Package: SparseMDC (via r-universe)

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Type Package

Title Implementation of SparseMDC Algorithm

Version 0.99.5

Description Implements the algorithm described in Barron, M., and Li, J. (Not yet published). This algorithm clusters samples from multiple ordered populations, links the clusters across the conditions and identifies marker genes for these changes. The package was designed for scRNA-Seq data but is also applicable to many other data types, just replace cells with samples and genes with variables. The package also contains functions for estimating the parameters for SparseMDC as outlined in the paper. We recommend that users further select their marker genes using the magnitude of the cluster centers.

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cell_type_biase

Biase Data Cell Type

Description

The cell type of each of the cells in the Biase data.

Usage

cell_type_biase

Format

An R.Data object containing a vector with the cell type of each of the cells in the Biase Data.

condition_biase 3

condition_biase

Biase Data Conditions

Description

The condition for each sample in the Biase data. To be used when splitting the data to demonstrate SparseDC.

Usage

condition_biase

Format

An R.Data object containing a vector with the condition of the 49 cells in the Biase data.

data_biase

Biase Data

Description

This dataset was created by Biase et al. to study cell fat inclination in mouse embryos. It contains FPKM gene expression measurements for 49 cells and 16,514 genes. There are three cell types in the dataset, zygote, two-cell embryo, and four-cell embryo cells.

Usage

data_biase

Format

An R.Data object storing FPKM gene expression measurements for each of the samples.

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generate_	uni	dat
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Uniform data generator For use with the gap statistic. Generates datasets drawn from the reference distribution where each reference feature is generated uniformly over the range of observed values for that feature.

Description

Uniform data generator For use with the gap statistic. Generates datasets drawn from the reference distribution where each reference feature is generated uniformly over the range of observed values for that feature.

Usage

```
generate_uni_dat(data)
```

Arguments

data

A dataset with rows as features and columns as samples.

Value

A dataset drawn from the reference distribution for use internally with the gap statistic.

lambda1_calculator

Lambda 1 Calcualtor

Description

Calculates the lambda 1 value for the SparseMDC algorithm. The lambda 1 value controls the number of marker genes selected for each cluster in the output from SparseMDC. It is calculated as the value of lambda 1 that results in no marker genes being selected when then are no meaningful clusters present in the data. Please see the original manuscript for further details.

```
lambda1_calculator(dat_l, dim, nclust, nboot = 1000, alpha1 = 0.5,
  delta = 1e-07)
```

lambda2_calculator 5

Arguments

dat_l	list with D entries, each entry contains data d , $p * n$ matrix. This data should be centered and log-transformed.
dim	Total number of conditions, D.
nclust	Total number of clusters.
nboot	The number of bootstrap repetitions used for estimating lambda 1, the default value is 1000.
alpha1	The quantile of the bootstrapped lambda 1 values to use, range is (0,1). The default value is 0.5, the median of the calculated lambda 1 values.
delta	Small value term added to ensure existance, default value is 0.0000001.

Value

The estimated value of lambda1 for use in main SparseMDC algorithm

Examples

```
set.seed(10)
# Select small dataset for example
data_test <- data_biase[1:100,]
# Split data into condition A and B
data_A <- data_test[ , which(condition_biase == "A")]
data_B <- data_test[ , which(condition_biase == "B")]
data_C <- data_test[ , which(condition_biase == "C")]
# Store data as list
dat_l <- list(data_A, data_B, data_C)
# Pre-process the data
pdat <- pre_proc_data(dat_l, dim=3, norm = FALSE, log = TRUE, center = TRUE)
lambda1 <- lambda1_calculator(pdat, dim = 3, nclust = 3 )</pre>
```

lambda2_calculator

Lambda 2 Calculator

Description

Calculates the lambda 2 values for use in the main SparseMDC algorithm, the lambda 2 value controls the number of genes that show condition-dependent expression within each cell type. That is it controls the number of different mean values across the conditions for each cluster. It is calculated by estimating the value of lambda 2 that would result in no difference in mean values across conditions when there are no meaningful differences across between the conditions. For further details please see the original manuscript.

```
lambda2_calculator(dat_1, dim, nclust, nboot = 1000, alpha2 = 0.5,
  delta = 1e-07, lambda1)
```

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Arguments

dat_l	list with D entries, each entry contains data d, p * n matrix. This data should be centered and log-transformed.
dim	Total number of conditions, D.
nclust	Total number of clusters.
nboot	The number of bootstrap repetitions for estimating lambda 2, the default value is 1000.
alpha2	The quantile of the bootstrapped lambda 2 values to use, range is (0,1). The default value is 0.5, the median of the calculated lambda 2 values.
delta	Small term to ensure existance of solution, default is 0.0000001.
lambda1	Calcualted penalty parameter for mean size.

