

# Package ‘lisaClust’

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**Type** Package

**Title** lisaClust: Clustering of Local Indicators of Spatial Association

**Version** 1.12.3

**Description** lisaClust provides a series of functions to identify and visualise regions of tissue where spatial associations between cell-types is similar. This package can be used to provide a high-level summary of cell-type colocalization in multiplexed imaging data that has been segmented at a single-cell resolution.

**License** GPL (>=2)

**biocViews** SingleCell, CellBasedAssays, Spatial

**Encoding** UTF-8

**Depends** R (>= 4.0)

**VignetteBuilder** knitr

**BugReports** <https://github.com/ellisppatrick/lisaClust/issues>

**URL** <https://ellisppatrick.github.io/lisaClust/>,

<https://github.com/ellisppatrick/lisaClust>

**Imports** ggplot2, class, concaveman, grid, BiocParallel,  
spatstat.explore, spatstat.geom, BiocGenerics, S4Vectors,  
methods, spicyR, purrr, stats, data.table, dplyr, tidyr,  
SingleCellExperiment, SpatialExperiment, SummarizedExperiment,  
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**Suggests** BiocStyle, knitr, rmarkdown, SpatialDatasets

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<i>hatchingPlot</i>	<i>hatchingPlot</i>
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## Description

The *hatchingPlot()* function is used to create hatching patterns for representing spatial regions and cell-types.

The hatching geom is used to create hatching patterns for representation of spatial regions.

## Usage

```

hatchingPlot(  

  data,  

  useImages = NULL,  

  region = "region",  

  imageID = "imageID",  

  cellType = "cellType",  

  spatialCoords = c("x", "y"),  

  window = "concave",  

  line.spacing = 21,  

  hatching.colour = 1,  

  nbp = 50,  

  window.length = NULL  

)  
  

geom_hatching(  

  mapping = NULL,  

  data = NULL,  

  stat = "identity",  

  position = "identity",

```

```

na.rm = FALSE,
show.legend = NA,
inherit.aes = TRUE,
line.spacing = 21,
hatching.colour = 1,
window = "concave",
window.length = NULL,
nbp = 250,
line.width = 1,
...
)

```

## Arguments

<code>data</code>	The data to be displayed in this layer. There are three options: If <code>NULL</code> , the default, the data is inherited from the plot data as specified in the call to <code>ggplot()</code> . A <code>data.frame</code> , or other object, will override the plot data. All objects will be fortified to produce a data frame. See <code>fortify()</code> for which variables will be created. A function will be called with a single argument, the plot data. The return value must be a <code>data.frame</code> , and will be used as the layer data. A function can be created from a formula (e.g. <code>~ head(.x, 10)</code> ).
<code>useImages</code>	A vector of images to plot.
<code>region</code>	The region column to plot.
<code>imageID</code>	The <code>imageIDs</code> column if using <code>data.frame</code> or <code>SingleCellExperiment</code> .
<code>cellType</code>	The <code>cellType</code> column if using <code>data.frame</code> or <code>SingleCellExperiment</code> .
<code>spatialCoords</code>	The spatial coordinates columns if using <code>data.frame</code> or <code>SingleCellExperiment</code> .
<code>window</code>	Should the window around the regions be 'square', 'convex' or 'concave'.
<code>line.spacing</code>	A integer indicating the spacing between hatching lines.
<code>hatching.colour</code>	A colour for the hatching.
<code>nbp</code>	An integer tuning the granularity of the grid used when defining regions
<code>window.length</code>	A tuning parameter for controlling the level of concavity when estimating concave windows.
<code>mapping</code>	Set of aesthetic mappings created by <code>aes()</code> or <code>aes_()</code> . If specified and <code>inherit.aes = TRUE</code> (the default), it is combined with the default mapping at the top level of the plot. You must supply <code>mapping</code> if there is no plot mapping.
<code>stat</code>	The statistical transformation to use on the data for this layer as a string.
<code>position</code>	adjustment, either as a string, or the result of a call to a position adjustment function.
<code>na.rm</code>	If <code>FALSE</code> , the default, missing values are removed with a warning. If <code>TRUE</code> , missing values are silently removed.
<code>show.legend</code>	logical. Should this layer be included in the legends? <code>NA</code> , the default, includes if any aesthetics are mapped. <code>FALSE</code> never includes, and <code>TRUE</code> always includes. It can also be a named logical vector to finely select the aesthetics to display.

