Package 'scImpute'

May 24, 2017

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Type Package
Title Accurate and robust imputation of single-cell RNA sequencing data
Version 0.0.1
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Description scRNA-seq analysis is complicated by the excess of zero or near zero counts in the data, which are the so-called dropouts due to low amounts of mRNA within each individual cell. scImpute is developed to simultaneously determine which expression values are affected by dropout events in scRNA-seq data and perform imputation only on dropout entries.
Depends R (>= 3.3.2), glmnet, maxLik, parallel, stats, utils
License GPL
RoxygenNote 6.0.1
Suggests knitr, rmarkdown
VignetteBuilder knitr
R topics documented: scimpute
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scimpute use SCimpute to impute dropout values in scRNA-seq data
Description
use SCimpute to impute dropout values in scRNA-seq data
Usage
<pre>scimpute(count_path, infile = "csv", outfile = "csv", out_dir, drop_thre = 0.5, celltype = FALSE, labels = NULL, ncores = 5)</pre>

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Arguments

count_path	A character specifying the full path of the raw count matrix;
infile	A character specifying the type of file storing the raw count matrix; can be either "csv" or "txt". The input file shoule have rows representing genes and columns representing cells, with its first row as cell names and first column as gene names.
outfile	A character specifying the type of file storing the imputed count matrix; can be either "csv" or "txt".
out_dir	A character specifying the full path of the output directory, which is used to store all intermdediate and final outputs.
drop_thre	A number between 0 and 1, specifying the threshold to determine dropout values.
celltype	A logical value indicating whether cell type information is available. labels must be specified if celltype = TRUE.
labels	A character vector specifying the cell type of each column in the raw count matrix. Only needed when celltype = TRUE. Each cell type should have at least two cells for imputation.
ncores	A integer specifying the number of cores used for parallel computation.

Value

 $Save the imputed count \ matrix \ to \ SC impute.csv \ or \ SC impute.txt \ (depending \ on \ out \texttt{file}) \ to \ out \texttt{_dir}.$

scimpute_quick	quick re-run of SCimpute with a different drop_thre	

Description

quick re-run of SCimpute with a different drop_thre

Usage

```
scimpute_quick(count_path, infile = "csv", outfile = "csv", out_dir,
  drop_thre = 0.5, celltype = FALSE, labels = NULL, ncores = 5)
```

Arguments

count_path	A character specifying the full path of the raw count matrix;
infile	A character specifying the type of file storing the raw count matrix; can be either "csv" or "txt". The input file shoule have rows representing genes and columns representing cells, with its first row as cell names and first column as gene names.
outfile	A character specifying the type of file storing the imputed count matrix; can be either "csv" or "txt".
out_dir	A character specifying the full path of the output directory, which is used to store all intermdediate and final outputs.
drop_thre	A number between 0 and 1, specifying the threshold to determine dropout values.

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celltype A logical value indicating whether cell type information is available. labels

must be specified if celltype = TRUE.

labels A character vector specifying the cell type of each column in the raw count

matrix. Only needed when celltype = TRUE. Each cell type should have at

least two cells for imputation.

ncores A integer specifying the number of cores used for parallel computation.

Value

Save the imputed count matrix to SC impute.csv or SC impute.txt (depending on outfile) to out_dir .

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