Value

The estimated value of lambda2

Examples

```
set.seed(10)
# Select small dataset for example
data_test <- data_biase[1:100,]
# Split data into condition A and B
data_A <- data_test[ , which(condition_biase == "A")]
data_B <- data_test[ , which(condition_biase == "B")]
data_C <- data_test[ , which(condition_biase == "C")]
# Store data as list
dat_l <- list(data_A, data_B, data_C)
# Pre-process the data
pdat <- pre_proc_data(dat_l, dim=3, norm = FALSE, log = TRUE,
center = TRUE)
lambda1 <- lambda1_calculator(pdat, dim = 3, nclust = 3)
lambda2 <- lambda2_calculator(pdat, dim = 3, nclust = 3, lambda1 = lambda1)</pre>
```

mu_calc

mu Calculator

Description

This function handles the calculations of the Mu Solver. This function runs inside sparse_mdc.

```
mu_calc(d, v, EQ, dim, x, nk, p1, p2, delta)
```

mu_solver 7

Arguments

d	- The current condition.
V	- Relationship matrix, describing relationship between d-1 and d.
EQ	- Equality matrix specifying the number of following terms to which mu_d is equal.
dim	Total number of conditions, D.
Х	- Matrix of mean values.
nk	- Vector with the number of samples in each dimension for this cluster.
p1	- Penalties on mean size.
p2	- Penalties on mean difference.
delta	Small term to ensure existance of solution.

Value

A list containing two vectors containing the calculated values of $mu_d \mid mu_d < mu_{d-1}$ and $mu_d \mid mu_d > mu_{d-1}$ respectively.

mu_solver Mu Solver

Description

Calculates the regularized center values for a cluster. This function runs inside sparse_mdc.

Usage

```
mu_solver(d, mu, v, EQ, dim, x, nk, p1, p2, delta)
```

Arguments

d	- The current condition.
mu	- A matrix of regularized mean values.
V	- Relationship matrix, describing relationship between d-1 and d.
EQ	- Equality matrix specifying the number of following terms to which mu_d is equal.
dim	- Total dimensions.
Х	- Matrix of mean values.
nk	- Vector with the number of samples in each condition for this cluster.
p1	- Penalties on mean size.
p2	- Penalties on mean difference.
delta	Small term to ensure existance of solution.

Value

A matrix containing the regularized cluster means for each dimension.

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y calculator	
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Description

Calcualtes the penalty terms for penalizing mean size and mean difference. This function runs inside sparse_mdc.

Usage

```
pen_calculator(lambda1, lambda2, nk, delta)
```

Arguments

lambda1 Calculated penalty parameter for mean size.

lambda2 Calculated penalty parameter for mean difference.

nk Vector containing the number of samples in each dimension.

delta Small term to ensure existance of solution, default is 0.0000001.

Value

a list with two vectors containing the penalty terms for mean size and mean difference respectively.

Description

This function centers on a gene-by-gene basis, normalizes and/or log transforms the data prior to the application of SparseMDC. For the sequencing depth normalization we recommend that users use one of the many methods developed for normalizing scRNA-Seq data prior to using SparseMDC and so can set norm = FALSE. However, here we normalize the data by dividing by the total number of reads. This function log transforms the data by applying log(x + 1) to each of the data sets. By far the most important pre-processing step for SparseMDC is the centralization of the data. Having centralized data is a core component of the SparseMDC algorithm and is necessary for both accurate clustering of the cells and identifying marker genes. We therefore recommend that all users centralize their data using this function and that only experienced users set center = FALSE.

```
pre_proc_data(dat_1, dim, norm = FALSE, log = TRUE, center = TRUE)
```

score_calc 9

Arguments

dat_l	list with D entries, each entry contains data d, $p * n$ matrix. The entries should be ordered according the condition of each dataset. The rows of the data matrix should contain samples while the columns contain features or genes.
dim	Total number of conditions, D.
norm	True/False on if the data should be normalized. This parameter controls whether the data is normalized for sequencing depth by dividing each column by the total number of reads for that sample. We recommend that user use one of the many methods for normalizing scRNA-Seq data and so set this as FALSE. The default value is FALSE
log	True/False of if the data should be transformed as $log(x + 1)$. The default value is TRUE.
center	This parameter controls whether the data is centered on a gene by gene basis. We recommend all users center their data prior to applying SparseMDC and only experienced users should set this as FALSE. The default value is TRUE.

Value

A list with D entries containing the pre-processed data.

