

Package ‘netSmooth’

July 16, 2025

Type Package

Title Network smoothing for scRNAseq

Version 1.29.0

Description netSmooth is an R package for network smoothing of single cell RNA sequencing data. Using bio networks such as protein-protein interactions as priors for gene co-expression, netsmooth improves cell type identification from noisy, sparse scRNAseq data.

biocViews Network, GraphAndNetwork, SingleCell, RNASeq, GeneExpression, Sequencing, Transcriptomics, Normalization, Preprocessing, Clustering, DimensionReduction

URL <https://github.com/BIMSBbioinfo/netSmooth>

BugReports <https://github.com/BIMSBbioinfo/netSmooth/issues>

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Encoding UTF-8

LazyData true

Depends R (>= 3.5), scatter (>= 1.15.11), clusterExperiment (>= 2.1.6)

Imports entropy, SummarizedExperiment, SingleCellExperiment, Matrix, cluster, data.table, stats, methods, DelayedArray, HDF5Array (>= 1.15.13)

Suggests knitr, testthat, Rtsne, biomaRt, igraph, STRINGdb, NMI, pheatmap, ggplot2, BiocStyle, rmarkdown, BiocParallel, uwot

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calc2DEntropy	<i>Calculate entropy in 2D data</i>
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Description

Calculate entropy in 2D data

Usage

```
calc2DEntropy(x, numBins1 = 20, numBins2 = 20)
```

Arguments

- | | |
|----------|---|
| x | the 2D data to get entropy from |
| numBins1 | the number of bins along the first dimension to discretize data into |
| numBins2 | the number of bins along the second dimension to discretize data into |

Value

The Shannon entropy in the 2D data x

`clusterExperimentWorkflow`*Performs clustering workflow using ‘clusterExperiment’ functions*

Description

Performs clustering workflow using ‘clusterExperiment’ functions

Usage

```
clusterExperimentWorkflow(  
  se,  
  dimReduceFlavor = c("pca", "tsne", "dm", "umap"),  
  cluster.ks = 5:10,  
  cluster.function = "pam",  
  nVarDims = c(100, 500, 1000),  
  makeConsensusProportion = 0.7,  
  makeConsensusMinSize = 4,  
  runMergeClusters = TRUE,  
  is.counts = TRUE,  
  random.seed = 1  
)
```

Arguments

<code>se</code>	SummarizedExperiment object
<code>dimReduceFlavor</code>	algorithm for reduced dimension embedding step
<code>cluster.ks</code>	range of Ks to cluster over
<code>cluster.function</code>	clustering algorithm to use for all clusterings
<code>nVarDims</code>	numbers of variable genes to perform clusterings over
<code>makeConsensusProportion</code>	proportion of times samples need to be co-clustered for co-clustering step
<code>makeConsensusMinSize</code>	minimum cluster size
<code>runMergeClusters</code>	logical: merge similar clusters
<code>is.counts</code>	logical: is data counts
<code>random.seed</code>	passed to clusterExperiment. set to NULL in order to not set a random seed.

Value

cluster assignments

clusterOne	<i>Run one clustering using kmeans o PAM</i>
------------	--

Description

Run one clustering using kmeans o PAM

Usage

```
clusterOne(x, algorithm = c("kmeans", "pam"), k = 5)
```

Value

kmeans or PAM cluster assignments

dimReduce	<i>Get lower dimension embedding</i>
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Description

Get lower dimension embedding

Usage

```
dimReduce(
  x,
  flavor = c("pca", "tsne", "umap"),
  k = 2,
  is.counts = TRUE,
  ntop = 500
)
```

Arguments

x	gene expresison matrix [GENES x SAMPLES]
flavor	the algorithm to use to obtain the dimensionality reduction must be in c('pca', 'tsne', 'umap')
k	the number of dimensions in the reduced dimension representation
is.counts	logical: is 'x' counts data
ntop	number of most variable genes to use for dimensionality reduction

Value

reduced dimensionality representation

`human.ppi`*Human Protein-Protein interaction graph*

Description

An adjacency matrix of the 10 percent highest confidence interactions between human proteins on STRINGdb.

Usage

```
human.ppi
```

Format

A square matrix where $A_{ij}=1$ if gene i interacts with gene j

Details

See the script in `'system.file(package="netSmooth", "data-raw", "make_ppi_from_string.R")'` for full details of how this object was made.

