

Package ‘scDDboost’

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Type Package

Title A compositional model to assess expression changes from single-cell rna-seq data

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Description

scDDboost is an R package to analyze changes in the distribution of single-cell expression data between two experimental conditions. Compared to other methods that assess differential expression, scDDboost benefits uniquely from information conveyed by the clustering of cells into cellular subtypes. Through a novel empirical Bayesian formulation it calculates gene-specific posterior probabilities that the marginal expression distribution is the same (or different) between the two conditions. The implementation in scDDboost treats gene-level expression data within each condition as a mixture of negative binomial distributions.

License GPL (>= 2)

Imports Rcpp (>= 0.12.11), RcppEigen (>= 0.3.2.9.0), EBSeq, BiocParallel, mclust, SingleCellExperiment, cluster, Oscope, SummarizedExperiment, stats, methods

biocViews SingleCell, Software, Clustering, Sequencing, GeneExpression, DifferentialExpression, Bayesian

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Author Xiuyu Ma [cre, aut],
Michael A. Newton [ctb]

Maintainer Xiuyu Ma <watsonforfun@gmail.com>

Contents

scDDboost-package	2
calD	3
clusHelper	4
detK	4
EBS	5
extractInfo	5
gCl	6
genRClus	6
getDD	7
getSizeofDD	7
getZ1Z2	8
gRef	8
isRef	9
LL	9
lpt1t2	10
lpzgt	10
mdd	11
pat	11
pdd	12
pddAggregate	13
pddRandom	13
rwMle	14
sim_dat	15

Index

16

Description

scDDboost is an R package to analyze changes in the distribution of single-cell expression data between two experimental conditions. Compared to other methods that assess differential expression, scDDboost benefits uniquely from information conveyed by the clustering of cells into cellular subtypes. Through a novel empirical Bayesian formulation it calculates gene-specific posterior probabilities that the marginal expression distribution is the same (or different) between the two conditions. The implementation in scDDboost treats gene-level expression data within each condition as a mixture of negative binomial distributions.

Details

The DESCRIPTION file: This package was not yet installed at build time.

Index: This package was not yet installed at build time.

Package used to score evidence of differential distribution in single-cell RNA-seq data

Author(s)

Xiuyu Ma [cre, aut], Michael A. Newton [ctb]

Maintainer: Xiuyu Ma <watsonforfun@gmail.com>

References

<https://projecteuclid.org/journals/annals-of-applied-statistics/volume-15/issue-2/A-compositional-model-to-assess-expression-changes-from-single-cell/10.1214/20-AOAS1423.short>

See Also

<https://github.com/wiscstatman/scDDboost/blob/master/DESCRIPTION>

Examples

```
data(sim_dat)
dat = extractInfo(sim_dat)
data_counts = dat$count_matrix
cd = dat$condition
bp <- BiocParallel::MulticoreParam(4)
D_c = calD(data_counts, bp)
pDD = pdd(data_counts, cd, bp, D_c)
```

calD *calculate distance matrix*

Description

calculate distance matrix

Usage

calD(data, bp)

Arguments

data	transcripts
bp	bioc parallel parameter

Value

distance matrix

Examples

```
data(sim_dat)
dat <- extractInfo(sim_dat)
data_counts <- dat$count_matrix
bp <- BiocParallel::MulticoreParam(4)
D_c <- calD(data_counts,bp)
```

clusHelper

function to get intra and inter distance for clusters

Description

function to get intra and inter distance for clusters

Usage

```
clusHelper(D, i)
```

Arguments

D	distance matrix
i	number of clusters

Value

vector of intra and inter distance

detK

determine the number of clusters

Description

determine the number of clusters

Usage

```
detK(D, epi = 1)
```

Arguments

D	distance matrix
epi	threshold for cutting off

Value

number of clusters

Examples

```
data(sim_dat)
dat <- extractInfo(sim_dat)
data_counts <- dat$count_matrix
bp <- BiocParallel::MulticoreParam(4)
D_c <- calD(data_counts, bp)
detK(D_c)
```

EBS

accelerated empirical bayesian

Description

accelerated empirical bayesian

Usage

```
EBS(data, conditions, gclus, sf, iter = 10, hyper, PP, stp1, stp2)
```

Arguments

data	single cell expression matrix, row as genes column as cells
conditions	partition of cells
gclus	partition of genes
sf	size factors
iter	maximum iteration step of EM
hyper	hyper parameters for beta distributions
PP	pattern of partitions
stp1	step size of hyperparameter alpha (shared by all units) in one step EM
stp2	step size of hyperparameter beta (unit specific) in one step EM