Examples

```
set.seed(10)
# Select small dataset for example
data_test <- data_biase[1:100,]
# Split data into condition A and B
data_A <- data_test[ , which(condition_biase == "A")]
data_B <- data_test[ , which(condition_biase == "B")]
data_C <- data_test[ , which(condition_biase == "C")]
# Store data as list
dat_l <- list(data_A, data_B, data_C)
# Pre-process the data
pdat <- pre_proc_data(dat_l, dim=3, norm = FALSE, log = TRUE, center = TRUE)</pre>
```

score_calc

Score calculator

Description

Calculates the score for each combination of cluster assignments and center values. This function runs inside sparse_mdc.

```
score_calc(pdat, clusters, mu, lambda1, lambda2, nclust, delta, dim)
```

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Arguments

pdat list with D entries, each entry contains data d, p * n matrix. This data should be

centered and log-transformed.

clusters List containing cluster assignments for each dimension as entries.

mu list with D entries, each entry contains centers for data d, p*k matrix.

lambda1 Penalty parameter for mean size.

lambda2 Penalty parameter for mean difference.

nclust Number of clusters in the data.

delta Small term to ensure existance of solution, default is 0.0000001.

dim Total number of conditions, D.

Value

The caluculated score.

sdc_mpar SparseDC Multi Parallel

Description

Applies sparse clustering to data from multiple conditions, linking the clusters across conditions and selecting a set of marker variables for each cluster and condition. This is a wrapper function to run SparseMDC in parallel and choose the solution with the mimimum score for each run.

Usage

```
sdc_mpar(pdat, nclust, dim, lambda1, lambda2, nitter = 20,
  nstarts = 50, init_iter = 5, delta = 1e-07, par_starts, cores)
```

Arguments

pdat list with D entries, each entry contains data d, p * n matrix. This data should be

centered and log-transformed.

nclust Number of clusters in the data.
dim Total number of conditions, D.

lambda1 The lambda 1 value to use in the SparseMDC function. This value controls the

number of marker genes detected for each of the clusters in the final result. This can be calculated using the "lambda1_calculator" function or supplied by the

user.

1ambda2 The lambda 2 value to use in the SparseMDC function. This value controls

the number of genes that show condition-dependent expression within each cell type. This can be calculated using the "lambda2_calculator" function or supplied

by the user.

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nitter	The max number of iterations for each of the start values, the default value is 20.
nstarts	The max number of possible starts. The default value is 50.
init_iter	The number of iterations used to initialize the algorithm. Higher values result in less starts but more accurate and vice versa. Default is 5.
delta	Small term to ensure existance of solution, default is 0.0000001.
par_starts	Number of parallel starts.
cores	Number of cores to use.

Value

A list containing cluster assignments, center values and the scores for each start.

Examples

```
set.seed(10)
# Select small dataset for example
data_test <- data_biase[1:100,]</pre>
# Split data into condition A and B
data_A <- data_test[ , which(condition_biase == "A")]</pre>
data_B <- data_test[ , which(condition_biase == "B")]</pre>
data_C <- data_test[ , which(condition_biase == "C")]</pre>
# Store data as list
dat_1 <- list(data_A, data_B, data_C)</pre>
# Pre-process the data
pdat <- pre_proc_data(dat_1, dim=3, norm = FALSE, log = TRUE,</pre>
center = TRUE)
# Calculate lambda1
lambda1 <- lambda1_calculator(pdat, dim = 3, nclust = 3)</pre>
# Calcualte lambda2
lambda2 <- lambda2_calculator(pdat, dim = 3, nclust = 3, lambda1 = lambda1)</pre>
# Prepare parallel enviornment
library(doParallel) # Load package
library(foreach) # Load the package
library(doRNG)
# Apply SparseMDC
smdc_res <- sdc_mpar(pdat, nclust = 3, dim = 3, lambda1 = lambda1,</pre>
lambda2 = lambda2, par_starts = 2, cores = 2)
```

sparsemdc_gap

Gap Statistic Calculator

Description

This function calculates the gap statistic for SparseMDC. For use when the number of clusters in the data is unknown. We recommend using alternate methods to infer the number of clusters in the data.

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Usage

```
sparsemdc_gap(pdat, dim, min_clus, max_clus, nboots = 200, nitter = 20,
  nstarts = 10, l1_boot = 50, l2_boot = 50)
```

Arguments

pdat	list with D entries, each entry contains data d, p * n matrix. This data should be centered and log-transformed.
dim	Total number of conditions, D.
min_clus	The minimum number of clusters to try, minimum value is 2.
max_clus	The maximum number of clusters to try.
nboots	The number of bootstrap repetitions to use, default = 200 .
nitter	The max number of iterations for each of the start values, the default value is 20.
nstarts	The number of start values to use for SparseDC. The default value is 10.
l1_boot	The number of bootstrap repetitions used for estimating lambda 1.
12_boot	The number of bootstrap repetitions used for estimating lambda 2.

