

Package ‘scapGNN’

June 27, 2022

Type Package

Title Graph Neural Network-Based Framework for Single Cell Active Pathways and Gene Modules Analysis

Version 0.1.1

Date 2022-06-27

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Description It is a single cell active pathway analysis tool based on the graph neural network (F. Scarselli (2009) <[doi:10.1109/TNN.2008.2005605](https://doi.org/10.1109/TNN.2008.2005605)>; Thomas N. Kipf (2017) <[arXiv:1609.02907v4](https://arxiv.org/abs/1609.02907v4)>) to construct the gene-cell association network, infer pathway activity scores from different single cell modalities data, integrate multiple modality data on the same cells into one pathway activity score matrix, identify cell phenotype activated gene modules and parse association networks of gene modules under multiple cell phenotype. In addition, abundant visualization programs are provided to display the results.

License GPL (>= 2)

Encoding UTF-8

LazyData true

Depends R (>= 4.1.0)

RoxygenNote 7.1.2

Imports ActivePathways, AdaptGauss, coop, igraph, mixtools,
reticulate, methods

Suggests rmarkdown, knitr

VignetteBuilder knitr

NeedsCompilation no

Repository CRAN

Date/Publication 2022-06-27 02:20:02 UTC

R topics documented:

ATAC_net	2
BIC_LTMG	3
BIC_ZIMG	3
ConNetGNN	4
ConNetGNN_data	6
cpGModule	6
create_scapGNN_env	8
Fit_LTMG	8
Global_Zcut	9
H9_0h_cpGM_data	9
H9_24h_cpGM_data	10
H9_36h_cpGM_data	10
Hv_exp	11
instPyModule	11
InteNet	12
isLoading	13
load_path_data	13
LTMG	14
LTMG-class	14
norm1	15
plotCCNetwork	15
plotGANetwork	17
plotMulPhenGM	19
Preprocessing	20
Pure_CDF	22
RNA_ATAC_IntNet	22
RNA_net	23
RunLTMG	23
RWR	24
scPathway	25
scPathway_data	26
Index	27

ATAC_net	<i>Results of ConNetGNN() for scATAC-seq data from SNARE-seq dataset</i>
----------	--

Description

A list to store the gene association network of scATAC-seq data. Case data from the SNARE-seq dataset.

Usage

ATAC_net

Format

a list of three adjacency matrices.

Examples

```
data(ATAC_net)
```

BIC_LTMG

BIC_LTMG

Description

The internal functions of the scapGNN package.

Usage

```
BIC_LTMG(y, rrr, Zcut)
```

Arguments

y	Internal parameters.
rrr	Internal parameters.
Zcut	Internal parameters.

Details

BIC_LTMG

BIC_ZIMG

BIC_ZIMG

Description

The internal functions of the scapGNN package.

Usage

```
BIC_ZIMG(y, rrr, Zcut)
```

Arguments

y	Internal parameters.
rrr	Internal parameters.
Zcut	Internal parameters.

Details

BIC_ZIMG

ConNetGNN	<i>Construct association networks for gene-gene, cell-cell, and gene-cell based on graph neural network (GNN)</i>
-----------	---

Description

This function implements a graph neural network with two autoencoders. 1. AutoEncoder (AE) based on deep neural network: Infer latent associations between genes and cells. 2. Graph AutoEncoder (GAE) based on graph convolutional neural network: Construct association networks for gene-gene, cell-cell.

Usage

```
ConNetGNN(
    Prep_data,
    python.path = NULL,
    miniconda.path = NULL,
    AE.epochs = 1000,
    AE.learning.rate = 0.001,
    AE.reg.alpha = 0.5,
    use.VGAE = TRUE,
    GAE.epochs = 300,
    GAE.learning.rate = 0.01,
    cell_val_ratio = 0.05,
    gene_val_ratio = 0.05,
    parallel = FALSE,
    seed = 125,
    verbose = TRUE
)
```

Arguments

Prep_data	The input data is the result from the Preprocessing function.
python.path	The path to a Python binary.
miniconda.path	The path in which miniconda will be installed. If the python.path is NULL and conda or miniconda is not installed in the system, the program will automatically install miniconda according to the path specified by miniconda.path.
AE.epochs	The number of epoch for the deep neural network (AE). Default: 1000.
AE.learning.rate	Initial learning rate of AE. Default: 0.001.
AE.reg.alpha	The LTMG regularized intensity. Default: 0.5.
use.VGAE	Whether to use Variational Graph AutoEncoder (VGAE). Default: TRUE.
GAE.epochs	The number of epoch for the GAE. Default: 300.
GAE.learning.rate	Initial learning rate of GAE. Default: 0.01.

