# **Package**

#### December 5, 2023

Title Granularity-Based Spatially Variable Genes Identifications

Version 0.0.2

#### **Description**

Identifying spatially variable genes is critical in linking molecular cell functions with tissue phenotypes. This package implemented a granularity-based dimension-agnostic tool for the identification of spatially variable genes. The detailed description of this method is available at Wang, J. and Li, J. et al. 2023 (Wang, J. and Li, J. (2023), <doi:10.1038/s41467-023-43256-5>).

License GPL (>= 2)
Encoding UTF-8
Imports Matrix,
sparseMatrixStats,
fitdistrplus,
RANN,
spam
Suggests knitr,
rmarkdown

RoxygenNote 7.2.1

## **R** topics documented:

GrabSVG		A Granularity-Based Approach to identify Spatially Variable Genes																		
Index																				4
	SpFilter		• • •							٠			•		٠	 •	•	•	 ٠	3
	LoadSpatial																			
	GrabSVG															 				1

#### **Description**

This function is designed to identify spatially variable genes through a granularity-based approach.

### Usage

```
GrabSVG(Coords, ExpMat_Sp, D_1 = 1.0, D_2 = 3.0,
Exp_Norm = TRUE, Coords_Norm_Method = c("Sliced", "Overall", "None"))
```

2 LoadSpatial

#### **Arguments**

Coords A M x D matrix representing D-dimensional coordinates for M spots

ExpMat\_Sp A sparse, N x M expression matrix in dgCMatrix class with N genes and M

spots

D\_1 Size of the small patchD\_2 Size of the big patch

Exp\_Norm A Boolean value indicating whether the expression matrix should be normalized

Coords\_Norm\_Method

Normalization method for the coordinates matrix, which can be "None", "Sliced",

or "Overall".

#### **Details**

This function utilizes a MxD matrix (Coords) representing D-dimensional coordinates with M spots and a sparse, NxM expression matrix (ExpMat\_Sp) with N genes and M spots.

#### Value

A data frame with the name of genes and corresponding p-values.

#### **Examples**

```
Coords <- expand.grid(1:100,1:100, 1:3)

RandFunc <- function(n) floor(10 * stats::rbeta(n, 1, 5))

Raw_Exp <- Matrix::rsparsematrix(nrow = 10^4, ncol = 3*10^4, density = 0.0001, rand.x = RandFunc)

Filtered_ExpMat <- SpFilter(Raw_Exp)

rownames(Filtered_ExpMat) <- paste0("Gene_", 1:nrow(Filtered_ExpMat))

P_values <- GrabSVG(Coords, Filtered_ExpMat)
```

LoadSpatial

Loading data from a Seurat object or a data frame.

#### Description

A function to load and filter data from a Seurat object or a data frame.

#### Usage

```
LoadSpatial(InputData, Dimension = 2)
```

#### **Arguments**

InputData A Seurat spatial object or a M x (D + N) data matrix representing the D-dimensional

coordinates and expressions of N genes on M spots. The coordinates should be

placed at the first D columns

Dimension The dimension of coordinates

SpFilter 3

#### Value

A list of two data frame:

Coords A M x D matrix representing D-dimensional coordinates for M spots

ExpMatrix A sparse, N x M expression matrix in dgCMatrix class with N genes and M

spots

SpFilter A function for filtering low expressed genes

#### **Description**

A function for filtering low expressed genes

#### Usage

```
SpFilter(ExpMat_Sp, Threshold = 5)
```

#### **Arguments**

ExpMat\_Sp A sparse, N x M expression matrix in dgCMatrix class with N genes and M

spots

Threshold A threshold set to filter out genes with a total read count below this specified

value

#### Value

A sparse expression matrix in dgCMatrix class

#### **Examples**

```
# create a sparse expression matrix
Raw_ExpMat <- Matrix::rsparsematrix(nrow = 10000, ncol = 2000,
density = 0.01, rand.x = function(n) rpois(n, 15))
Filtered_ExpMat <- SpFilter(Raw_ExpMat)</pre>
```

# Index

GrabSVG, 1
LoadSpatial, 2

SpFilter, 3