# Package 'BayesSpace'

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Title Clustering and Resolution Enhancement of Spatial Transcriptomes

**Description** Tools for clustering and enhancing the resolution of spatial gene expression experiments. BayesSpace clusters a low-dimensional representation of the gene expression matrix, incorporating a spatial prior to encourage neighboring spots to cluster together. The method can enhance the resolution of the low-dimensional representation into ``sub-spots", for which features such as gene expression or cell type composition can be imputed.

**Depends** R (>= 4.0.0), SingleCellExperiment

Imports Rcpp (>= 1.0.4.6), stats, purrr, scater, scran, SummarizedExperiment, coda, rhdf5, S4Vectors, Matrix, assertthat, mclust, RCurl, DirichletReg, xgboost, utils, ggplot2, scales, BiocFileCache, BiocSingular

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BugReports https://github.com/edward130603/BayesSpace/issues

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```
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clusterPlot

Plot spatial cluster assignments.

# **Description**

Plot spatial cluster assignments.

# Usage

```
clusterPlot(
    sce,
    label = "spatial.cluster",
    palette = NULL,
    color = NULL,
    platform = NULL,
    is.enhanced = NULL,
    ...
)
```

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# Arguments

sce	SingleCellExperiment. If fill is specified and is a string, it must exist as a column in colData(sce).
label	Labels used to color each spot. May be the name of a column in colData(sce), or a vector of discrete values.
palette	Optional vector of hex codes to use for discrete spot values.
color	Optional hex code to set color of borders around spots. Set to NA to remove borders.
platform	Spatial sequencing platform. If "Visium", the hex spot layout will be used, otherwise square spots will be plotted.  NOTE: specifying this argument is only necessary if sce was not created by spatialCluster() or spatialEnhance().
is.enhanced	True if sce contains subspot-level data instead of spots. Spatial sequencing platform. If true, the respective subspot lattice for each platform will be plotted. NOTE: specifying this argument is only necessary if sce was not created by spatialCluster() or spatialEnhance().
	Additional arguments for geom_polygon(). size, to specify the linewidth of these borders, is likely the most useful.

# Value

Returns a ggplot object.

# See Also

Other spatial plotting functions: featurePlot()

# Examples

```
sce <- exampleSCE()
clusterPlot(sce)</pre>
```

enhanceFea	atures
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Predict feature vectors from enhanced PCs.

# Description

Predict feature vectors from enhanced PCs.

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#### Usage

```
enhanceFeatures(
   sce.enhanced,
   sce.ref,
   feature_names = NULL,
   model = c("xgboost", "dirichlet", "lm"),
   use.dimred = "PCA",
   assay.type = "logcounts",
   altExp.type = NULL,
   feature.matrix = NULL,
   nrounds = 0,
   train.n = round(ncol(sce.ref) * 2/3)
)
```

#### **Arguments**

sce.enhanced SingleCellExperiment object with enhanced PCs.

sce.ref SingleCellExperiment object with original PCs and expression.

feature\_names List of genes/features to predict expression/values for.

model Model used to predict enhanced values.

use.dimred Name of dimension reduction to use.

assay.type Expression matrix in assays(sce.ref) to predict.

altExp.type Expression matrix in altExps(sce.ref) to predict. Overrides assay.type if

specified.

feature.matrix Expression/feature matrix to predict, if not directly attached to sce.ref. Must

have columns corresponding to the spots in sce.ref. Overrides assay.type

and altExp. type if specified.

nrounds Nonnegative integer to set the nrounds parameter (max number of boosting

iterations) for xgboost. nrounds = 100 works reasonably well in most cases. If nrounds is set to 0, the parameter will be tuned using a train-test split. We recommend tuning nrounds for improved feature prediction, but note this will

increase runtime.

train.n Number of spots to use in the training dataset for tuning nrounds. By default,

2/3 the total number of spots are used.

#### **Details**

Enhanced features are computed by fitting a predictive model to a low-dimensional representation of the original expression vectors. By default, a linear model is fit for each gene using the top 15 principal components from each spot, i.e. lm(gene ~ PCs), and the fitted model is used to predict the enhanced expression for each gene from the subspots' principal components.

