Package 'scKWARN'

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Type Package	
Title Single-cell Technique	RNA Sequencing Normalization Using A Local Average
Version 1.0.7	
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Description A n	ormalization method for single-cell RNA sequencing data.
License GPL-2	GPL-3
Imports Rcpp (>	= 1.0.1), Matrix, stats, methods
LinkingTo Rcpp	
LazyData true	
RoxygenNote 6.	1.1
Archs x64	
R topics do	cumentea: N
LocASN	Single-cell RNA sequencing normalization using a local average technique
Description	
A function of	f normalizing single cell RNA-seq gene expression.
Usage	
gene	ntmatrix, conditions = NULL, filter = FALSE, e_num_gezero = 3, cell_num_gezero = 10, GeneforEst = 2000, divideforFast = TRUE, numDivide = NULL)

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Arguments

countmatrix Input. Unnormalized count (sparse) matrix (genes by cells).

conditions Input (Optional). Indicate which cells are sampled from the same conditions.

The default value, NULL, denotes all the cells are sampled from the same con-

dition.

filter Input (Optional). A logic value to indicate if need data filtering. If yes, please

see the details of gene_num_gezero and cell_num_gezero for input. The default

value if FALSE.

gene_num_gezero

Input (Optional). A threshold (interger) to determine the inclusion of a gene. The gene included needs to be expressed in at least <code>gene_num_gezero</code> cells. The

default value is 3.

cell_num_gezero

Input (Optional). A threshold (interger) to determine the inclusion of a cell. The cell included needs to contain at least *cell_num_gezero* expressed genes. The

default value is 10.

numGeneforEst Input (Optional). Use top *numGeneforEst* (integer) genes detected in most cells

to estimate scaling factors. The default value is 2000.

divideforFast Input (Optional). A logic value to indicate if speeding up computation by ran-

domly dividing cells in each condition into *numDivide* smaller groups. Please input an integer in *numDivide* below if *divideforFast* = TRUE. The default value

is TRUE.

numDivide Input (Optional). An integer is required if divideforFast = TRUE. numDivide =

NULL denotes # of cells in each condition divided by 5K (i.e., no division for

less than 10K cells).

Value

NormalizedData Matrix (genes by cells). Data matrix after normalization.

scalingFactor Vector. Cell-specific scaling factors.

delete_genes Vector. Indeice of the genes deleted.

delete_cells Vector. Indeice of the cells deleted.

Examples

```
set.seed(12345)
G <- 2000; n <- 600 # G: number of genes, n: number of cells
NB_cell <- function(j) rnbinom(G, size = 0.1, mu = rgamma(G, shape = 2, rate = 2))
countsimdata <- sapply(1:n, NB_cell)
colnames(countsimdata) <- paste("cell", 1:n, sep = "_")
rownames(countsimdata) <- paste("gene", 1:G, sep = "_")
Result <- LocASN(countmatrix = as(countsimdata, "sparseMatrix"))
Result$NormalizedData[1:10,1:10]; Result$scalingFactor[1:10]

#conditions <- c(rep(1,n/2), rep(2,n/2))
#Result2 <- LocASN(countmatrix = countsimdata, conditions = conditions)
#Result2$NormalizedData[1:10,1:10]; Result2$scalingFactor[1:10]</pre>
```

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