Package 'SC.MEB'

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Description Spatial clustering with hidden markov random field fitted via EM algorithm, details of which can be found in Yi Yang (2021) <doi:10.1101 2021.06.05.447181="">. It is not only computationally efficient and scalable to the sample size increment, but also is capable of choosing the smoothness parameter and the number of clusters as well.</doi:10.1101>
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 ${\tt ClusterPlot}$

ClusterPlot.

Description

The function ClusterPlot is used to Visualize spatial clusters.

Usage

```
ClusterPlot(out, pos, size = 5, shape = 15)
```

Arguments

out	is the output of function selectK.
pos	is a n-by-2 matrix of position.
size	is a positive value for characterizing the size of point in the plot, which is the same as size in ggplot2.
shape	is a positive value for characterizing the shape of point in the plot, which is the same as shape in ggplot2.

Details

The function ClusterPlot is used to Visualize spatial clusters.

Value

```
a ggplot2 object.
```

```
pos = cbind(rep(1:5, each=5), rep(1:5, 5))
out = list()
out[[1]] = ""
out[[2]] = rep(1:5, each = 5)
ClusterPlot(out, pos)
```

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find_neighbors2

find_neighbors2.

Description

find_neighbors2 was used to find the neighborhood of spot.

Usage

```
find_neighbors2(sce, platform)
```

Arguments

sce

is a SingleCellExperiment object containing PCA and position informatin.

platform

is the name of spatial transcriptomic platform. Specify 'Visium' for hex lattice geometry or 'ST' for square lattice geometry. Specifying this parameter is

optional as this information is included in their metadata.

Details

find_neighbors2 was used to find the neighborhood of spot.

Value

a sparse matrix recording the information of neighborhood.

Examples

```
data(sce)
platform = "ST"
Adj <- find_neighbors2(sce, platform)
```

```
getneighborhood_fast getneighborhood_fast
```

Description

an efficient function to find the neighborhood based on the matrix of position and a pre-defined cutoff

Usage

```
getneighborhood_fast(x, cutoff)
```

ICMEM

Arguments

x is a n-by-2 matrix of position.

cutoff is a threashold of Euclidean distance to decide whether a spot is an neighborhood

of another spot. For example, if the Euclidean distance between spot A and B is

less than cutoff, then A is taken as the neighbourhood of B.

Value

A sparse matrix containing the neighbourhood

Examples

```
pos = cbind(rep(1:5, each=5), rep(1:5, 5))
Adj = getneighborhood_fast(pos, 2)
```

ICMEM

ICMEM.

Description

The function ICMEM was used to conduct spatial clustering with hidden Markov random field for a sequence of beta and fixed number of clusters

Usage

```
ICMEM(
   y,
   x_int,
   Adj,
   mu_int,
   sigma_int,
   alpha,
   beta_grid,
   PX,
   maxIter_ICM,
   maxIter
)
```

Arguments

y is a matrix of PCs containing gene expression.

x_int is a vector of initial cluster label.

Adj is a matrix containing neighborhood information generated by find_neighbors2.

mu_int is a initial mean vector. we often generated it by Gaussian mixture model.

sigma_int is a initial co-variance matrix. we often generated it by Gaussian mixture model.

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alpha is a intercept.

beta_grid is a sequence of smoothing parameter that can be specified by user.

PX is a logical value specifying the parameter expansion in EM algorithm.

maxIter_ICM is the maximum iteration of ICM algorithm.

maxIter is the maximum iteration of EM algorithm.

Details

The function ICMEM was used to conduct spatial clustering with hidden Markov random field for fixed beta and fixed number of clusters

Value

a list.

The item 'x' is the clustering result.

The item 'gam' is the posterior probability matrix.

The item 'ell' is the opposite log-likelihood.

The item 'mu' is the mean of each component.

The item 'sigma' is the variance of each component.

Examples

```
y = matrix(rnorm(50, 0, 1), 25,2)
pos = cbind(rep(1:5, each=5), rep(1:5, 5))
Adj = getneighborhood_fast(pos, 1.2)
beta_grid = c(0.5,1)
G = 2
fit_int = Mclust(y, G = G)
x_gmm <- fit_int$classification
mu_int <- unname(fit_int$parameter$mean)
sigma_int <- unname(fit_int$parameter$variance$sigma)
alpha <- -log(fit_int$parameter$pro)*0
reslist <- ICMEM(y = y, x_int = x_gmm, Adj = Adj, mu_int = mu_int, sigma_int = sigma_int,
alpha = alpha, beta_grid = beta_grid,
PX = TRUE, maxIter_ICM = 10, maxIter = 50)</pre>
```

parafun parafun.