Diagnostic measures, such as RMSE for xgboost or R.squared for linear regression, are added to the 'rowData' of the enhanced experiment if the features are an assay of the original experiment. Otherwise they are stored as an attribute of the returned matrix/altExp.

Note that feature matrices will be returned and are expected to be input as  $p \times n$  matrices of p-dimensional feature vectors over the n spots.

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# Value

If assay.type or altExp.type are specified, the enhanced features are stored in the corresponding slot of sce.enhanced and the modified SingleCellExperiment object is returned.

If feature.matrix is specified, or if a subset of features are requested, the enhanced features are returned directly as a matrix.

# **Examples**

```
set.seed(149)
sce <- exampleSCE()
sce <- spatialCluster(sce, 7, nrep=100, burn.in=10)
enhanced <- spatialEnhance(sce, 7, init=sce$spatial.cluster, nrep=100, burn.in=10)
enhanced <- enhanceFeatures(enhanced, sce, feature_names=c("gene_1", "gene_2"))</pre>
```

exampleSCE

Create minimal SingleCellExperiment for documentation examples.

# Description

Create minimal SingleCellExperiment for documentation examples.

#### Usage

```
exampleSCE(nrow = 8, ncol = 12, n_genes = 100, n_PCs = 10)
```

#### **Arguments**

nrow	Number of rows of spots
ncol	Number of columns of spots
n_genes	Number of genes to simulate

n\_PCs Number of principal components to include

#### **Details**

Inspired by scuttle's mockSCE().

#### Value

A SingleCellExperiment object with simulated counts, corresponding logcounts and PCs, and positional data in colData. Spots are distributed over an (nrow x ncol) rectangle.

```
set.seed(149)
sce <- exampleSCE()</pre>
```

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featurePlot

Plot spatial gene expression.

# Description

Plot spatial gene expression.

# Usage

```
featurePlot(
    sce,
    feature,
    assay.type = "logcounts",
    diverging = FALSE,
    low = NULL,
    high = NULL,
    mid = NULL,
    color = NULL,
    platform = NULL,
    is.enhanced = NULL,
    ...
)
```

# Arguments

sce	SingleCellExperiment. If feature is specified and is a string, it must exist as a row in the specified assay of sce.
feature	Feature vector used to color each spot. May be the name of a gene/row in an assay of sce, or a vector of continuous values.
assay.type	String indicating which assay in sce the expression vector should be taken from.
diverging	If true, use a diverging color gradient in featurePlot() (e.g. when plotting a fold change) instead of a sequential gradient (e.g. when plotting expression).
low, mid, high	Optional hex codes for low, mid, and high values of the color gradient used for continuous spot values.
color	Optional hex code to set color of borders around spots. Set to NA to remove borders.
platform	Spatial sequencing platform. If "Visium", the hex spot layout will be used, otherwise square spots will be plotted.
	NOTE: specifying this argument is only necessary if sce was not created by spatialCluster() or spatialEnhance().
is.enhanced	True if sce contains subspot-level data instead of spots. Spatial sequencing platform. If true, the respective subspot lattice for each platform will be plotted. NOTE: specifying this argument is only necessary if sce was not created by spatialCluster() or spatialEnhance().
• • •	Additional arguments for geom_polygon(). size, to specify the linewidth of these borders, is likely the most useful.

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# Value

Returns a ggplot object.

#### See Also

Other spatial plotting functions: clusterPlot()

# **Examples**

```
sce <- exampleSCE()
featurePlot(sce, "gene_2")</pre>
```

getRDS

Download a processed sample from our S3 bucket

# Description

Datasets are cached locally using BiocFileCache. The first time using this function, you may need to consent to creating a BiocFileCache directory if one does not already exist.

# Usage

```
getRDS(dataset, sample, cache = TRUE)
```

# **Arguments**

dataset Dataset identifier sample Sample identifier

cache If true, cache the dataset locally with BiocFileCache. Otherwise, download

directly from our S3 bucket. Caching saves time on subsequent loads, but con-

sumes disk space.

