

Package ‘Cepo’

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Title Cepo for the identification of differentially stable genes

Version 1.4.0

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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Imports DelayedMatrixStats, DelayedArray, HDF5Array, S4Vectors, methods, SingleCellExperiment, SummarizedExperiment, ggplot2, rlang, grDevices, patchwork, reshape2, BiocParallel, stats, dplyr

biocViews Classification, GeneExpression, SingleCell, Software, Sequencing, DifferentialExpression

Suggests knitr, rmarkdown, BiocStyle, testthat, covr, UpSetR, scater, scMerge, fgsea, escape, pheatmap, patchwork

VignetteBuilder knitr

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cellbench *cellbench*

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 *Computing Cepo cell identity genes*

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 = NULL,
  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>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> .

block Vector of batch labels
... Additional arguments passed to BiocParallel::MulticoreParam and BiocParallel::SnowParam.
prefilter_pzeros Numeric value indicating threshold relating to the percentage of zero expression of genes. Used with prefilter_sd.

Value

Returns a list of key genes.

Examples

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

plotDensities *Plot densities*

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 colData(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>
----------	---

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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