Package 'jazzPanda'

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Description This package contains the function to find marker genes for image-based spatial transcriptomics data. There are functions to create spatial vectors from the cell and transcript coordiantes, which are passed as inputs to find marker genes. Marker genes are detected for every cluster by two approaches. The first approach is by permutation testing, which is implemented in parallel for finding marker genes for one sample study. The other approach is to build a linear model for every gene. This approach can account for multiple samples and backgound noise.

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URL https://github.com/phipsonlab/jazzPanda,

https://bhuvad.github.io/jazzPanda/

BugReports https://github.com/phipsonlab/jazzPanda/issues

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Contents

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jazzPanda-package

jazzPanda: A hybrid approach to find spatially relevant marker genes in image-based spatial transcriptomics data

Description

jazzPanda pacakge provides hybrid approaches to prioritize marker genes that uses the spatial coordinates of gene detections and cells making up clusters. We propose a binning approach get_vectors that summarises the number of genes and cells within a cluster as spatial vectors. We have developed two approaches to detect and prioritize marker genes. The first approach compute_permp is based on correlation coefficients between genes and cluster spatial vectors, where significance of the marker genes are assessed through permutation. The second approach lasso_markers is based on lasso regularisation and linear modeling of our defined spatial vectors. This second approach is more flexible and can account for multiple samples and background noise.

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.check_binning

helper function to check the input of binning

Description

helper function to check the input of binning

Usage

.check_binning(bin_param, bin_type, range_list)

Arguments

bin_param	A numeric vector indicating the size of the bin. If the bin_type is "square" or "rectangle", this will be a vector of length two giving the numbers of rectangular quadrats in the x and y directions. If the bin_type is "hexagonal", this will be a number giving the side length of hexagons. Positive numbers only.
bin_type	A string indicating which bin shape is to be used for vectorization. One of "square" (default), "rectangle", or "hexagon".
range_list	A named list of spatial ranges for each sample. Each element should be a list with two components: w_x and w_y , which are numeric vectors of length 2 specifying the x- and y-axis ranges (e.g., from cell or transcript coordinates). The range is calculated with 5 within the window.

Value

the length of total bins

.check_valid_input helper function to check the inputs passed to marker detection function

Description

helper function to check the inputs passed to marker detection function

Usage

```
.check_valid_input(
  gene_mt,
  cluster_mt,
  sample_names,
  n_fold = 10,
  background = NULL
)
```

Arguments

gene_mt	A matrix contains the transcript count in each grid. Each row refers to a grid, and each column refers to a gene. The column names must be specified and refer to the genes. This can be the output from the function get_vectors.
cluster_mt	A matrix contains the number of cells in a specific cluster in each grid. Each row refers to a grid, and each column refers to a cluster. The column names must be specified and refer to the clusters. Please do not assign integers as column names. This can be the output from the function get_vectors.
sample_names	A vector specifying the names for the samples.
n_fold	Optional. A positive number giving the number of folds used for cross valida- tion. This parameter will pass to cv.glmnet to calculate a penalty term for every gene.
background	Optional. A matrix providing the background information. Each row refers to a grid, and each column refers to one category of background information. Number of rows must equal to the number of rows in gene_mt and cluster_mt. Can be obtained by only providing coordinates matrices cluster_info. to function get_vectors.

Value

a list of two matrices with the following components

n_clusters	Number of clusters
cluster_names	a vector of strings giving the name of the clusters

.check_valid_names helper function to check the names of gene/cluster/sample

Description

helper function to check the names of gene/cluster/sample

Usage

```
.check_valid_names(x, x_name)
```

Arguments

х	A character vector to check naming
x_name	A name specifying the type of x, for message purpose only

Value

A character vector of same length with valid names

.compute_observation Compute observation statistic for permutation framework

Description

Compute observation statistic for permutation framework

```
.compute_observation(
    x,
    cluster_info,
    correlation_method,
    n_cores,
    test_genes,
    bin_type,
    bin_param,
    use_cm
)
```