#### Value

sce A SingleCellExperiment with positional information in colData and PCs based on the top 2000 HVGs

```
sce <- getRDS("2018_thrane_melanoma", "ST_mel1_rep2", cache=FALSE)</pre>
```

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mcmcChain	Read MCMC chain associated with a BayesSpace clustering or enhancement

# **Description**

BayesSpace stores the MCMC chain associated with a clustering or enhancement on disk in an HDF5 file. The mcmcChain() function reads any parameters specified by the user into a coda::mcmc object compatible with TidyBayes.

#### Usage

```
mcmcChain(sce, params = NULL)
removeChain(sce)
```

# **Arguments**

sce SingleCellExperiment with a file path stored in its metadata.

params List of model parameters to read

#### **Details**

To interact with the HDF5 file directly, obtain the filename from the SingleCellExperiment's metadata: metadata(sce)\$chain.h5. Each parameter is stored as a separate dataset in the file, and is represented as a matrix of size (n\_iterations x n\_parameter\_indices). Parameter choices for the spot-level clustering include:

- z (cluster assignments)
- weights  $(w_i)$
- mu (mean vectors)
- lambda (precision matrix)
- plogLik (pseudo-log-likelihood)

Parameter choices for the subspot-level enhanced clustering include:

- z (cluster assignments)
- weights  $(w_i)$
- Y (enhanced PCs)
- mu (mean vectors)
- lambda (precision matrix)
- Ychange (acceptance rate for the jittering of PCs)

#### Value

Returns an mcmc object containing the values of the requested parameters over the constructed chain.

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#### **Examples**

```
set.seed(149)
sce <- exampleSCE()
sce <- spatialCluster(sce, 7, nrep=100, burn.in=10, save.chain=TRUE)
chain <- mcmcChain(sce)
removeChain(sce)</pre>
```

qTune

Tuning the choice of q (number of clusters) before running spatial-Cluster

# Description

Before running spatialCluster(), we recommend tuning the choice of q by choosing the q that maximizes the model's negative log likelihood over early iterations. qTune() computes the average negative log likelihood for a range of q values over iterations 100:1000, and qPlot() displays the results.

# Usage

```
qPlot(sce, qs = seq(3, 7), force.retune = FALSE, ...)
qTune(sce, qs = seq(3, 7), burn.in = 100, nrep = 1000, ...)
```

#### **Arguments**

sce A SingleCellExperiment object containing the spatial data.

The values of q to evaluate.

force.retune If specified, existing tuning values in sce will be overwritten.

... Other parameters are passed to spatialCluster(). burn.in, nrep
Integers specifying the range of repetitions to compute.

#### **Details**

qTune() takes the same parameters as spatialCluster() and will run the MCMC clustering algorithm up to nrep iterations for each value of q. The first burn. in iterations are discarded as burn-in and the log likelihood is averaged over the remaining iterations.

qPlot() plots the computed negative log likelihoods as a function of q. If qTune() was run previously, i.e. there exists an attribute of sce named "q.logliks", the pre-computed results are displayed. Otherwise, or if force.retune is specified, qplot() will automatically run qTune() before plotting (and can take the same parameters as spatialCluster().

#### Value

qTune() returns a modified sce with tuning log likelihoods stored as an attribute named "q.logliks". qPlot() returns a ggplot object.

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#### **Examples**

```
set.seed(149)
sce <- exampleSCE()
sce <- qTune(sce, seq(3, 7), burn.in=10, nrep=100)
qPlot(sce)</pre>
```

readVisium

Load a Visium spatial dataset as a SingleCellExperiment.

# **Description**

Load a Visium spatial dataset as a SingleCellExperiment.

#### Usage

```
readVisium(dirname)
```

# **Arguments**

dirname

Path to spaceranger output directory (e.g. "sampleID/outs/"). This directory must contain the counts matrix and feature/barcode TSVs in filtered\_feature\_bc\_matrix/, and the spot positions at spatial/tissue\_positions\_list.csv. (These are default locations for spaceranger outputs.)

#### **Details**

We store two variables associated with downstream BayesSpace functions in a list called BayesSpace. data in the SingleCellExperiment's metadata.

- platform is set to "Visium", and is used to determine spot layout and neighborhood structure.
- is.enhanced is set to FALSE to denote the object contains spot-level data.