<code>inherit.aes</code>	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <code>borders()</code> .
<code>line.width</code>	A numeric controlling the width of the hatching lines
<code>...</code>	Other arguments passed on to <code>layer()</code> . These are often aesthetics, used to set an aesthetic to a fixed value, like <code>colour = "red"</code> or <code>size = 3</code> . They may also be parameters to the paired geom/stat.

## Value

A ggplot object  
A ggplot geom

## Examples

```
## Generate toy data
set.seed(51773)
x <- round(c(
  runif(200), runif(200) + 1, runif(200) + 2, runif(200) + 3,
  runif(200) + 3, runif(200) + 2, runif(200) + 1, runif(200)
), 4) * 100
y <- round(c(
  runif(200), runif(200) + 1, runif(200) + 2, runif(200) + 3,
  runif(200), runif(200) + 1, runif(200) + 2, runif(200) + 3
), 4) * 100
cellType <- factor(paste("c", rep(rep(c(1:2), rep(200, 2)), 4), sep = ""))
imageID <- rep(c("s1", "s2"), c(800, 800))
cells <- data.frame(x, y, cellType, imageID)
cells <- SingleCellExperiment::SingleCellExperiment(colData = cells)

## Generate regions
cells <- lisaClust(cells, k = 2)

## Plot regions
hatchingPlot(cells)

## Generate toy data
set.seed(51773)
library(ggplot2)
x <- round(c(
  runif(200), runif(200) + 1, runif(200) + 2, runif(200) + 3,
  runif(200) + 3, runif(200) + 2, runif(200) + 1, runif(200)
), 4) * 100
y <- round(c(
  runif(200), runif(200) + 1, runif(200) + 2, runif(200) + 3,
  runif(200), runif(200) + 1, runif(200) + 2, runif(200) + 3
), 4) * 100
cellType <- factor(paste("c", rep(rep(c(1:2), rep(200, 2)), 4), sep = ""))
imageID <- rep(c("s1", "s2"), c(800, 800))
cells <- data.frame(x, y, cellType, imageID)
## Generate regions
```

```

cells <- lisaClust(cells, k = 2)

# Plot the regions with geom_hatching()
ggplot(
  cells, aes(x = x, y = y, colour = cellType, region = region)
) +
  geom_point() +
  facet_wrap(~imageID) +
  geom_hatching()

```

**inhomLocalK***Calculate the inhomogenous local K function.***Description**

Calculate the inhomogenous local K function.

**Usage**

```

inhomLocalK(
  data,
  Rs = c(20, 50, 100, 200),
  sigma = 10000,
  window = "convex",
  window.length = NULL,
  minLambda = 0.05,
  lisaFunc = "K"
)

```

**Arguments**

<code>data</code>	The data.
<code>Rs</code>	A vector of the radii that the measures of association should be calculated.
<code>sigma</code>	A numeric variable used for scaling when fitting inhomogeneous L-curves.
<code>window</code>	Should the window around the regions be 'square', 'convex' or 'concave'.
<code>window.length</code>	A tuning parameter for controlling the level of concavity.
<code>minLambda</code>	Minimum value for density for scaling when fitting inhomogeneous L-curves.
<code>lisaFunc</code>	Either "K" or "L" curve.