Description

The function parafun implements the model SC-MEB for fixed number of clusters and a sequence of beta with initial value from Gaussian mixture model

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Usage

```
parafun(
   y,
   Adj,
   G,
   beta_grid = seq(0, 4, 0.2),
   PX = TRUE,
   maxIter_ICM = 10,
   maxIter = 50
)
```

Arguments

y is n-by-d PCs.

Adj is a sparse matrix of neighborhood.

G is an integer specifying the numbers of clusters.

beta_grid is a numeric vector specifying the smoothness parameter of Random Markov

Field. The default is seq(0,4,0.2).

PX is a logical value specifying the parameter expansion in EM algorithm.

maxIter_ICM is the maximum iteration of ICM algorithm. The default is 10.
maxIter is the maximum iteration of EM algorithm. The default is 50.

Details

The function parafun implements the model SC-MEB for fixed number of clusters and a sequence of beta with initial value from Gaussian mixture model

Value

a list, We briefly explain the output of the SC.MEB.

The item 'x' storing clustering results.

The item 'gam' is the posterior probability matrix.

The item 'ell' is the opposite log-likelihood.

The item 'mu' is the mean of each component.

The item 'sigma' is the variance of each component.

```
y = matrix(rnorm(50, 0, 1), 25,2)
pos = cbind(rep(1:5, each=5), rep(1:5, 5))
Adj_sp = getneighborhood_fast(pos, 1.2)
beta_grid = c(0.5,1)
G = 2
out = parafun(y, Adj_sp, G, beta_grid)
```

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PC

simulated PCs

Description

A dataset containing PCs

Usage

```
data(PC)
```

Format

It is a matrix containing 5 PCs the variables are listed as following

```
PC1 The 1th PC
PC2 The 2th PC ...
PC5 The 5th PC
```

Examples

```
## run the PC with the Gaussian mixture model
data(PC)
out1 = mclust::Mclust(PC,G = 2)
```

SC.MEB

SC.MEB.

Description

SC.MEB implements the model SC-MEB, spatial clustering with hidden Markov random field using empirical Bayes.

Usage

```
SC.MEB(
   y,
   Adj_sp,
   beta_grid = seq(0, 4, 0.2),
   K_set = 2:10,
   parallel = TRUE,
   num_core = 5,
   PX = TRUE,
   maxIter_ICM = 10,
   maxIter = 50
)
```

SC.MEB

Arguments

у	is n-by-d PCs.
Adj_sp	is a sparse matrix of neighborhood. It is often generated from function find_neighbors2 or getneighborhood_fast.
beta_grid	is a numeric vector specifying the smoothness parameter of Random Markov Field. The default is $seq(0,4,0.2)$.
K_set	is an integer vector specifying the numbers of mixture components (clusters) for which the BIC is to be calculated. The default is $K = 2:10$.
parallel	is a logical value to decide whether the function SC.MEB run in parallel. The default is TRUE.
num_core	is an integer value to decide how many cores are used to run SC.MEB in parallel.
PX	is a logical value to decide whether to use parameter expansion in EM algorithm
maxIter_ICM	is the maximum iteration of ICM algorithm. The default is 10.
maxIter	is the maximum iteration of EM algorithm. The default is 50.

Details

SC.MEB can implements the model SC-MEB in parallel which can improve the speed of the computation.

Value

a list, We briefly explain the output of the SC.MEB.

The item 'x' contains clustering results.

The item 'gam' is the posterior probability matrix.

The item 'ell' is the opposite log-likelihood.

The item 'mu' is the mean of each component.

The item 'sigma' is the variance of each component.

References

Yang Y, Shi X, Zhou Q, et al. SC-MEB: spatial clustering with hidden Markov random field using empirical Bayes[J]. bioRxiv, 2021.

```
y = matrix(rnorm(50, 0, 1), 25,2)
pos = cbind(rep(1:5, each=5), rep(1:5, 5))
Adj_sp = getneighborhood_fast(pos, 1.2)
beta_grid = c(0.5,1)
K_set = 2:3
out = SC.MEB(y, Adj_sp, beta_grid, K_set, TRUE, 2)
```

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sce

A simulated SingleCellExperiment

Description

A dataset of SingleCellExperiment

Usage

```
data(sce)
```

Format

It is a SingleCellExperiment object with gene expression and meta information

References

Amezquita R A, Lun A T L, Becht E, et al. Orchestrating single-cell analysis with Bioconductor[J]. Nature methods, 2020, 17(2): 137-145.

Examples

```
## find the neighborhood of spots in SingleCellExperiment
data(sce)
out = find_neighbors2(sce, "ST")
```

selectK

selectK.