# Value

SingleCellExperiment containing the counts matrix in counts and spatial data in colData. Array coordinates for each spot are stored in columns row and col, while image coordinates are stored in columns imagerow and imagecol.

```
## Not run:
sce <- readVisium("path/to/outs/")
## End(Not run)</pre>
```

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spatialCluster

Spatial clustering

# Description

Cluster a spatial expression dataset.

# Usage

```
spatialCluster(
  sce,
 q,
 use.dimred = "PCA",
 d = 15,
 platform = c("Visium", "ST"),
  init = NULL,
  init.method = c("mclust", "kmeans"),
 model = c("t", "normal"),
 precision = c("equal", "variable"),
 nrep = 50000,
 burn.in = 1000,
 gamma = NULL,
 mu0 = NULL,
 lambda0 = NULL,
 alpha = 1,
 beta = 0.01,
  save.chain = FALSE,
 chain.fname = NULL
)
```

# **Arguments**

sce	A SingleCellExperiment object containing the spatial data.
q	The number of clusters.
use.dimred	Name of a reduced dimensionality result in reducedDims(sce). If provided, cluster on these features directly.
d	Number of top principal components to use when clustering.
platform	Spatial transcriptomic platform. Specify 'Visium' for hex lattice geometry or 'ST' for square lattice geometry. Specifying this parameter is optional when analyzing SingleCellExperiments processed using readVisium or spatialPreprocess, as this information is included in their metadata.
init	Initial cluster assignments for spots.
init.method	If init is not provided, cluster the top d PCs with this method to obtain initial cluster assignments.
model	Error model. ('normal' or 't')

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precision Covariance structure. ('equal' or 'variable' for EEE and VVV covariance mod-

els, respectively.)

nrep The number of MCMC iterations.

burn.in The number of MCMC iterations to exclude as burn-in period.

gamma Smoothing parameter. Defaults to 2 for platform="ST" and 3 for platform="Visium".

(Values in range of 1-3 seem to work well.)

mu0 Prior mean hyperparameter for mu. If not provided, mu0 is set to the mean of

PCs over all spots.

lambda0 Prior precision hyperparam for mu. If not provided, lambda0 is set to a diagonal

matrix 0.01I.

alpha Hyperparameter for Wishart distributed precision lambda.

beta Hyperparameter for Wishart distributed precision lambda.

save.chain If true, save the MCMC chain to an HDF5 file.

chain. fname File path for saved chain. Tempfile used if not provided.

#### **Details**

The input SCE must have row and col columns in its colData, corresponding to the array row and column coordinates of each spot. These are automatically parsed by readVisium or can be added manually when creating the SCE.

Cluster labels are stored in the spatial.cluster column of the SCE, and the cluster initialization is stored in cluster.init.

# Value

Returns a modified sce with cluster assignments stored in colData under the name spatial.cluster.

#### See Also

spatialPreprocess for preparing the SCE for clustering, spatialEnhance for enhancing the clustering resolution, clusterPlot for visualizing the cluster assignments, featurePlot for visualizing expression levels in spatial context, and mcmcChain for examining the full MCMC chain associated with the clustering.

```
set.seed(149)
sce <- exampleSCE()
sce <- spatialCluster(sce, 7, nrep=100, burn.in=10)</pre>
```

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spatialEnhance

Enhance spot resolution

# **Description**

Enhanced clustering of a spatial expression dataset to subspot resolution.

# Usage

```
spatialEnhance(
  sce,
 platform = c("Visium", "ST"),
 use.dimred = "PCA",
 d = 15,
  init = NULL,
  init.method = c("spatialCluster", "mclust", "kmeans"),
 model = c("t", "normal"),
 nrep = 2e + 05,
 gamma = NULL,
 mu0 = NULL,
 lambda0 = NULL,
 alpha = 1,
 beta = 0.01,
  save.chain = FALSE,
  chain.fname = NULL,
 burn.in = 10000,
  jitter_scale = 5,
  jitter_prior = 0.3,
 verbose = FALSE
)
```

# **Arguments**

sce	A SingleCellExperiment object containing the spatial data.
q	The number of clusters.
platform	Spatial transcriptomic platform. Specify 'Visium' for hex lattice geometry or 'ST' for square lattice geometry. Specifying this parameter is optional when analyzing SingleCellExperiments processed using readVisium, spatialPreprocess, or spatialCluster, as this information is included in their metadata.
use.dimred	Name of a reduced dimensionality result in reducedDims(sce). If provided, cluster on these features directly.
d	Number of top principal components to use when clustering.
init	Initial cluster assignments for spots.
init.method	If init is not provided, cluster the top d PCs with this method to obtain initial cluster assignments.

