# Package 'SiPSiC'

May 10, 2024

Type Package

Title Calculate Pathway Scores for Each Cell in scRNA-Seq Data Version 1.4.0 Date 2022-12-29 **Description** Infer biological pathway activity of cells from single-cell RNA-sequencing data by calculating a pathway score for each cell (pathway genes are specified by the user). It is recommended to have the data in Transcripts-Per-Million (TPM) or Counts-Per-Million (CPM) units for best results. Scores may change when adding cells to or removing cells off the data. SiPSiC stands for Single Pathway analysis in Single Cells. License file LICENSE URL https://doi.org/10.1101/2023.03.27.534310 biocViews Software, DifferentialExpression, GeneSetEnrichment, BiomedicalInformatics, CellBiology, Transcriptomics, RNASeq, SingleCell, Transcription, Sequencing, ImmunoOncology, DataImport Depends Matrix, SingleCellExperiment Suggests knitr, rmarkdown, BiocStyle VignetteBuilder knitr BugReports https://github.com/DanielDavis12/SiPSiC/issues git\_url https://git.bioconductor.org/packages/SiPSiC git\_branch RELEASE\_3\_19 git\_last\_commit 092f808 git\_last\_commit\_date 2024-04-30 Repository Bioconductor 3.19 Date/Publication 2024-05-09

**Author** Daniel Davis [aut, cre] (<a href="https://orcid.org/0000-0001-7863-1521">https://orcid.org/0000-0001-7863-1521</a>),

Maintainer Daniel Davis < Daniel Davis 000@gmail.com>

Yotam Drier [aut]

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#### **Description**

Infer biological pathway activity of cells from single-cell RNA-sequencing data by calculating a pathway score for each cell (pathway genes are specified by the user). It is recommended to have the data in Transcripts-Per-Million (TPM) or Counts-Per-Million (CPM) units for best results. Scores may change when adding cells to or removing cells off the data. SiPSiC stands for Single Pathway analysis in Single Cells.

# **Details**

Use this package to calculate per-cell scores for a biological pathway of your choice, from single-cell RNA-seq data. Transcripts-Per-Million (TPM) or Counts-Per-Million (CPM) units of the data are recommended for best results.

#### Author(s)

Daniel Davis, Yotam Drier. Maintainer: Daniel Davis

#### References

https://medicine.ekmd.huji.ac.il/en/research/yotamd/Pages/default.aspx

# See Also

getPathwayScores

getPathwayScores 3

getPathwayScores Calculate Pathway Scores for all Cells in a scRNA-Seq Data

# **Description**

Calculate a pathway score for each cell included in the input scRNA-seq data, using the cell's transcription level of the pathway's genes.

#### Usage

getPathwayScores(dataMatrix, pathwayGenes, percentForNormalization)

# **Arguments**

a list of three items:

dataMatrix a matrix whose rows and columns represent genes and cells, respectively, con-

taining the scRNA-seq data; Counts-Per-Million (CPM) or Transcripts-Per-Million (TPM) units should be used for best results. The matrix should be of type sparse

matrix (dgCMatrix), otherwise errors might be raised.

pathwayGenes a character vector of the gene names of which the relevant biological pathway

consists.

percentForNormalization

The percent of top cells for each gene whose median is used to normalize the

gene's expression values (5 by default).

#### Value

pathwayScores an array (type double) of the pathway score of each cell in the input dataMatrix,

less than two of the pathway genes are found in the data, in which case NA is

returned.

index a numeric array of the row indices in the dataMatrix where genes of the pathway

were found.

#### Author(s)

Daniel Davis, Yotam Drier

# References

https://medicine.ekmd.huji.ac.il/en/research/yotamd/Pages/default.aspx

normalizeCountsForGene

# **Examples**

```
library(SiPSiC)
geneCountsMatrix <- matrix(rpois(16, lambda = 10), ncol = 4, nrow = 4)
geneCountsMatrix <- as(geneCountsMatrix, "dgCMatrix")
rownames(geneCountsMatrix) <- c("Gene1", "Gene2", "Gene3", "Gene4")
colnames(geneCountsMatrix) <- c("Cell1", "Cell2", "Cell3", "Cell4")
assayData <- SingleCellExperiment(assays = list(counts = geneCountsMatrix))
pathwayGenesList <- c("Gene1", "Gene2", "Gene4")
percentForNormalization <- 7
scoresAndIndices <- getPathwayScores(counts(assayData), pathwayGenesList, percentForNormalization)
pathwayScoresOfCells <- scoresAndIndices$pathwayScores
pathwayGeneIndices <- scoresAndIndices$index</pre>
```

normalizeCountsForGene

Gene counts normalization

#### **Description**

Get the counts of a single gene normalized by the median of the top 5 percent cells, unless it's zero; In this case, the counts are all divided by the maximum value across all cells. If all counts are zeros, they are returned untouched.

# Usage

normalizeCountsForGene(expressionValues, percentForNormalization)

## **Arguments**

expressionValues

An array of type double, containing the counts (in any units, e.g. CPM or TPM) of a single gene across different cells.

percentForNormalization

The percent of top cells for each gene whose median is used to normalized the gene's expression values.

# Value

An array (type double) of the normalized input counts.

## Author(s)

Daniel Davis, Yotam Drier

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