Value

posterior probability of mean expression pattern

extractInfo

extract count matrix from SingleCellExperiment object

Description

extract count matrix from SingleCellExperiment object

Usage

```
extractInfo(data)
```

Arguments

data SingleCellExperiment object

Value

list of count matrix and condition vector

Examples

```
data(sim_dat)
dat <- extractInfo(sim_dat)
```

gCl

gene_level cluster

Description

gene_level cluster

Usage

```
gCl(data, bp)
```

Arguments

data	transcripts
bp	bioc parallel parameter

Value

return a matrix whose row represent gene specific cluster

genRClus

generate random clusterings

Description

generate random clusterings

Usage

```
genRClus(D, a, K)
```

Arguments

D	distance matrix of cells
a	paramter for weights
K	number of subtypes

Value

random generated clustering of cells

getDD	<i>index of DD genes under FDR control</i>
-------	--

Description

index of DD genes under FDR control

Usage

```
getDD(pDD, FDR = 0.01)
```

Arguments

pDD	probability of genes being DD
FDR	fdr to be controlled

Value

index of positive genes

Examples

```
p_dd <- c(0.01, 0.99, 0.7, 0.5)
getDD(p_dd)
```

getSizeofDD	<i>number of DD genes under FDR control</i>
-------------	---

Description

number of DD genes under FDR control

Usage

```
getSizeofDD(pDD, FDR = 0.01)
```

Arguments

pDD	estimated probability of being DD
FDR	fdr to be controlled

Value

number of positive genes

Examples

```
p_dd <- c(0.1, 0.99, 1, 0.05, 0.05)
getSizeofDD(p_dd)
```

getZ1Z2*function to get counts of cluster sizes at two conditions***Description**

function to get counts of cluster sizes at two conditions

Usage

```
getZ1Z2(ccl, cd)
```

Arguments

ccl	clustering label
cd	condition label

Value

return list of counts

gRef*generate reference matrix***Description**

generate reference matrix

Usage

```
gRef(Posp)
```

Arguments

Posp	possible partition of data
------	----------------------------

Value

return a matrix indicate the refinement relation between different partitions.

isRef	<i>check refinement relation between two clusters</i>
-------	---

Description

check refinement relation between two clusters

Usage

```
isRef(x, y)
```

Arguments

x	a cluster
y	a cluster

Value

whether x refines y

LL	<i>likelihood function for hyperparameters estimation</i>
----	---

Description

likelihood function for hyperparameters estimation

Usage

```
LL(param, x, d0)
```

Arguments

param	parameters to be determined by MLE
x	distance matrix of cells
d0	rate parameter of prior of 1 / true distance

Value

return hyperparameteres a.

lpt1t2*log likelihood of z1,z2 given t1,t2***Description**

log likelihood of z1,z2 given t1,t2

Usage`lpt1t2(z1, z2, pp, alpha1, alpha2)`**Arguments**

<code>z1</code>	counts of each group in condition 1
<code>z2</code>	counts of each group in condition 2
<code>pp</code>	a partition
<code>alpha1</code>	parameter of double dirichlet prior
<code>alpha2</code>	parameter of double dirichlet prior

Value

log likelihood of z1,z2 given t1,t2

lpzgt*log likelihood of aggregated multinomial counts z given aggregated proportions t***Description**

log likelihood of aggregated multinomial counts z given aggregated proportions t

Usage`lpzgt(z, pp, alpha)`**Arguments**

<code>z</code>	counts of each group in one condition
<code>pp</code>	a partition
<code>alpha</code>	parameter of double dirichlet prior

Value

log likelihood of aggregated multinomial counts z given aggregated proportions t

mdd	<i>posterior of proportion change given mixture double dirichlet prior</i>
-----	--

Description

posterior of proportion change given mixture double dirichlet prior

Usage

```
mdd(z1, z2, pat, alpha1, alpha2)
```

Arguments

z1	counts of each group in condition 1
z2	counts of each group in condition 2
pat	partition patterns
alpha1	parameter of double dirichlet prior
alpha2	parameter of double dirichlet prior

Value

posterior of proportion change

pat	<i>generating partition patterns</i>
-----	--------------------------------------

Description

generating partition patterns

Usage

```
pat(K)
```

Arguments

K	number of elements
---	--------------------

Value

all possible partition of K elements

Examples

```
pat(3)
```

pdd	<i>calculate posterior probabilities of a gene to be differential distributed</i>
-----	---