Source

<http://www.string-db.org/>

`l1NormalizeColumns`*Column-normalize a sparse, symmetric matrix (using the l1 norm) so that each column sums to 1.*

Description

Column-normalize a sparse, symmetric matrix (using the l1 norm) so that each column sums to 1.

Usage

```
l1NormalizeColumns(A)
```

Arguments

`A` matrix

Value

column-normalized sparse matrix object

l1NormalizeRows	<i>Row-normalize a sparse, symmetric matrix (using the l1 norm) so that each row sums to 1.</i>
-----------------	---

Description

Row-normalize a sparse, symmetric matrix (using the l1 norm) so that each row sums to 1.

Usage

```
l1NormalizeRows(A)
```

Arguments

A matrix

Value

row-normalized sparse matrix object

mouse.ppi	<i>Mouse Protein-Protein interaction graph</i>
-----------	--

Description

An adjacency matrix of the 10 percent highest confidence interactions between mouse proteins on STRINGdb.

Usage

```
mouse.ppi
```

Format

A square matrix where $A_{ij}=1$ if gene i interacts with gene j

Details

See the script in ‘system.file(package="netSmooth", "data-raw", "make_ppi_from_string.R")’ for full details of how this object was made.

Source

<http://www.string-db.org/>

`netSmooth,matrix-method`*Perform network smoothing of gene expression or other omics data*

Description

Perform network smoothing of gene expression or other omics data

Usage

```
## S4 method for signature 'matrix'
netSmooth(
  x,
  adjMatrix,
  alpha = "auto",
  normalizeAdjMatrix = c("rows", "columns"),
  autoAlphaMethod = c("robustness", "entropy"),
  autoAlphaRange = 0.1 * (seq_len(9)),
  autoAlphaDimReduceFlavor = "auto",
  is.counts = TRUE,
  bpparam = BiocParallel::SerialParam(),
  ...
)

## S4 method for signature 'SummarizedExperiment'
netSmooth(x, ...)

## S4 method for signature 'SingleCellExperiment'
netSmooth(x, ...)

## S4 method for signature 'Matrix'
netSmooth(
  x,
  adjMatrix,
  alpha = "auto",
  normalizeAdjMatrix = c("rows", "columns"),
  autoAlphaMethod = c("robustness", "entropy"),
  autoAlphaRange = 0.1 * (seq_len(9)),
  autoAlphaDimReduceFlavor = "auto",
  is.counts = TRUE,
  bpparam = BiocParallel::SerialParam(),
  ...
)

## S4 method for signature 'DelayedMatrix'
netSmooth(
  x,
  adjMatrix,
  alpha = "auto",
  normalizeAdjMatrix = c("rows", "columns"),
  autoAlphaMethod = c("robustness", "entropy"),
```

```

autoAlphaRange = 0.1 * (seq_len(9)),
autoAlphaDimReduceFlavor = "auto",
is.counts = TRUE,
bpparam = BiocParallel::SerialParam(),
filepath = NULL,
...
)

```

Arguments

<code>x</code>	matrix or SummarizedExperiment
<code>adjMatrix</code>	adjacency matrix of gene network to use
<code>alpha</code>	numeric in [0,1] or 'auto'. if 'auto', the optimal value for alpha will be automatically chosen among the values specified in 'autoAlphaRange', using the strategy specified in 'autoAlphaMethod'
<code>normalizeAdjMatrix</code>	how to normalize the adjacency matrix possible values are 'rows' (in-degree) and 'columns' (out-degree)
<code>autoAlphaMethod</code>	if 'robustness', pick alpha that gives the highest proportion of samples in robust clusters if 'entropy', pick alpha that gives highest Shannon entropy in 2D PCA embedding
<code>autoAlphaRange</code>	if 'alpha='optimal'', search these values for the best alpha
<code>autoAlphaDimReduceFlavor</code>	algorithm for dimensionality reduction that will be used to pick the optimal value for alpha. Either the 2D embedding to calculate the Shannon entropy for (if 'autoAlphaMethod='entropy''), or the dimensionality reduction algorithm to be used in robust clustering (if 'autoAlphamethod='robustness'')
<code>is.counts</code>	logical: is the assay count data
<code>bpparam</code>	instance of bpparam, for parallel computation with the 'alpha='auto'' option. See the BiocParallel manual.
<code>...</code>	arguments passed on to 'robustClusters' if using the robustness criterion for optimizing alpha
<code>filepath</code>	String: Path to location where hdf5 output file is supposed to be saved. Will be ignored when regular matrices or SummarizedExperiment are used as input.