Value

A list containing the optimal number of clusters, as well as gap statistics and the calculated standard error for each number of clusters.

Examples

```
set.seed(10)
# Select small dataset for example
data_test <- data_biase[1:100,]
# Split data into condition A and B
data_A <- data_test[ , which(condition_biase == "A")]
data_B <- data_test[ , which(condition_biase == "B")]
data_C <- data_test[ , which(condition_biase == "C")]
# Store data as list
dat_l <- list(data_A, data_B, data_C)
# Pre-process the data
pdat <- pre_proc_data(dat_l, dim=3, norm = FALSE, log = TRUE,
center = TRUE)
# Run with one bootstrap sample for example
gap_stat <- sparsemdc_gap(pdat, dim=3, min_clus = 2, max_clus =3, nboots =2,
nitter = 2, nstarts = 1, l1_boot = 5, l2_boot = 5)</pre>
```

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sparse_mdc	SparseDC Multi		
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Description

Applies sparse clustering to data from multiple conditions, linking the clusters across conditions and selecting a set of marker variables for each cluster and condition. See the manuscript for descriptions of the different categories of marker genes.

Usage

```
sparse_mdc(pdat, nclust, dim, lambda1, lambda2, nitter = 20,
  nstarts = 50, init_iter = 5, delta = 1e-07)
```

Arguments

pdat	list with D entries, each entry contains data d, p \ast n matrix. This data should be centered and log-transformed.
nclust	Number of clusters in the data.
dim	Total number of conditions, D.
lambda1	The lambda 1 value to use in the SparseMDC function. This value controls the number of marker genes detected for each of the clusters in the final result. This can be calculated using the "lambda1_calculator" function or supplied by the user.
lambda2	The lambda 2 value to use in the SparseMDC function. This value controls the number of genes that show condition-dependent expression within each cell type. This can be calculated using the "lambda2_calculator" function or supplied by the user.
nitter	The max number of iterations for each of the start values, the default value is 20.
nstarts	The max number of possible starts. The default value is 50.
init_iter	The number of iterations used to initialize the algorithm. Higher values result in less starts but more accurate and vice versa. Default is 5.
delta	Small term to ensure existance of solution, default is 0.0000001.

Value

A list containing cluster assignments, center values and the scores for each start.

Examples

```
set.seed(10)
# Select small dataset for example
data_test <- data_biase[1:100,]
# Split data into condition A and B
data_A <- data_test[ , which(condition_biase == "A")]</pre>
```

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```
data_B <- data_test[ , which(condition_biase == "B")]
data_C <- data_test[ , which(condition_biase == "C")]
# Store data as list
dat_l <- list(data_A, data_B, data_C)
# Pre-process the data
pdat <- pre_proc_data(dat_l, dim=3, norm = FALSE, log = TRUE,
center = TRUE)
lambda1 <- lambda1_calculator(pdat, dim = 3, nclust = 3)
lambda2 <- lambda2_calculator(pdat, dim = 3, nclust = 3, lambda1 = lambda1)
smdc_res <- sparse_mdc(pdat, nclust = 3, dim = 3, lambda1 = lambda1,
lambda2 = lambda2)</pre>
```

 S_func

The soft thresholding operator

Description

Function to solve the soft thresholding problem

Usage

```
S_func(x, a)
```

Arguments

x The data value

a The lambda value

Value

The solution to the soft thresholding operator.

update_c

update_c

Description

Assigns samples to clusters. This function runs inside sparse_mdc.

```
update_c(mu, pdat, nclust, dim, lambda1, lambda2, delta)
```

update_mu 15

Arguments

mu	list with D entries, each entry contains centers for data d, p*k matrix.
pdat	list with D entries, each entry contains data d, p \ast n matrix. This data should be centered and log-transformed.
_	

nclust Total number of clusters.

dim Total number of conditions, D.

lambda1 Calculated penalty parameter for mean size.
 lambda2 Calculated penalty parameter for mean difference.

delta Small term to ensure existance of solution.

Value

A list with D entries containing cluster assignments for each sample.

update_mu	update_mu		
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Description

Update the mean/center values for each cluster and dimension. This function runs inside sparse_mdc.

Usage

```
update_mu(clusters, pdat, nclust, dim, lambda1, lambda2, ngenes, delta)
```

Arguments

clusters	List containing cluster assignments for each dimension as entries.
pdat	list with D entries, each entry contains data d, p \ast n matrix. This data should be centered and log-transformed.
nclust	Number of clusters in the data.
dim	Total number of conditions, D.
lambda1	Calculated penalty parameter for mean size.
lambda2	Calculated penalty parameter for mean difference.
ngenes	The number of genes in the data.
delta	Small term to ensure existance of solution.

Value

A list containing the center values for the clusters in each dimensions as entries.

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