Package 'DCATS'

November 27, 2024

Title Differential Composition Analysis Transformed by a Similarity

Type Package

```
matrix
Version 1.5.0
Description Methods to detect the differential composition abundances between
      conditions in singel-cell RNA-seq experiments, with or without replicates.
      It aims to correct bias introduced by missclaisification and enable
      controlling of confounding covariates. To avoid the influence of proportion
      change from big cell types, DCATS can use either total cell number or
      specific reference group as normalization term.
Depends R (>= 4.1.0), stats
License MIT + file LICENSE
Imports MCMCpack, matrixStats, robustbase, aod, e1071
Suggests testthat (>= 3.0.0), knitr, Seurat, SeuratObject, tidyverse,
      rmarkdown, BiocStyle
VignetteBuilder knitr
RoxygenNote 7.2.3
biocViews SingleCell, Normalization
Encoding UTF-8
Config/testthat/edition 3
git_url https://git.bioconductor.org/packages/DCATS
git_branch devel
git_last_commit 61682fd
git_last_commit_date 2024-10-29
Repository Bioconductor 3.21
Date/Publication 2024-11-27
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```

create_simMat

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Description

Create a similarity matrix assuming the misclassification error distribute uniformly in all clusters

Usage

```
create_simMat(K, confuse_rate)
```

Arguments

K A integer for number of clusterconfuse_rate A float for confusion rate, uniformly to none-self clusters

Value

a similarity matrix with uniform confusion with other cluster

```
create_simMat(4, 0.1)
```

dcats_GLM 3

dcats_GLM	A Generalised linear model based likelihood ratio testing

Description

GLM supports both beta-binomial and negative binomial from aod package.

Usage

```
dcats_GLM(
  count_mat,
  design_mat,
  similarity_mat = NULL,
  pseudo_count = NULL,
  base_model = "NULL",
  fix_phi = NULL,
  reference = NULL
)
```

Arguments

count_mat	A matrix of composition sizes (n_sample, n_cluster) for each cluster in each sample
design_mat	A matrix or a data frame of testing candidate factors (n_sample, n_factor) with same sample order as count_mat. All factors should be continous and categorical with only two levels.
similarity_mat	A matrix of floats (n_cluster, n_cluster) for the similarity matrix between cluster group pair. The order of cluster should be consistent with those in 'count_mat'.
pseudo_count	A pseudo count to add for counts in all cell types Default NULL means 0 except if a cell type is empty in one condition, otherwise pseudo_count will be: 0.01 * rowMeans for each condition
base_model	A string value: 'NULL' for 1 factor vs NULL factor testing; 'FULL' for FULL factors vs $n\text{-}1$ factors testing.
fix_phi	A numeric used to provided a fixed phi value for the GLM for all cell types
reference	A vector of characters indicating which cell types are used as reference for normalization. 'NULL' indicates using total count for normalization.

Value

a list of significance p values for each cluster

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Examples

```
K <- 3
totals1 = c(100, 800, 1300, 600)
totals2 = c(250, 700, 1100)
diri_s1 = rep(1, K) * 20
diri_s2 = rep(1, K) * 20
simil_mat = DCATS::create_simMat(K, confuse_rate=0.2)
sim_dat <- DCATS::simulator_base(totals1, totals2, diri_s1, diri_s2, simil_mat)</pre>
sim_count = rbind(sim_dat\numb_cond1, sim_dat\numb_cond2)
sim_design = data.frame(condition = c("g1", "g1", "g1", "g1", "g2", "g2", "g2"),
gender = sample(c("Female", "Male"), 7, replace = TRUE))
## Using 1 factor vs NULL factor testing
dcats_GLM(sim_count, sim_design, similarity_mat = simil_mat)
## Using full factors vs n-1 factors testing with intercept term
dcats_GLM(sim_count, sim_design, similarity_mat = simil_mat, base_model='FULL')
## Fix phi
dcats_GLM(sim_count, sim_design, similarity_mat = simil_mat, fix_phi = 1/61)
## Specify reference cell type
colnames(sim_count) <- c("celltypeA", "celltypeB", "celltypeC")</pre>
```

detect_reference

Calculate a global phi for all cell types

Description

Assuming all cell types share the same phi. This global phi can be calculate by pooling all cell types together to fit a beta binomial distribution.

Usage