Value

network-smoothed gene expression matrix or SummarizedExperiment object

Examples

```

x <- matrix(rnbinom(12000, size=1, prob = .1), ncol=60)
rownames(x) <- paste0('gene', seq_len(dim(x)[1]))

adj_matrix <- matrix(as.numeric(rnorm(200*200)>.8), ncol=200)
rownames(adj_matrix) <- colnames(adj_matrix) <- paste0('gene', seq_len(dim(x)[1]))
x.smoothed <- netSmooth(x, adj_matrix, alpha=0.5)

```

pickDimReduction,matrix-method

Pick the dimensionality reduction method for a dataset that gives the 2D embedding with the highest entropy

Description

Pick the dimensionality reduction method for a dataset that gives the 2D embedding with the highest entropy

Usage

```
## S4 method for signature 'matrix'
pickDimReduction(x, flavors = c("pca", "tsne", "umap"), is.counts = TRUE)

## S4 method for signature 'SummarizedExperiment'
pickDimReduction(x)

## S4 method for signature 'Matrix'
pickDimReduction(x, flavors = c("pca", "tsne", "umap"), is.counts = TRUE)

## S4 method for signature 'DelayedMatrix'
pickDimReduction(x, flavors = c("pca", "tsne", "umap"), is.counts = TRUE)
```

Arguments

x	matrix or SummarizedExperiment object [GENES x SAMPLES]
flavors	list of dimensionality reduction algorithms to try. Currently the options are "pca", "tsne" and "umap"
is.counts	logical: is exprs count data

Value

name of dimensionality reduction method that gives the highest 2d entropy

Examples

```
x <- matrix(rnbinom(60000, size=1, prob = .1), ncol=100)
pickDimReduction(x)
```

projectFromNetworkRecombine,matrix-method

Combine gene expression from smoothed space (that of the network) with the expression of genes that were not smoothed (not present in network)

Description

Combine gene expression from smoothed space (that of the network) with the expression of genes that were not smoothed (not present in network)

Usage

```
## S4 method for signature 'matrix'
projectFromNetworkRecombine(original_expression, smoothed_expression)
```

Arguments

`original_expression`
the non-smoothed expression

`smoothed_expression`
the smoothed gene expression, in the space of the genes defined by the network

`filepath` String: Path to location where hdf5 output file is supposed to be saved. Will be ignored when regular matrices or SummarizedExperiment are used as input.

Value

a matrix in the dimensions of `original_expression`, where values that are present in `smoothed_expression` are copied from there.

projectOnNetwork,matrix-method

Project the gene expression matrix onto a lower space of the genes defined in the smoothing network

Description

Project the gene expression matrix onto a lower space of the genes defined in the smoothing network

Usage

```
## S4 method for signature 'matrix'
projectOnNetwork(gene_expression, new_features, missing.value = 0)
```

Arguments

`gene_expression`
gene expression matrix

`new_features` the genes in the network, on which to project the gene expression matrix

`missing.value` value to assign to genes that are in network, but missing from gene expression matrix

Value

the gene expression matrix projected onto the gene space defined by `new_features`

randomWalkByIterations

Smooth data on graph by computing iterations

Description

Smooth data on graph by computing iterations

Usage

```
randomWalkByIterations(
  f0,
  adjMatrix,
  alpha,
  normalizeAdjMatrix = c("rows", "columns"),
  tol = 1e-06,
  max.iter = 100
)
```

Arguments

f0	initial data matrix [NxM]
adjMatrix	adjacency matrix of graph to network smooth on will be column-normalized.
alpha	smoothing coefficient (1 - restart probability of random walk)
tol	the tolerance (stopping criterion)
max.iter	the maximum number of iterations before terminating

Value

network-smoothed gene expression

randomWalkByMatrixInv, matrix-method

Smooth data on graph by computing the closed-form steady state distribution of the random walk with restarts process.

Description

The closed-form solution is given by $f_{ss} = (I - \alpha) * (I - \alpha * A)^{-1} * f_0$ and is computed by matrix inversion in this function.