Description

calculate posterior probabilities of a gene to be differential distributed

Usage

```
pdd(
  data,
  cd,
  bp,
  D,
  random = TRUE,
  norm = TRUE,
  epi = 1,
  Upper = 1000,
  nrandom = 50,
  iter = 20,
  reltol = 0.001,
  stp1 = 1e-06,
  stp2 = 0.01,
  K = 0
)
```

Arguments

<code>data</code>	normalized preprocessed transcripts
<code>cd</code>	conditions label
<code>bp</code>	bioc parallel parameter
<code>D</code>	distance matrix of cells or cluster of cells or a given clustering
<code>random</code>	boolean indicator of whether randomization has been implemented on distance matrix
<code>norm</code>	boolean indicator of whether the input expression data is normalized
<code>epi</code>	tol for change of validity score in determining number of clusters
<code>Upper</code>	bound for hyper parameters optimization
<code>nrandom</code>	number of random generated distance matrix
<code>iter</code>	max number of iterations for EM
<code>reltol</code>	relative tolerance for optimization on weighting parameters
<code>stp1</code>	step size of hyperparameter alpha (shared by all units) in one step EM
<code>stp2</code>	step size of hyperparameter beta (unit specific) in one step EM
<code>K</code>	number of subtypes, could be user specified or determined internally(set to 0)

Value

posterior probabilities of a gene to be differential distributed

Examples

```
data(sim_dat)
dat <- extractInfo(sim_dat)
data_counts <- dat$count_matrix
cd <- dat$condition
bp <- BiocParallel::MulticoreParam(4)
D_c <- calD(data_counts, bp)
pDD <- pdd(data_counts, cd, bp, D_c)
```

pddAggregate

*function to aggregate intermediate results and get prob of DD***Description**

function to aggregate intermediate results and get prob of DD

Usage

```
pddAggregate(z1, z2, Posp, DE, K, REF)
```

Arguments

z1	counts of cluster sizes in condition 1
z2	counts of cluster sizes in condition 2
Posp	partition of cells
DE	posterior probabilities of DE patterns
K	number of clusters
REF	reference matrix indicating relation of nested partitions

Value

return vector of prob of DD

pddRandom

*calculate PDD when add random noise in distance matrix***Description**

calculate PDD when add random noise in distance matrix

Usage

```
pddRandom(data, cd, K, D, a, sz, hp, Posp, iter, REF, stp1, stp2)
```

Arguments

<code>data</code>	normalized preprocessed transcripts
<code>cd</code>	condition label
<code>K</code>	number of subgroups
<code>D</code>	distance matrix of cells
<code>a</code>	shape param for weights
<code>sz</code>	size factors
<code>hp</code>	hyper parameters for EBSeq
<code>Posp</code>	partition patterns
<code>iter</code>	max number of iterations for EM in EBSeq
<code>REF</code>	refinement relation matrix
<code>stp1</code>	step size of hyperparameter alpha (shared by all units) in one step EM
<code>stp2</code>	step size of hyperparameter beta (unit specific) in one step EM

Value

posterior probabilities under random distance matrix

<code>rwMle</code>	<i>MLE for random weighting parameter</i>
--------------------	---

Description

MLE for random weighting parameter

Usage

```
rwMle(D, reltol)
```

Arguments

<code>D</code>	distance matrix of cells
<code>reltol</code>	tolerance of convergence

Value

MLE of random weighting parameter

<code>sim_dat</code>	<i>scDDboost</i>
----------------------	------------------

Description

simulated data for demonstration, data are mixture negative binomial distributed

Usage

```
data(sim_dat)
```

Format

An object of class "list".

Examples

```
data(sim_dat)
```

Index

- * **Empirical Bayes, clustering, random weighting, local false discovery rate**
 - scDDboost-package, 2
- * **datasets**
 - sim_dat, 15
- * **internal**
 - pddRandom, 13
- calD, 3
- clusHelper, 4
- detK, 4
- EBS, 5
- extractInfo, 5
- gCl, 6
- genRClus, 6
- getDD, 7
- getSizeofDD, 7
- getZ1Z2, 8
- gRef, 8
- isRef, 9
- LL, 9
- lpt1t2, 10
- lpzgt, 10
- mdd, 11
- pat, 11
- pdd, 12
- pddAggregate, 13
- pddRandom, 13
- rwMle, 14

scDDboost (scDDboost-package), 2
scDDboost-package, 2
sim_dat, 15