<code>cell_val_ratio</code>	For GAE that construct cell-cell association networks, the proportion of edges that are extracted as the validation set. Default: 0.05.
<code>gene_val_ratio</code>	As with parameter <code>cell_val_ratio</code> , it is simply applied with the construction of gene-gene association networks.
<code>parallel</code>	Whether to use multiple processors to run GAE. Default: FALSE When <code>parallel=TRUE</code> (default), tow processors will be used to run GAE.
<code>seed</code>	Random number generator seed.
<code>verbose</code>	Gives information about each step. Default: TRUE.

Details

ConNetGNN

The ConNetGNN function establishes a graph neural network (GNN) framework to mine latent relationships between genes and cells and within themselves. This framework mainly includes two capabilities:

- 1.Deep neural network-based AutoEncoder inferring associations between genes and cells and generating gene features and cell features for the GAE.
- 2.The GAE takes the gene feature and cell feature as the node features of the initial gene correlation network and cell correlation network, and constructs the gene association network and cell association network through the graph convolution process.

The GNN is implemented based on pytorch, so an appropriate python environment is required:

- python \geq 3.9.7
- pytorch \geq 1.10.0
- sklearn \geq 0.0
- scipy \geq 1.7.3
- numpy \geq 1.19.5

If the user has already configured the python environment, the path of the python binary file can be directly entered into `python.path`. If the parameter `python.path` is NULL, the program will build a miniconda environment called `scapGNN_env` and configure python. We also provide environment files for conda: `/inst/extdata/scapGNN_env.yaml`. Users can install it with the command: `conda env create -f scapGNN_env.yaml`.

Value

A list:

cell_network Constructed cell association network.

gene_network Constructed gene association network.

cell_gene_network Constructed gene-cell association network.

Examples

```

require(coop)
require(reticulate)
require(parallel)
# Data preprocessing
data("Hv_exp")
Prep_data <- Preprocessing(Hv_exp[1:300,])
## Not run:
# Specify the python path
ConNetGNN_data <- ConNetGNN(Prep_data,python.path="../miniconda3/envs/scapGNN_env/python.exe")

## End(Not run)

```

ConNetGNN_data	<i>The results of ConNetGNN() function</i>
----------------	--

Description

Results of ConNetGNN() function with Hv_exp as input.

Usage

```
ConNetGNN_data
```

Format

a list.

Examples

```
data(ConNetGNN_data)
```

cpGModule	<i>Identify cell phenotype activated gene module</i>
-----------	--

Description

Mining activated gene modules in specific cell phenotype.

Usage

```
cpGModule(
  network.data,
  cellset,
  nperm = 100,
  cut.pvalue = 0.01,
  cut.fdr = 0.05,
  parallel.cores = 2,
  rwr.gamma = 0.7,
  verbose = TRUE
)
```

Arguments

<code>network.data</code>	Network data constructed by the ConNetGNN function.
<code>cellset</code>	A vector of cell id. The specified cell set, which will be used as the restart set.
<code>nperm</code>	Number of random permutations. Default: 100.
<code>cut.pvalue</code>	The threshold of P-value, and genes below this threshold are regarded as gene modules activated by the cell set. Default: 0.01.
<code>cut.fdr</code>	The threshold of false discovery rate (FDR), and genes below this threshold are regarded as gene modules activated by the cell set. Default: 0.05.
<code>parallel.cores</code>	Number of processors to use when doing the calculations in parallel (default: 2). If <code>parallel.cores=0</code> , then it will use all available core processors unless we set this argument with a smaller number.
<code>rwr.gamma</code>	Restart parameter. Default: 0.7.
<code>verbose</code>	Gives information about each step. Default: TRUE.