**Value**

A matrix of LISA curves

## Examples

```
library(spicyR)
# Read in data
isletFile <- system.file("extdata", "isletCells.txt.gz", package = "spicyR")
cells <- read.table(isletFile, header = TRUE)
cells$x <- cells$AreaShape_Center_X
cells$y <- cells$AreaShape_Center_Y
cells$cellType <- as.factor(sample(
  c("big", "medium", "small"),
  length(cells$AreaShape_Center_Y),
  replace = TRUE
))
cells$cellID <- as.factor(cells$ObjectNumber)

inhom <- inhomLocalK(cells[1:100, ])
```

**lisa**

*Generate local indicators of spatial association*

## Description

Generate local indicators of spatial association

## Usage

```
lisa(
  cells,
  Rs = NULL,
  BPPARAM = BiocParallel::SerialParam(),
  window = "convex",
  window.length = NULL,
  whichParallel = "imageID",
  sigma = NULL,
  lisaFunc = "K",
  minLambda = 0.05,
  fast = TRUE,
  spatialCoords = c("x", "y"),
  cellType = "cellType",
  imageID = "imageID"
)
```

## Arguments

cells	A SingleCellExperiment, SpatialExperiment or data frame that contains at least the variables x and y, giving the coordinates of each cell, imageID and cellType.
Rs	A vector of the radii that the measures of association should be calculated.

BPPARAM	A BiocParallelParam object.
window	Should the window around the regions be 'square', 'convex' or 'concave'.
window.length	A tuning parameter for controlling the level of concavity when estimating concave windows.
whichParallel	Should the function use parallelization on the imageID or the cellType.
sigma	A numeric variable used for scaling when fitting inhomogeneous L-curves.
lisaFunc	Either "K" or "L" curve.
minLambda	Minimum value for density for scaling when fitting inhomogeneous L-curves.
fast	A logical describing whether to use a fast approximation of the inhomogeneous local L-curves.
spatialCoords	The columns which contain the x and y spatial coordinates.
cellType	The column which contains the cell types.
imageID	The column which contains image identifiers.

### Value

A matrix of LISA curves

### Examples

```
library(spicyR)
library(SingleCellExperiment)
# Read in data
isletFile <- system.file("extdata", "isletCells.txt.gz", package = "spicyR")
cells <- read.table(isletFile, header = TRUE)
cellExp <- SingleCellExperiment(
  assay = list(intensities = t(cells[, grepl(names(cells), pattern = "Intensity_")])),
  colData = cells[, !grepl(names(cells), pattern = "Intensity_")]
)

# Cluster cell types
markers <- t(assay(cellExp, "intensities"))
kM <- kmeans(markers, 8)
colData(cellExp)$cluster <- paste("cluster", kM$cluster, sep = "")

# Generate LISA
lisaCurves <- lisa(
  cellExp,
  spatialCoords = c("Location_Center_X", "Location_Center_Y"),
  cellType = "cluster", imageID = "ImageNumber"
)

# Cluster the LISA curves
kM <- kmeans(lisaCurves, 2)
```

---

lisaClust	<i>Use k-means clustering to cluster local indicators of spatial association. For other clustering use lisa.</i>
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---

## Description

Use k-means clustering to cluster local indicators of spatial association. For other clustering use lisa.

## Usage

```
lisaClust(
  cells,
  k = 2,
  Rs = NULL,
  spatialCoords = c("x", "y"),
  cellType = "cellType",
  imageID = "imageID",
  regionName = "region",
  BPPARAM = BiocParallel::SerialParam(),
  window = "convex",
  window.length = NULL,
  whichParallel = "imageID",
  sigma = NULL,
  lisaFunc = "K",
  minLambda = 0.05,
  fast = TRUE
)
```

## Arguments

cells	A SingleCellExperiment, SpatialExperiment or data frame that contains at least the variables x and y, giving the coordinates of each cell, imageID and cellType.
k	The number of regions to cluster.
Rs	A vector of the radii that the measures of association should be calculated.
spatialCoords	The columns which contain the x and y spatial coordinates.
cellType	The column which contains the cell types.
imageID	The column which contains image identifiers.
regionName	The output column for the lisaClust regions.
BPPARAM	A BiocParallelParam object.
window	Should the window around the regions be 'square', 'convex' or 'concave'.
window.length	A tuning parameter for controlling the level of concavity when estimating concave windows.
whichParallel	Should the function use parallelization on the imageID or the cellType.

<code>sigma</code>	A numeric variable used for scaling when fitting inhomogeneous L-curves.
<code>lisaFunc</code>	Either "K" or "L" curve.
<code>minLambda</code>	Minimum value for density for scaling when fitting inhomogeneous L-curves.
<code>fast</code>	A logical describing whether to use a fast approximation of the inhomogeneous local L-curves.