Description

The function selectK is used to select the best K according to BIC or Modified BIC criterion.

Usage

```
selectK(SCobject, K_set = 2:10, criterion = "BIC", c = 1)
```

Arguments

SCobject is an object generated from SC.MEB function.

K_set is a integer vector used in SC.MEB. The default is 2:10

criterion is a character specifying the criterion for selecting K. The default value is BIC.

The alternative value MBIC can also be used.

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С

is a positive value in the modified BIC. The default is 1. Here we briefly explain how to choose the parameter c in the modified BIC. In general, For the ST or Visium dataset, it often ranges from 0.4 to 1 while for the MERFISH dataset with large number of cells, it often becomes larger, for example 10,20. Most importantly, SC-MEB is fast, scaling well in terms of sample size, which allow the user to tune the c based on their prior knowledge about the tissues or cells.

Details

The function selectK is used to select the best K according to BIC or Modified BIC criterion.

Value

a list contains two items. one is for the best K and the other is the clustering labels of n spots.

Examples

```
y = matrix(rnorm(50, 0, 1), 25,2)
pos = cbind(rep(1:5, each=5), rep(1:5, 5))
Adj_sp = getneighborhood_fast(pos, 1.2)
beta_grid = c(0.5,1)
K_set = 2:3
out = SC.MEB(y, Adj_sp, beta_grid, K_set, TRUE, 2)
selectK(out, K_set)
```

selectKPlot

selectKPlot.

Description

The function selectKPlot is used to demonstrate the scatter plot of BIC or Modified BIC vs K for selecting the best K.

Usage

```
selectKPlot(SCobject, K_set = 2:10, criterion = "BIC", c = 1)
```

Arguments

SCobject is a object generated from SC.MEB function.

K_set is the corresponding K_set used in your previous function SC.MEB.

criterion is a character specifying the criterion for selecting K. The default is BIC, the

alternative criterion MBIC can also be used.

c is a positive value in modified BIC. The default is 1. Here we briefly explain

how to choose the parameter c in the modified BIC. In general, For the ST or Visium dataset, it often ranges from 0.4 to 1 while for the MERFISH dataset with large number of cells, it often becomes larger, for example 10,20. Most importantly, SC-MEB is fast, scaling well in terms of sample size, which allow the user to tune the c based on their prior knowledge about the tissues or cells.

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Details

The function selectKPlot is used to demonstrate the scatter plot of BIC or Modified BIC vs K for selecting the best K.

Value

```
a ggplot2 object.
```

Examples

```
y = matrix(rnorm(50, 0, 1), 25,2)
pos = cbind(rep(1:5, each=5), rep(1:5, 5))
Adj_sp = getneighborhood_fast(pos, 1.2)
beta_grid = c(0.5,1)
K_set = 2:3
out = SC.MEB(y, Adj_sp, beta_grid, K_set, TRUE, 2)
selectKPlot(out, K_set)
```

spatialPreprocess

Preprocess a spatial dataset for SC-MEB

Description

Adds metadata required for downstream analyses, and (optionally) performs PCA on log-normalized expression of top HVGs.

Usage

```
spatialPreprocess(
    sce,
    platform = c("Visium", "ST"),
    n.PCs = 15,
    n.HVGs = 2000,
    skip.PCA = FALSE,
    log.normalize = TRUE,
    assay.type = "logcounts",
    BSPARAM = BiocSingular::ExactParam()
)
```

Arguments

sce	SingleCellExperiment to preprocess
platform	Spatial sequencing platform. Used to determine spot layout and neighborhood structure (Visium = hex, ST = square).
n.PCs	Number of principal components to compute. We suggest using the top 15 PCs in most cases.
n.HVGs	Number of highly variable genes to run PCA upon.

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skip.PCA Skip PCA (if dimensionality reduction was previously computed.)

log.normalize Whether to log-normalize the input data with scater. May be omitted if log-

normalization previously computed.

assay.type Name of assay in sce containing normalized counts. Leave as "logcounts" un-

less you explicitly pre-computed a different normalization and added it to sce under another assay. Note that we do not recommend running BayesSpace on

PCs computed from raw counts.

BSPARAM A BiocSingularParam object specifying which algorithm should be used to per-

form the PCA. By default, an exact PCA is performed, as current spatial datasets are generally small (<10,000 spots). To perform a faster approximate PCA, please specify FastAutoParam() and set a random seed to ensure reproducibil-

ity.

Value

SingleCellExperiment with PCA and SC.MEB metadata

```
## read the simulated data
data(sce)
platform = "ST"
out = find_neighbors2(sce, platform)
```

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