Details**cpGModule**

The `cpGModule` function takes a user-defined cell set as a restart set to automatically identify activated gene modules. A perturbation analysis was used to calculate a significant P-value for each gene. The Benjamini & Hochberg (BH) method was used to adjust the P-value to obtain the FDR. Genes with a significance level less than the set threshold are considered as cell phenotype activated gene modules.

Value

A data frame contains four columns:

Genes Gene ID.

AS Activity score.

Pvalue Significant P-value.

FDR False discovery rate.

Examples

```

require(parallel)
require(stats)

# Load the result of the ConNetGNN function.
data(ConNetGNN_data)
data(Hv_exp)

# Construct the cell set corresponding to 0h.
index<-grep("0h",colnames(Hv_exp))
cellset<-colnames(Hv_exp)[index]
cpGModule_data<-cpGModule(ConNetGNN_data,cellset,nperm=10,parallel.cores=1)

```

```

create_scapGNN_env      Create the create_scapGNN_env environment on miniconda

```

Description

The internal functions of the scapGNN package.

Usage

```
create_scapGNN_env()
```

Details

```
create_scapGNN_env
```

```

Fit_LTMG                Fitting function for Left-truncated mixed Gaussian

```

Description

The internal functions of the scapGNN package.

Usage

```
Fit_LTMG(x, n, q, k, err = 1e-10)
```

Arguments

x	Internal parameters.
n	Internal parameters.
q	Internal parameters.
k	Internal parameters.
err	Internal parameters.

Details

Fit_LTMG

Global_Zcut	<i>Global_Zcut</i>
-------------	--------------------

Description

The internal functions of the scapGNN package.

Usage

```
Global_Zcut(MAT, seed = 123)
```

Arguments

MAT	Internal parameters.
seed	Random number generator seed.

Details

Global_Zcut

H9_0h_cpGM_data	<i>Cell-activated gene modules under the 0-hour phenotype</i>
-----------------	---

Description

Results of cpGModule() function.

Usage

```
H9_0h_cpGM_data
```

Format

a list.

Examples

```
data(H9_0h_cpGM_data)
```

H9_24h_cpGM_data

Cell-activated gene modules under the 24-hour phenotype

Description

Results of cpGModule() function.

Usage

H9_24h_cpGM_data

Format

a list.

Examples

```
data(H9_24h_cpGM_data)
```

H9_36h_cpGM_data

Cell-activated gene modules under the 36-hour phenotype

Description

Results of cpGModule() function.

Usage

H9_36h_cpGM_data

Format

a list.

Examples

```
data(H9_36h_cpGM_data)
```

`Hv_exp`*Single-cell gene expression profiles*

Description

A log-transformed gene-cell matrix containing highly variable features.

Usage`Hv_exp`**Format**

a matrix.

Examples`data(Hv_exp)`

`instPyModule`*Install the pyhton module through the reticulate R package*

Description

The internal functions of the scapGNN package.

Usage`instPyModule(module)`**Arguments**

`module` Internal parameters.

Details`instPyModule`

 InteNet

Integrate network data from single-cell RNA-seq and ATAC-seq

Description

For the SNARE-seq dataset, a droplet-based method to simultaneously profile gene expression and chromatin accessibility in each of thousands of single nuclei, the InteNet function can integrate network data of scRNA-seq data and scATAC-seq data (results of the ConNetGNN function) to into a gene-cell network.

Usage

```
InteNet(RNA_net, ATAC_net, parallel.cores = 2, verbose = TRUE)
```

Arguments

RNA_net	Network data for RNA datasets. Produced by the ConNetGNN function.
ATAC_net	Network data for ATAC datasets. Produced by the ConNetGNN function.
parallel.cores	Number of processors to use when doing the calculations in parallel (default: 2). If parallel.cores=0, then it will use all available core processors unless we set this argument with a smaller number.
verbose	Gives information about each step. Default: TRUE.