Usage

```
## S4 method for signature 'matrix'
randomWalkByMatrixInv(
  f0,
  adjMatrix,
  alpha,
  normalizeAdjMatrix = c("rows", "columns")
)
```

Arguments

`f0` initial data matrix [NxM]
`adjMatrix` adjacency matrix of graph to network smooth on will be column-normalized.
`alpha` smoothing coefficient (1 - restart probability of random walk)

Value

network-smoothed gene expression

randomWalkBySolve, matrix-method

*Smooth data on graph by solving the linear equation $(I - \alpha * A) * E_{sm} = E * (1 - \alpha)$*

Description

Smooth data on graph by solving the linear equation $(I - \alpha * A) * E_{sm} = E * (1 - \alpha)$

Usage

```
## S4 method for signature 'matrix'
randomWalkBySolve(E, A, alpha, normalizeAdjMatrix = c("rows", "columns"))
```

Arguments

`E` initial data matrix [NxM]
`A` adjacency matrix of graph to network smooth on will be column-normalized.
`alpha` smoothing coefficient (1 - restart probability of random walk)

Value

network-smoothed gene expression

robustClusters, SummarizedExperiment-method

Perform robust clustering on dataset, and calculate the proportion of samples in robust clusters

Description

Perform robust clustering on dataset, and calculate the proportion of samples in robust clusters

Usage

```
## S4 method for signature 'SummarizedExperiment'
robustClusters(x, dimReduceFlavor = "auto", is.counts = TRUE, ...)

## S4 method for signature 'matrix'
robustClusters(x, ...)
```

Arguments

<code>x</code>	matrix or SummarizedExperiment object
<code>dimReduceFlavor</code>	algorithm for dimensionality reduction step of clustering procedure. May be 'pca', 'tsne', 'dm', 'umap' or 'auto', which uses shannon entropy to pick the algorithm.
<code>is.counts</code>	logical: is the data counts
<code>...</code>	arguments passed on to 'clusterExperimentWorkflow'

Value

list(clusters, proportion.robust)

Examples

```
data("smallscRNAseq")
robustClusters(smallscRNAseq, dimReduceFlavor='pca')
```

scoreSmoothing	<i>Calculate a score for a smoothing result, for picking the best alpha value</i>
----------------	---

Description

Calculate a score for a smoothing result, for picking the best alpha value

Usage

```
scoreSmoothing(x, method = c("entropy", "robustness"), is.counts = TRUE, ...)
```

Arguments

<code>x</code>	the network-smoothed expression matrix
<code>method</code>	the scoring method. 'entropy' calculates shannon entropy in a 2D PCA of the data. 'robustness' performs robust clustering and reports the proportion of samples in robust clusters

Value

the score

smallPPI

A small human Protein-Protein interaction graph for use in examples.

Description

Contains a synthetic PPI of human genes.

Usage

smallPPI

Format

An object of class `matrix` with 611 rows and 611 columns.

smallscRNAseq

A small single cell RNA-seq dataset for use in examples.

Description

Contains scRNAseq profiles of human blastomeres.

Usage

smallscRNAseq

Format

SingleCellExperiment

Source

<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE44183>

smoothAndRecombine,matrix-method

Perform network smoothing on network when the network genes and the experiment genes aren't exactly the same.

Description

The gene network might be defined only on a subset of genes that are measured in any experiment. Further, an experiment might not measure all genes that are present in the network. This function projects the experiment data onto the gene space defined by the network prior to smoothing. Then, it projects the smoothed data back into the original dimensions.

Usage

```
## S4 method for signature 'matrix'
smoothAndRecombine(
  gene_expression,
  adj_matrix,
  alpha,
  smoothing.function = randomWalkBySolve,
  normalizeAdjMatrix = c("rows", "columns")
)
```

Arguments

gene_expression	gene expression data to be smoothed [N_genes x M_samples]
adj_matrix	adjacency matrix of network to perform smoothing over. Will be column-normalized. Rownames and colnames should be genes.
alpha	network smoothing parameter (1 - restart probability in random walk model.
smoothing.function	must be a function that takes in data, adjacency matrix, and alpha. Will be used to perform the actual smoothing.
normalizeAdjMatrix	which dimension (rows or columns) should the adjacency matrix be normalized by. rows corresponds to in-degree, columns to out-degree.
filepath	String: Path to location where hdf5 output file is supposed to be saved. Will be ignored when regular matrices or SummarizedExperiment are used as input.

Value

matrix with network-smoothed gene expression data. Genes that are not present in smoothing network will retain original values.

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