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The number of MCMC iterations. nrep Smoothing parameter. (Values in range of 1-3 seem to work well.) gamma mu0 Prior mean hyperparameter for mu. If not provided, mu0 is set to the mean of PCs over all spots. lambda0 Prior precision hyperparam for mu. If not provided, lambda0 is set to a diagonal matrix 0.01I. Hyperparameter for Wishart distributed precision lambda. alpha beta Hyperparameter for Wishart distributed precision lambda. If true, save the MCMC chain to an HDF5 file. save.chain

Error model. ('normal' or 't')

File path for saved chain. Tempfile used if not provided. Number of iterations to exclude as burn-in period. The MCMC iterations are burn.in

currently thinned to every 100; accordingly burn. in is rounded down to the

nearest multiple of 100.

jitter\_scale Controls the amount of jittering. Small amounts of jittering are more likely to

be accepted but result in exploring the space more slowly. We suggest tuning

jitter\_scale so that Ychange is on average around 25%-40%.

Scale factor for the prior variance, parameterized as the proportion (default = jitter\_prior

> 0.3) of the mean variance of the PCs. We suggest making jitter\_prior smaller if the jittered values are not expected to vary much from the overall mean of the

spot.

verbose Log progress to stderr.

#### **Details**

model

chain.fname

The enhanced SingleCellExperiment has most of the properties of the input SCE - rowData, colData, reducedDims - but does not include expression data in counts or logcounts. To impute enhanced expression vectors, please use [enhanceFeatures()] after running spatialEnhance.

The colData of the enhanced SingleCellExperiment includes the following columns to permit referencing the subspots in spatial context and linking back to the original spots:

- spot.idx: Index of the spot this subspot belongs to (with respect to the input SCE).
- subspot.idx: Index of the subspot within its parent spot.
- spot.row: Array row of the subspot's parent spot.
- spot.col: Array col of the subspot's parent spot.
- row: Array row of the subspot. This is the parent spot's row plus an offset based on the subspot's position within the spot.
- col: Array col of the subspot. This is the parent spot's col plus an offset based on the subspot's position within the spot.
- imagerow: Pixel row of the subspot. This is the parent spot's row plus an offset based on the subspot's position within the spot.
- imagecol: Pixel col of the subspot. This is the parent spot's col plus an offset based on the subspot's position within the spot.

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#### Value

Returns a new SingleCellExperiment object. By default, the assays of this object are empty, and the enhanced resolution PCs are stored as a reduced dimensionality result accessible with reducedDim(sce, 'PCA').

#### See Also

spatialCluster for clustering at the spot level before enhancing, clusterPlot for visualizing the cluster assignments, enhanceFeatures for imputing enhanced expression, and mcmcChain for examining the full MCMC chain associated with the enhanced clustering.

#### **Examples**

```
set.seed(149)
sce <- exampleSCE()
sce <- spatialCluster(sce, 7, nrep=100, burn.in=10)
enhanced <- spatialEnhance(sce, 7, nrep=100, burn.in=10)</pre>
```

spatialPreprocess

Preprocess a spatial dataset for BayesSpace

#### **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 = ExactParam()
)
```

#### **Arguments**

sce SingleCellExperiment to preprocess

Platform Spatial sequencing platform. Used to determine spot layout and neighborhood structure (Visium = hex, ST = square).

Number of principal components to compute. We suggest using the top 15 PCs in most cases.

spatialPreprocess

n.HVGs Number of highly variable genes to run PCA upon.

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 BayesSpace metadata

```
sce <- exampleSCE()
sce <- spatialPreprocess(sce)</pre>
```

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