Details

InteNet

The scATAC-seq dataset needs to be converted into a gene activity matrix according to the process of Signac(<https://satijalab.org/signac/articles/snareseq.html>). The subsequent process is consistent with the scRNA-seq dataset. The InteNet function then integrates the network data of RNA-seq data and ATAC-seq data into a gene-cell network. With integrated network data as input, scPathway and cpGModule functions will infer pathway activity score matrix and gene modules supported by single-cell multi-omics.

Value

A list.

Examples

```
require(ActivePathways)
require(parallel)
data(RNA_net)
data(ATAC_net)
## Not run:
RNA_ATAC_IntNet<-InteNet(RNA_net,ATAC_net,parallel.cores=1)

## End(Not run)
```

```
# View data
data(RNA_ATAC_IntNet)
summary(RNA_ATAC_IntNet)
```

isLoaded *The internal functions of the scapGNN package*

Description

Determine if the package is loaded.

Usage

```
isLoaded(name)
```

Arguments

name Internal parameters.

Details

isLoaded

load_path_data *load pathway or gene set's gmt file*

Description

The internal functions of the scapGNN package.

file format: 1. first index: pathway's name or ID. 2. second index: pathway's url or others, it doesn't matter. 3. third to all: gene symbols in pathway.

Usage

```
load_path_data(gmt_file_path)
```

Arguments

gmt_file_path Internal parameters.

Details

load_path_data

Value

a list

LTMG	<i>Left-truncated mixed Gaussian</i>
------	--------------------------------------

Description

Functional implementation of Left-truncated mixed Gaussian. The internal functions of the scapGNN package.

Usage

```
LTMG(VEC, Zcut_G, k = 5)
```

Arguments

VEC	Internal parameters.
Zcut_G	Internal parameters.
k	Internal parameters.

Details

LTMG

LTMG-class	<i>An S4 class to represent the input data for LTMG.</i>
------------	--

Description

An S4 class to represent the input data for LTMG.

Slots

InputData	Input data for LTMG.
OrdinalMatrix	LTMG output data.

norm1	<i>Function that computes the norm 1 of a numeric vector</i>
-------	--

Description

Function that computes the norm 1 of a numeric vector

Usage

```
norm1(x)
```

Arguments

x : numeric vector

Value

a single real value (the norm1 of the input vector)

plotCCNetwork	<i>Visualize cell cluster association network graph</i>
---------------	---

Description

The plotCCNetwork function takes cells belonging to the same phenotype as a cluster. When cell phenotypes are not provided, the plotCCNetwork functions identify cell clusters based on edge betweenness. Cell interactions between cell clusters are merged into one edge by mean. The thickness of the edge indicates the strength of interaction between cell clusters.

Usage

```
plotCCNetwork(  
  network.data,  
  cell_id = NULL,  
  cell_cluster = FALSE,  
  vertex.colors = NULL,  
  vertex.size = 10,  
  vertex.label.cex = 0.8,  
  vertex.label.dist = 1,  
  vertex.label.color = "black",  
  edge.width = 5,  
  margin = 0,  
  layout = layout_with_lgl,  
  legend.cex = 1.5,  
  legend.pt.cex = 3,  
  proportion = 1,  
  plotgraph = TRUE  
)
```

Arguments

<code>network.data</code>	The input network data is the result from the ConNetGNN function.
<code>cell_id</code>	A vector of cell phenotype.
<code>cell_cluster</code>	A binary value. Whether to automatically identify cell clusters based on edge betweenness. Default: FALSE.
<code>vertex.colors</code>	The fill color of the vertex. The number of colors should match the number of cell phenotypes. If NULL (default), the system will automatically assign colors.
<code>vertex.size</code>	The size of the vertex. Default: 10.
<code>vertex.label.cex</code>	The font size for vertex labels. Default: 0.8.
<code>vertex.label.dist</code>	The distance of the label from the center of the vertex. If it is 0 then the label is centered on the vertex. Default: 1.
<code>vertex.label.color</code>	The color of the labels. Default: black.
<code>edge.width</code>	The width of the edge. This does not affect the relative size of the edge weights. Default: 5.
<code>margin</code>	The amount of empty space below, over, at the left and right of the plot, it is a numeric vector of length four. Usually values between 0 and 0.5 are meaningful, but negative values are also possible, that will make the plot zoom in to a part of the graph. If it is shorter than four then it is recycled. Default: 0.
<code>layout</code>	Either a function or a numeric matrix. It specifies how the vertices will be placed on the plot. For details, please refer to the <code>igraphPackage</code> . Default: <code>layout_with_lgl</code> .
<code>legend.cex</code>	The font size of legend. Default: 1.5.
<code>legend.pt.cex</code>	Expansion factor(s) for the points. Default: 3.
<code>proportion</code>	This parameter specifies what percentage of edges to display (edges are sorted by their weight in descending order). Default: 1, all edges are used.
<code>plotgraph</code>	Whether to draw the picture. Default: TRUE. If FALSE, the image will not be displayed but the network data will be returned in the <code>igraph</code> data format.