**Value**

A matrix of LISA curves

**Examples**

```
library(spicyR)
library(SingleCellExperiment)
# Read in data
isletFile <- system.file("extdata", "isletCells.txt.gz", package = "spicyR")
cells <- read.table(isletFile, header = TRUE)
cellExp <- SingleCellExperiment(
  assay = list(intensities = t(cells[, grepl(names(cells), pattern = "Intensity_")])),
  colData = cells[, !grepl(names(cells), pattern = "Intensity_")]
)

# Cluster cell types
markers <- t(assay(cellExp, "intensities"))
kM <- kmeans(markers, 8)
colData(cellExp)$cluster <- paste("cluster", kM$cluster, sep = "")

# Generate LISA
cellExp <- lisaClust(cellExp,
  k = 2,
  imageID = "ImageNumber",
  cellType = "cluster",
  spatialCoords = c("Location_Center_X", "Location_Center_Y")
)
```

regionMap

*Plot heatmap of cell type enrichment for lisaClust regions*

**Description**

Plot heatmap of cell type enrichment for lisaClust regions

**Usage**

```
regionMap(
  cells,
  type = "bubble",
```

```

cellType = "cellType",
region = "region",
limit = c(0.33, 3),
...
)

```

### Arguments

cells	SingleCellExperiment, SpatialExperiment or data.frame
type	Make a "bubble" or "heatmap" plot.
cellType	The column storing the cell types
region	The column storing the regions
limit	limits to the lower and upper relative frequencies
...	Any arguments to be passed to the pheatmap package

### Value

A bubble plot or heatmap

### Examples

```

set.seed(51773)
x <- round(c(
  runif(200), runif(200) + 1, runif(200) + 2, runif(200) + 3,
  runif(200) + 3, runif(200) + 2, runif(200) + 1, runif(200)
), 4) * 100
y <- round(c(
  runif(200), runif(200) + 1, runif(200) + 2, runif(200) + 3,
  runif(200), runif(200) + 1, runif(200) + 2, runif(200) + 3
), 4) * 100
cellType <- factor(paste("c", rep(rep(c(1:2), rep(200, 2)), 4), sep = ""))
imageID <- rep(c("s1", "s2"), c(800, 800))

cells <- data.frame(x, y, cellType, imageID)

cells <- lisaClust(cells, k = 2)

regionMap(cells)

```

### Description

Region scale constructor.

**Usage**

```
scale_region(aesthetics = "region", ..., guide = "legend")

scale_region_manual(..., values)
```

**Arguments**

aesthetics	The names of the aesthetics that this scale works with
...	Arguments passed on to discrete_scale
guide	A function used to create a guide or its name. See guides() for more info.
values	a set of aesthetic values to map data values to. If this is a named vector, then the values will be matched based on the names. If unnamed, values will be matched in order (usually alphabetical) with the limits of the scale. Any data values that don't match will be given na.value.

**Value**

a ggplot guide

**Examples**

```
library(spicyR)
## Generate toy data
set.seed(51773)
x <- round(c(
  runif(200), runif(200) + 1, runif(200) + 2, runif(200) + 3,
  runif(200) + 3, runif(200) + 2, runif(200) + 1, runif(200)
), 4) * 100
y <- round(c(
  runif(200), runif(200) + 1, runif(200) + 2, runif(200) + 3,
  runif(200), runif(200) + 1, runif(200) + 2, runif(200) + 3
), 4) * 100
cellType <- factor(paste("c", rep(rep(c(1:2), rep(200, 2)), 4), sep = ""))
imageID <- rep(c("s1", "s2"), c(800, 800))
cells <- data.frame(x, y, cellType, imageID)
cells <- SingleCellExperiment::SingleCellExperiment(colData = cells)

## Generate regions
cells <- lisaClust(cells, k = 2)

# Plot the regions with hatchingPlot()
hatchingPlot(cells) +
  scale_region_manual(
    values = c(1, 4), labels = c("Region A", "Region B"),
    name = "Regions"
  )
```

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