Package 'SpaTopic'

January 20, 2025

```
Type Package
Title Topic Inference to Identify Tissue Architecture in Multiplexed
     Images
Version 1.1.0
Date 2024-04-22
Description A novel spatial topic model to integrate both cell type and spatial information to iden-
     tify the complex spatial tissue architecture on multiplexed tissue images without human interven-
     tion. The Package implements a collapsed Gibbs sampling algorithm for infer-
     ence. 'SpaTopic' is scalable to large-scale image datasets without extracting neighborhood infor-
     mation for every single cell. For more details on the methodol-
     ogy, see <https://xiyupeng.github.io/SpaTopic/>.
License GPL (>= 3)
Depends R (>= 3.5.0),
Imports Rcpp (>= 0.12.0), RANN (>= 2.6.0), sf (>= 1.0-12), methods (>=
     3.4), foreach (>= 1.5.0), iterators (>= 1.0),
LinkingTo Rcpp, RcppArmadillo, RcppProgress,
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     (>=1.0),
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```

gibbs.res-class

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		_
gibbs	.res-class A class of the output from 'SpaTopic'	

Description

Outputs from function SpaTopic_inference. A list contains the following members:

- \$Perplexity. The perplexity is for the training data. Let N be the total number of cells across all images. Perplexity = exp(-loglikelihood/N)
- \$Deviance. Deviance = -2loglikelihood.
- \$loglikelihood. The model log-likelihood.
- \$loglike.trace. The log-likelihood for every collected posterior sample. NULL if trace = FALSE.
- \$Beta. Topic content matrix with rows as celltypes and columns as topics
- \$Theta. Topic prevalent matrix with rows as regions and columns as topics
- \$Ndk. Number of cells per topic (col) per region (row).
- \$Nwk. Number of cells per topic (col) per celltype (row).
- \$Z. trace. Number of times cell being assigned to each topic across all posterior samples. We can further compute the posterior distributions of Z (topic assignment) for individual cells.
- \$doc.trace. Ndk for every collected posterior sample. NULL if trace = FALSE.
- \$word.trace. Nwk for every collected posterior sample. NULL if trace = FALSE.

See Also

SpaTopic_inference

lung5

lung5

Example input data for 'SpaTopic'

Description

multiplexed image data on tumor tissue sample from non small cell lung cancer patient

Usage

lung5

Format

'lung5' A data frame with 100149 rows and 4 columns:

image Image ID

X X coordinate of the cell

Y Y coordinate of the cell

type cell type

Source

https://nanostring.com/products/cosmx-spatial-molecular-imager/ffpe-dataset/nsclc-ffpe-dataset/

See Also

```
SpaTopic_inference,Seurat5obj_to_SpaTopic
```

```
Seurat5obj_to_SpaTopic
```

Convert a Seurat v5 object as the input of 'SpaTopic'

Description

Prepare 'SpaTopic' input from one Seurat v5 object

Usage

```
Seurat5obj_to_SpaTopic(object, group.by, image = "image1")
```

Arguments

object Seurat v5 object

group.by character. The name of the column that contains celltype information in the

Seurat object.

image character. The name of the image. Default is "image1".

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Value

Return a data frame as the input of 'SpaTopic'

See Also

lung5

Examples

```
## nano.obj is a Seurat v5 object
#dataset<-Seurat5obj_to_SpaTopic(object = nano.obj,
# group.by = "predicted.annotation.l1",image = "image1")
## Expect output
data("lung5")</pre>
```

SpaTopic-Package

'SpaTopic' R package

Description

The 'SpaTopic' R package is centered around the 'SpaTopic' algorithm to infer the spatial tissue architectures from multiplexed images.

Details

The package implements a Collapsed Gibbs sampling algorithm to infer topics, corresponding to distinct tissue microenvironments across multiple tissue images. Without obtaining the cell neighborhood info of every single cell, 'SpaTopic' runs much faster than other KNN-based methods on large-scale images.

The main functions in the 'SpaTopic' package

- Prepare input Seurat5obj_to_SpaTopic
- Model Inference SpaTopic_inference

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SpaTopic_inference

'SpaTopic': fast topic inference to identify tissue architecture in multiplexed images

Description

This is the main function of 'SpaTopic', implementing a Collapsed Gibbs Sampling algorithm to learn topics, which referred to different tissue microenvironments, across multiple multiplexed tissue images. The function takes cell labels and coordinates on tissue images as input, and returns the inferred topic labels for every cell, as well as topic contents, a distribution over celltypes. The function recovers spatial tissue architectures across images, as well as indicating cell-cell interactions in each domain.

Usage