Details

`plotCCNetwork`

Value

Graph or network data.

Examples

```
require(igraph)
require(graphics)
```



```

data(ConNetGNN_data)

# Construct the cell phenotype vector.
cell_id<-colnames(ConNetGNN_data[["cell_network"]])
temp<-unlist(strsplit(cell_id,"_"))
cell_phen<-temp[seq(2,length(temp)-1,by=3)]
names(cell_id)<-cell_phen
head(cell_id)
plotCCNetwork(ConNetGNN_data,cell_id,edge.width=10)

```

plotGANetwork	<i>Visualize gene association network graph of a gene module or pathway at the specified cell phenotype</i>
---------------	---

Description

Based on the gene set input by the user, plotGANetwork functional draws the gene association network in the specified cell phenotype. The node size in the network reflects the activation strength of the gene. The thickness of the edge indicates the strength of interaction between genes.

Usage

```

plotGANetwork(
  network.data,
  cellset,
  geneset,
  rwr.gamma = 0.7,
  vertex.colors = NULL,
  vertex.size = 10,
  vertex.label.cex = 0.8,
  vertex.label.dist = 1,
  vertex.label.color = "black",
  edge.width = 5,
  margin = 0,
  layout = layout_as_star,
  main = NULL,
  plotgraph = TRUE
)

```

Arguments

network.data	Network data constructed by the ConNetGNN function.
cellset	A vector of cell id. A cell set corresponding to the specified cell phenotype.
geneset	A vector of gene id. A gene module or pathway.
rwr.gamma	Restart parameter. Default: 0.7.
vertex.colors	he fill color of the vertex. The number of colors should match the number of cell phenotypes. If NULL (default), the system will automatically assign colors.

<code>vertex.size</code>	The size of the vertex. Default: 10.
<code>vertex.label.cex</code>	The font size for vertex labels. Default: 0.8.
<code>vertex.label.dist</code>	The distance of the label from the center of the vertex. If it is 0 then the label is centered on the vertex. Default: 1.
<code>vertex.label.color</code>	The color of the labels. Default: black.
<code>edge.width</code>	The width of the edge. This does not affect the relative size of the edge weights. Default: 5.
<code>margin</code>	The amount of empty space below, over, at the left and right of the plot, it is a numeric vector of length four. Usually values between 0 and 0.5 are meaningful, but negative values are also possible, that will make the plot zoom in to a part of the graph. If it is shorter than four then it is recycled. Default: 0.
<code>layout</code>	Either a function or a numeric matrix. It specifies how the vertices will be placed on the plot. For details, please refer to the <code>igraphPackage</code> . Default: <code>layout_as_star</code> .
<code>main</code>	A main title for the plot.
<code>plotgraph</code>	Whether to draw the picture. Default: TRUE. If FALSE, the image will not be displayed but the network data will be returned in the <code>igraph</code> data format.

Details

`plotGANetwork`

Value

A graph or list.

