package = 'Cepo')
cellbench
cepoOutput <- Cepo(logcounts(cellbench), cellbench$celltype)

plotDensities(
  x = cellbench,
  cepoOutput = cepoOutput,
  assay = 'logcounts',
  plotType = 'histogram',
  celltypeColumn = 'celltype'
)

plotDensities(
  x = cellbench,
  cepoOutput = cepoOutput,
  genes = c('PLTP', 'CPT1C', 'MEG3', 'SYCE1', 'MICOS10P3', 'HOXB7'),
  assay = 'logcounts',
  plotType = 'histogram',
  celltypeColumn = 'celltype'
)
```

sce_pancreas

sce_pancreas

Description

A subsampled single-cell RNA-seq dataset

Usage

```
data(sce_pancreas)
```

Format

An object of SingleCellExperiment class with 528 cells and 1358 genes.

setCepoBPPARAM

Setting parallel params based on operating platform

Description

Setting parallel params based on operating platform

Usage

```
setCepoBPPARAM(workers = 1L, ...)
```

Arguments

- workers Number of cores to use. Default to 1, which invokes BiocParallel::SerialParam. For workers greater than 1, see the workers argument in BiocParallel::MulticoreParam and BiocParallel::SnowParam.
- ... Additional arguments passed to BiocParallel::MulticoreParam and BiocParallel::SnowParam.

Value

Parameters for parallel computing depending on OS

Examples

```
# system.time(BiocParallel::bplapply(1:3, FUN = function(i){Sys.sleep(i)},
# BPPARAM = setCepoBPPARAM(workers = 1)))
# system.time(BiocParallel::bplapply(1:3, FUN = function(i){Sys.sleep(i)},
# BPPARAM = setCepoBPPARAM(workers = 3)))
```

topGenes	<i>Extract the top genes from the Cepo output</i>
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Description

Extract the top genes from the Cepo output

Usage

```
topGenes(object, n = 5, returnValues = FALSE)
```

Arguments

- object Output from the Cepo function
- n Number of top genes to extract
- returnValues Whether to return the numeric value associated with the top selected genes

Value

Returns a list of key genes.

Examples

```
set.seed(1234)
n <- 50 ## genes, rows
p <- 100 ## cells, cols
exprsMat <- matrix(rpois(n * p, lambda = 5), nrow = n)
rownames(exprsMat) <- paste0('gene', 1:n)
colnames(exprsMat) <- paste0('cell', 1:p)
cellTypes <- sample(letters[1:3], size = p, replace = TRUE)
cepo_output <- Cepo(exprsMat = exprsMat, cellTypes = cellTypes)
cepo_output
topGenes(cepo_output, n = 2)
topGenes(cepo_output, n = 2, returnValues = TRUE)
```

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