```
detect_reference(count_mat, design_mat, similarity_mat = NULL, fix_phi = NULL)
```

Arguments

count_mat	A matrix of composition sizes (n_sample, n_cluster) for each cluster in each sample.
design_mat	A matrix or a data frame of testing candidate factors (n_sample, n_factor) with same sample order as count_mat. All factors should be continous and categorical with only two levels.
~	A matrix of floats (n_cluster, n_cluster) for the similarity matrix between cluster group pair. The order of cluster should be consistent with those in 'count_mat'.
fix_phi	A numeric used to provided a fixed phi value for the GLM for all cell types.

Value

A data frame with ordered cell types and their p-value. Cell types are ordered by their p-values. The order indicating how they are recommended to be selected as reference cell types.

getPhi 5

Examples

```
K <- 3
totals1 = c(100, 800, 1300, 600)
totals2 = c(250, 700, 1100)
diri_s1 = rep(1, K) * 20
diri_s2 = rep(1, K) * 20
simil_mat = DCATS::create_simMat(K, confuse_rate=0.2)
sim_dat <- DCATS::simulator_base(totals1, totals2, diri_s1, diri_s2, simil_mat)
sim_count = rbind(sim_dat$numb_cond1, sim_dat$numb_cond2)
sim_design = data.frame(condition = c("g1", "g1", "g1", "g1", "g2", "g2", "g2"),
gender = sample(c("Female", "Male"), 7, replace = TRUE))
## Using 1 factor vs NULL factor testing
detect_reference(sim_count, sim_design)</pre>
```

getPhi

Calculate a global phi for all cell types

Description

Assuming all cell types share the same phi. This global phi can be calculate by pooling all cell types together to fit a beta binomial distribution.

Usage

```
getPhi(count_mat, design_mat)
```

Arguments

count_mat A matrix of composition sizes (n_sample, n_cluster) for each cluster in each

sample

design_mat A matrix of testing candidate factors (n_sample, n_factor) with same sample

order as count_mat

Value

A number indicating a global phi for all cell types

```
K <- 3
totals1 = c(100, 800, 1300, 600)
totals2 = c(250, 700, 1100)
diri_s1 = rep(1, K) * 20
diri_s2 = rep(1, K) * 20
simil_mat = DCATS::create_simMat(K, confuse_rate=0.2)
sim_dat <- DCATS::simulator_base(totals1, totals2, diri_s1, diri_s2, simil_mat)
sim_count = rbind(sim_dat$numb_cond1, sim_dat$numb_cond2)
sim_design = data.frame(condition = c("g1", "g1", "g1", "g1", "g2", "g2", "g2"))
phi = DCATS::getPhi(sim_count, sim_design)</pre>
```

6 Haber2017

Haber2017	Count matrices of intestinal epithelial scRNA-seq data from three con-
	ditions

Description

A data containing the count matrices, the similarity matrix and other variables used to generate the similarity matrix from intestinal epithelial single cell RNA sequencing data with three condition. Count matrices are calculated based on the number of cells in each cell type. The similarity matrix is calculated by support vector machine classifiers using 5-fold cross validation. Top 30 PCs are used as predictors.

Usage

Haber2017

Format

A list with 7 items:

```
count_ctrl the count matrix for the control group
count_Hpoly3 the count matrix for three days after H.polygyrus infection
count_Hpoly10 the count matrix for ten days after H.polygyrus infection
count_Salma the count matrix for two days after Salmonella infection
svm_mat the similarity matrix
source the source of this dataset
```

Source

```
https://www.nature.com/articles/nature24489
```

```
library(DCATS)
data(Haber2017)
```

Kang2017 7

Kang2017	Count matrices of 8 pooled lupus patient samples within two conditions

Description

A data containing the count matrices, the similarity matrix and other variables used to generate the similarity matrix from single cell RNA sequencing data of 8 pooled lupus patient samples within two conditions. Count matrices are calculated based on the number of cells in each cell type. The similarity matrix is calculated by support vector machine classifiers using 5-fold cross validation. Top 30 PCs are used as predictors. The symDF contains 30 PCs generated by standard Seurat pipeline and the condition, cell type information collected from the original paper.

Usage

Kang2017

Format

A list with 5 items:

count_ctrl the count matrix for three days after H.polygyrus infection
count_stim the count matrix for ten days after H.polygyrus infection
svm_mat the simularity matrix
svmDF the data frame used to calculate the similarity matrix.
source the source of this dataset

Source

https://www.nature.com/articles/nbt.4042

knn_simMat	Calculate stochastic transition matrix between clusters from a KNN connection matrix

Description

The transition probability from cluster i to j is the fraction of neighbours of all samples in cluster i that belongs to cluster j. Note, this matrix is asymmetric, so as the input KNN connection matrix.

Usage