Examples

```
require(igraph)

# Load the result of the ConNetGNN function.
data(ConNetGNN_data)

data("Hv_exp")
index<-grep("0h",colnames(Hv_exp))
cellset<-colnames(Hv_exp)[index]
pathways<-load_path_data(system.file("extdata", "KEGG_human.gmt", package = "scapGNN"))
geneset<-pathways[[which(names(pathways)=="Tight junction [PATH:hsa04530]")]
plotGANetwork(ConNetGNN_data,cellset,geneset,main = "Tight junction [PATH:hsa04530]")
```

plotMulPhenGM	<i>Visualize gene association network graph for activated gene modules under multiple cell phenotypes</i>
---------------	---

Description

For multiple cell phenotypes, the plotMulPhenGM function will display the activated gene modules for each phenotype and show the connection and status of genes in different cell phenotypes.

Usage

```
plotMulPhenGM(
  data.list,
  network.data,
  vertex.colors = NULL,
  vertex.size = 10,
  vertex.label.cex = 0.8,
  vertex.label.dist = 1,
  vertex.label.color = "black",
  edge.width = 5,
  margin = 0,
  layout = layout_with_lgl,
  legend.position = "bottomright",
  legend.cex = 1.5,
  legend.pt.cex = 3,
  plotgraph = TRUE
)
```

Arguments

data.list	a list. Each element represents the cpGModule function result of a cell phenotype and the names of the lists are the corresponding cell phenotype.
network.data	Network data constructed by the ConNetGNN function.
vertex.colors	The fill color of the vertex. The number of colors should match the number of cell phenotypes. If NULL (default), the system will automatically assign colors.
vertex.size	The size of the vertex. Default: 10.
vertex.label.cex	The font size for vertex labels. Default: 0.8.
vertex.label.dist	The distance of the label from the center of the vertex. If it is 0 then the label is centered on the vertex. Default: 1.
vertex.label.color	The color of the labels. Default: black.
edge.width	The width of the edge. This does not affect the relative size of the edge weights. Default: 5.

margin	The amount of empty space below, over, at the left and right of the plot, it is a numeric vector of length four. Usually values between 0 and 0.5 are meaningful, but negative values are also possible, that will make the plot zoom in to a part of the graph. If it is shorter than four then it is recycled. Default: 0.
layout	Either a function or a numeric matrix. It specifies how the vertices will be placed on the plot. For details, please refer to the igraph Package. Default: layout_with_lgl.
legend.position	This places the legend on the inside of the plot frame at the given location. See the legend() function for details.
legend.cex	The font size of legend. Default: 1.5.
legend.pt.cex	Expansion factor(s) for the points. Default: 3.
plotgraph	Whether to draw the picture. Default: TRUE. If FALSE, the image will not be displayed but the network data will be returned in the igraph data format.

Details

plotMulPhenGM

If a gene is significantly activated in more than one cell phenotype, we call it a co-activated gene. These co-activated genes are shown on the sector diagram. Each interval of the sector diagram represents the activation strength of the gene in this cell phenotype relative to other cell phenotypes.

Value

A graph or list.

Examples

```
require(igraph)
require(grDevices)
# Load the result of the ConNetGNN function.
data(ConNetGNN_data)
# Obtain cpGModule results for each cell phenotype.
data(H9_0h_cpGM_data)
data(H9_24h_cpGM_data)
data(H9_36h_cpGM_data)
data.list<-list(H9_0h=H9_0h_cpGM_data,H9_24h=H9_24h_cpGM_data,H9_36h=H9_36h_cpGM_data)
plotMulPhenGM(data.list,ConNetGNN_data)
```

Description

This function is to prepare data for the ConNetGNN function.

Usage

```
Preprocessing(data, verbose = TRUE)
```

Arguments

<code>data</code>	The input data should be a data frame or a matrix where the rows are genes and the columns are cells. The <code>seurat</code> object are also accepted.
<code>verbose</code>	Gives information about each step. Default: TRUE.

Details**Preprocessing**

The function is able to interface with the `seurat` framework. The process of `seurat` data processing refers to `Examples`. The input data should be containing hypervariable genes and log-transformed. Left-truncated mixed Gaussian (LTMG) modeling to calculate gene regulatory signal matrix. Positively correlated gene-gene and cell-cell are used as the initial gene correlation matrix and cell correlation matrix.