```
SpaTopic_inference(
  tissue,
  ntopics,
  sigma = 50,
  region_radius = 400,
  kneigh = 5,
  npoints_selected = 1,
  ini_LDA = TRUE,
  ninit = 10,
  niter_init = 100,
  beta = 0.05,
  alpha = 0.01,
  trace = FALSE,
  seed = 123,
  thin = 20,
  burnin = 1000,
  niter = 200,
  display_progress = TRUE,
  do.parallel = FALSE,
  n.cores = 1,
  axis = "2D"
)
```

Arguments

tissue

(Required). A data frame or a list of data frames. One for each image. Each row represent a cell with its image ID, X, Y coordinates on the image, celltype, with column names (image, X, Y, type), respectively. You may add another column Y2 for 3D tissue image.

ntopics

(Required). Number of topics. Topics will be obtained as distributions of cell types.

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sigma Default is 50. The lengthscale of the Nearest-neighbor Exponential Kernel. Sigma controls the strength of decay of correlation with distance in the kernel function. Please check the paper for more information. Need to be adjusted based on the image resolution region_radius Default is 400. The radius for each grid square when sampling region centers for each image. Need to be adjusted based on the image resolution and pattern complexity. kneigh Default is 5. Only consider the top 5 closest region centers for each cell. npoints_selected Default is 1. Number of points sampled for each grid square when sampling region centers for each image. Used with region_radius. ini_LDA Default is TRUE. Use warm start strategy for initialization and choose the best one to continue. If 0, it simply uses the first initialization. ninit Default is 10. Number of initialization. Only retain the initialization with the highest log likelihood (perplexity). niter_init Default is 100. Warm start with 100 iterations in the Gibbs sampling during initialization. Default is 0.05. A hyperparameter to control the sparsity of topic content (topicbeta celltype) matrix Beta. A smaller value introduces more sparse in Beta. alpha Default is 0.01. A hyperparameter to control the sparsity of document (region) content (region-topic) matrix Theta. For our application, we keep it very small for the sparsity in Theta. Default is FALSE. Compute and save log likelihood, Ndk, Nwk for every posterior trace samples. Useful when you want to use DIC to select number of topics, but it is time consuming to compute the likelihood for every posterior samples. Default is 123. Random seed. seed thin Default is 20. Key parameter in Gibbs sampling. Collect a posterior sample for every thin=20 iterations. burnin Default is 1000. Key parameter in Gibbs sampling. Start to collect posterior samples after 1000 iterations. You may increase the number of iterations for burn-in for highly complex tissue images. niter Default is 200. Key parameter in Gibbs sampling. Number of posterior samples collected for model inference. display_progress Default is TRUE. Display the progress bar. do.parallel Default is FALSE. Use parallel computing through R package for each. Default is 1. Number of cores used in parallel computing. n.cores

Default is "2D". You may switch to "3D" for 3D tissue images. However, the

Value

axis

Return a gibbs.res-class object. A list of outputs from Gibbs sampling.

model inference for 3D tissue is still under test.

stratified_sampling_sf

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See Also

```
gibbs.res-class
```

Examples

```
stratified_sampling_sf
```

Spatially stratified random sample points from an image.

Description

Spatially stratified random sample points from an image by R package sf

Usage

```
stratified_sampling_sf(
  points,
  cellsize = c(600, 600),
  num_samples_per_stratum = 1
)
```

Arguments

```
points a data frame contains all points in a image with X, Y coordinates.

cellsize a vector of length 2 contains the size of each grid square. Default c(600,600).

num_samples_per_stratum

number of point selected from each grid square. Default 1.
```

Value

Return a vector contains index of sampled points.

Examples

```
data("lung5")
pt_idx<-stratified_sampling_sf(lung5, cellsize = c(600,600))</pre>
```

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