```
knn_simMat(KNN_matrix, clusters)
```

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Arguments

KNN_matrix a sparse binary matrix with size (n_sample, n_sample). x_ij=1 means sample

j is a neighbour of sample i. As definition, we expect sum(KNN_matrix) =

n_sample * K, where K is the number neighbours.

clusters a (n_sample,) vector of cluster id for each sample.

Value

a similarity matrix calculated based on the knn graph.

Examples

```
data(simulation)
knn_mat = knn_simMat(simulation$knnGraphs, simulation$labels)
```

multinom_EM

An EM algorithm to fit a multinomial with maximum likelihood

Description

An EM algorithm to fit a multinomial with maximum likelihood

Usage

```
multinom_EM(X, simMM, min_iter = 10, max_iter = 1000, logLik_threshold = 0.01)
```

Arguments

X A vector of commopent sizes

simMM A matrix of floats (n_cluster, n_cluster) for the similarity matrix between clus-

ters. simMM[i,j] means the proportion of cluster i will be assigned to cluster j,

hence colSums(simMM) are ones.

min_iter integer(1). number of minimum iterations
max_iter integer(1). number of maximum iterations

logLik_threshold

A float. The threshold of logLikelihood increase for detecting convergence

Value

a list containing mu, a vector for estimated latent proportion of each cluster, logLik, a float for the estimated log likelihood, simMM, the input of simMM, codeX, the input of X, X_prop, the proportion of clusters in the input X, predict_X_prop, and the predicted proportion of clusters based on mu and simMM.

Ren2021 9

Examples

```
X = c(100, 300, 1500, 500, 1000)
simMM = create_simMat(5, confuse_rate=0.2)
multinom_EM(X, simMM)
```

Ren2021

Count matrix and metadata of a large COVID-19 scRNA-seq data co-hort.

Description

A data containing the count matrix, metadata from a large COVID-19 cohort. Count matrices are calculated based on the number of cells in each cell type. The information in the design matrix is collected in the original paper.

Usage

Ren2021

Format

A list with 7 items:

countM the count matrix for all samples comming from different condition
 designM the corresponding metadata related to each sample
 source the source of this dataset

Source

```
https://www.nature.com/articles/nature24489
```

simulation

Simulated dataset with two conditions

Description

A data containing the count matrices, the similarity matrix and other variables used to generate the similarity matrix from a simulated single cell RNA sequencing data with two conditions. Dirichlet distribution was used to generate a proportion vector for cell types based on the defined true proportions. Multinomial distribution was used to generate simulated cell numbers. Cells were selected from cell pools based on cell numbers and gene expression matrices were processed by Seurat. The count matrix were calculated by the clustering results, and the similarity matrix was calculated using knn graph. The labels are the groud true annotation of each cell.

10 simulator_base

Usage

simulation

Format

```
A list with 5 items:
```

```
numb_cond1 the count matrix of condition 1numb_cond2 the count matrix of condition 2
```

knn_mat the similariy matrix

knnGraphs the knn graphs information used to calculate the similarity matrix.

labels the clusters' label for each simulated single cell

simulator_base

Composition size simulator

Description

Directly simulating the composition size from a Dirichlet-Multinomial distribution with replicates for two conditions.

Usage

```
simulator_base(
  totals_cond1,
  totals_cond2,
  dirichlet_s1,
  dirichlet_s2,
  similarity_mat = NULL
)
```

Arguments

Value

a list of two matrices for composition sizes in each replicate and each cluster in both conditions.

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Examples

```
K <- 2
totals1 = c(100, 800, 1300, 600)
totals2 = c(250, 700, 1100)
diri_s1 = rep(1, K) * 20
diri_s2 = rep(1, K) * 20
confuse_rate = 0.2
simil_mat = create_simMat(2, 0.2)
sim_dat <- simulator_base(totals1, totals2, diri_s1, diri_s2, simil_mat)</pre>
```

svm_simMat

Calculate stochastic transition matrix between clusters from a data frame including information about clustering

Description

The transition probability from cluster i to j is calculated based on the information used to cluster cells. It is estimated by the misclassification rate from cluster i to j comparing the original labels with the labels predicted by support vector machine with 5-fold cross validation.

Usage

```
svm_simMat(dataframe)
```

Arguments

dataframe

a data frame contains the information used for clustering and the original label of each cell. The original labels should have the column name 'clusterRes'.

Value

a similarity matrix estimated by 5-fold cross validation support vector machine.

```
data(Kang2017)
svm_mat = svm_simMat(Kang2017$svmDF)
```

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