Value

A list:

orig_dara User-submitted raw data, rows are highly variable genes and columns are cells.

cell_features Cell feature matrix.

gene_features Gene feature matrix.

ltmg_matrix Gene regulatory signal matrix for LTMG.

cell_adj The adjacency matrix of the cell correlation network.

gene_adj The adjacency matrix of the gene correlation network.

Examples

```
## Not run:
# Load dependent packages.
require(coop)

# Seurat data processing.
require(Seurat)

# Load the PBMC dataset (Case data for seurat)
pbmc.data <- Read10X(data.dir = "../data/pbmc3k/filtered_gene_bc_matrices/hg19/")

# Our recommended data filtering is that only genes expressed as non-zero in more than
# 1% of cells, and cells expressed as non-zero in more than 1% of genes are kept.
# In addition, users can also filter mitochondrial genes according to their own needs.
pbmc <- CreateSeuratObject(counts = pbmc.data, project = "case",
                           min.cells = 3, min.features = 200)
pbmc[["percent.mt"]] <- PercentageFeatureSet(pbmc, pattern = "^MT-")
pbmc <- subset(pbmc, subset = nFeature_RNA > 200 & nFeature_RNA < 2500 & percent.mt < 5)
```

```

# Normalizing the data.
pbmc <- NormalizeData(pbmc, normalization.method = "LogNormalize")

# Identification of highly variable features.
pbmc <- FindVariableFeatures(pbmc, selection.method = 'vst', nfeatures = 2000)

# Run Preprocessing.
Prep_data <- Preprocessing(pbmc)

## End(Not run)

# Users can also directly input data
# in data frame or matrix format
# containing highly variable genes.
data("Hv_exp")
Prep_data <- Preprocessing(Hv_exp[1:300,])

```

Pure_CDF

Pure_CDF

Description

The internal functions of the scapGNN package.

Usage

Pure_CDF(Vec)

Arguments

Vec Internal parameters.

Details

Pure_CDF

RNA_ATAC_IntNet

Results of InteNet() for integrating scRNA-seq and scATAC-seq data.

Description

An integrated network of scRNA-seq and scATAC-seq data from SNARE-seq.

Usage

RNA_ATAC_IntNet

Format

a list of three adjacency matrices.

Examples

```
data(RNA_ATAC_IntNet)
```

RNA_net

Results of ConNetGNN() for scRNA-seq data from SNARE-seq dataset

Description

A list to store the gene association network of scRNA-seq data. Case data from the SNARE-seq dataset.

Usage

```
RNA_net
```

Format

a list of three adjacency matrices.

Examples

```
data(RNA_net)
```

RunLTMG

Run Left-truncated mixed Gaussian

Description

Functional implementation of Left-truncated mixed Gaussian. The internal functions of the scapGNN package.

Usage

```
.RunLTMG(object, Gene_use = NULL, k = 5, verbose, seed = 123)
```

```
RunLTMG(object, Gene_use = NULL, k = 5, verbose, seed = 123)
```

```
## S4 method for signature 'LTMG'
```

```
RunLTMG(object, Gene_use = NULL, k = 5, verbose, seed = 123)
```

Arguments

object	A LTMG object
Gene_use	using X numebr of top variant gene. input a number, recommend 2000.
k	Constant, defaults 5.
verbose	Gives information about each step.
seed	Random number generator seed.

Details

RunLTMG

For more information, please refer to: DOI: 10.1093/nar/gkz655 and <https://github.com/zy26/LTMGSCA>.

Value

A list contains raw input data and LTMG results.

RWR	<i>Function that performs a random Walk with restart (RWR) on a given graph</i>
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Description

Function that performs a random Walk with restart (RWR) on a given graph

Usage

```
RWR(W, ind.positives, gamma = 0.6, tmax = 1000, eps = 1e-10, norm = TRUE)
```

Arguments

W	: adjacency matrix of the graph
ind.positives	: indices of the "core" positive examples of the graph. They represent to the indices of W corresponding to the positive examples
gamma	: restart parammer (def: 0.6)
tmax	: maximum number of iterations (def: 1000)
eps	: maximum allowed difference between the computed probabilities at the steady state
norm	: if TRUE (def) the adjacency matrix W of the graph is normalized to $M = D^{-1} * W$, otherwise it is assumed that the matrix W is just normalized

Value

a list with three elements: - p : the probability at the steady state - ind.positives : indices of the "core" positive examples of the graph (it is equal to the same input parameter - n.iter : number of performed iterations

`scPathway`*Infer pathway activation score matrix at single-cell resolution*

Description

Calculate pathway activity score of single-cell by random walk with restart (RWR).

Usage

```
scPathway(  
  network.data,  
  gmt.path = NULL,  
  pathway.min = 10,  
  pathway.max = 500,  
  nperm = 100,  
  parallel.cores = 2,  
  rwr.gamma = 0.7,  
  verbose = TRUE  
)
```

Arguments

<code>network.data</code>	The input network data is the result from the ConNetGNN function.
<code>gmt.path</code>	Pathway database in GMT format.
<code>pathway.min</code>	Minimum size (in genes) for pathway to be considered. Default: 10.
<code>pathway.max</code>	Maximum size (in genes) for database gene sets to be considered. Default: 500.
<code>nperm</code>	Number of random permutations. Default: 100.
<code>parallel.cores</code>	Number of processors to use when doing the calculations in parallel (default: 2). If <code>parallel.cores=0</code> , then it will use all available core processors unless we set this argument with a smaller number.
<code>rwr.gamma</code>	Restart parameter. Default: 0.7.
<code>verbose</code>	Gives information about each step. Default: TRUE.

Details

`scPathway`

The `scPathway` function integrates the results of ConNetGNN into a gene-cell association network. The genes included in each pathway are used as a restart set in the gene-cell association network to calculate the strength of its association with each cell through RWR. Perturbation analysis was performed to remove noise effects in the network and to obtain the final single-cell pathway activity score matrix.

Value

A matrix of single-cell pathway activity score.

Examples

```
require(parallel)
require(utils)
# Load the result of the ConNetGNN function.
data(ConNetGNN_data)
kegg.path<-system.file("extdata", "KEGG_human.gmt", package = "scapGNN")
scPathway_data<-scPathway(ConNetGNN_data,gmt.path=kegg.path,
                          pathway.min=25,nperm=6,parallel.cores=1)
```

scPathway_data	<i>Single cell pathway activity matrix</i>
----------------	--

Description

Results of scPathway() function.

Usage

```
scPathway_data
```

Format

a matrix.

Examples

```
data(scPathway_data)
```

Index

* datasets

- ATAC_net, [2](#)
- ConNetGNN_data, [6](#)
- H9_0h_cpGM_data, [9](#)
- H9_24h_cpGM_data, [10](#)
- H9_36h_cpGM_data, [10](#)
- Hv_exp, [11](#)
- RNA_ATAC_IntNet, [22](#)
- RNA_net, [23](#)
- scPathway_data, [26](#)
- .RunLTMG (RunLTMG), [23](#)

ATAC_net, [2](#)

BIC_LTMG, [3](#)
BIC_ZIMG, [3](#)

ConNetGNN, [4](#)
ConNetGNN_data, [6](#)
cpGModule, [6](#)
create_scapGNN_env, [8](#)

Fit_LTMG, [8](#)

Global_Zcut, [9](#)

H9_0h_cpGM_data, [9](#)
H9_24h_cpGM_data, [10](#)
H9_36h_cpGM_data, [10](#)
Hv_exp, [11](#)

instPyModule, [11](#)
InteNet, [12](#)
isLoading, [13](#)

load_path_data, [13](#)
LTMG, [14](#)
LTMG-class, [14](#)

norm1, [15](#)

plotCCNetwork, [15](#)

plotGANetwork, [17](#)
plotMulPhenGM, [19](#)
Preprocessing, [20](#)
Pure_CDF, [22](#)

RNA_ATAC_IntNet, [22](#)
RNA_net, [23](#)
RunLTMG, [23](#)
RunLTMG, LTMG-method (RunLTMG), [23](#)
RWR, [24](#)

scPathway, [25](#)
scPathway_data, [26](#)