X	a named list (of transcript detection coordinates) or SingleCellExperiment or SpatialExperiment or SpatialFeatureExperiment object. If a named list is pro- vided, every list element is a dataframe containing the transcript detection co- ordinates and column names must include "feature_name" (gene name), "x" (x coordinate), "y" (y coordinate). The list names must match samples in clus- ter_info.
cluster_info	A dataframe/matrix containing the centroid coordinates, cluster label and sample for each cell.The column names must include "x" (x coordinate), "y" (y coordinate), "cluster" (cluster label) and "sample" (sample). It is strongly recommended to use syntactically valid names for columns clusters and samples. If invalid names are detected, the function make.names will be employed to generate valid names. A message will also be displayed to indicate this change.
correlation_me	thod
	A parameter pass to cor, indicating which correlation coefficient is to be com- puted. One of "pearson" (default), "kendall", or "spearman": can be abbreviated.
n_cores	A positive number specifying number of cores used for parallelizing permuta- tion testing. Default is one core (sequential processing).
test_genes	A vector of strings giving the name of the genes you want to test correlation for.
bin_type	A string indicating which bin shape is to be used for vectorization. One of "square" (default), "rectangle", or "hexagon".
bin_param	A numeric vector indicating the size of the bin. If the bin_type is "square" or "rectangle", this will be a vector of length two giving the numbers of rectangular quadrats in the x and y directions. If the bin_type is "hexagonal", this will be a number giving the side length of hexagons. Positive numbers only.
use_cm	A boolean value that specifies whether to create spatial vectors for genes using the count matrix and cell coordinates instead of the transcript coordinates when both types of information are available. The default setting is FALSE.

Value

A named list with the following components	
obs.stat	A matrix contains the observation statistic for every gene and every cluster. Each row refers to a gene, and each column refers to a cluster
gene_mt	contains the transcript count in each grid. Each row refers to a grid, and each column refers to a gene.

.compute_permutation Compute permutation statistics for permutation framework

Description

Compute permutation statistics for permutation framework

.convert_data

Usage

```
.compute_permutation(
   cluster_info,
   perm.size = 1000,
   correlation_method = "pearson",
   bin_type,
   bin_param,
   n_cores = 1,
   gene_mt,
   cluster_names
)
```

Arguments

cluster_info	A dataframe/matrix containing the centroid coordinates and cluster label for each cell.The column names should include "x" (x coordinate), "y" (y coordinate), and "cluster" (cluster label).
perm.size	A positive number specifying permutation times
correlation_me	thod
	A parameter pass to cor , indicating which correlation coefficient is to be com- puted. One of "pearson" (default), "kendall", or "spearman": can be abbreviated.
bin_type	A string indicating which bin shape is to be used for vectorization. One of "square" (default), "rectangle", or "hexagon".
bin_param	A numeric vector indicating the size of the bin. If the bin_type is "square" or "rectangle", this will be a vector of length two giving the numbers of rectangular quadrats in the x and y directions. If the bin_type is "hexagonal", this will be a number giving the side length of hexagons. Positive numbers only.
n_cores	A positive number specifying number of cores used for parallelizing permuta- tion testing. Default is one core (sequential processing).
gene_mt	A matrix contains the transcript count in each grid. Each row refers to a grid, and each column refers to a gene.
cluster_names	A list of strings giving the name and order of the clusters

Value

A matrix with permutation statistics

.convert_data	ConvertSingleCell Experiment/Spatial Experiment/Spatial Feature Experiment
	objects to list object for jazzPanda.

Description

This function takes an object of class SingleCellExperiment, SpatialExperiment or SpatialFeature-Experimentreturns and returns a list object that is expected for the get_vector functions.

Usage

```
.convert_data(x, sample_names, test_genes)
```

Arguments

х	a SingleCellExperiment or SpatialExperiment or SpatialFeatureExperiment object
sample_names	a vector of strings giving the sample names
test_genes	A vector of strings giving the name of the genes you want to create gene vector.

Value

outputs a list object with the following components

trans_lst	A list of named dataframes. Each dataframe refers to one sample and shows
	the transcript detection coordinates for each gene. The name matches the input
	sample_names
cm_lst	A list of named dataframes containing the count matrix for each sample. The name matches the input sample_names

.create_cor_mg_result Create a marker gene result object for correlation approach

Description

This function creates a structured output object named 'cor_mg_result' for storing the permutation results. The object contains three matrices:

Usage

```
.create_cor_mg_result(obs.stat, perm.pval, perm.pval.adj)
```

Arguments

obs.stat	A matrix containing the correlation coefficients for each pair of genes and cluster vectors.
perm.pval	A matrix containing the raw permutation p-value for each pair of genes and cluster.
perm.pval.adj	A matrix containing the adjusted permutation p-value for each pair of genes and cluster.

Value

An S3 object of class 'cor_mg_result' which includes three matrices.

.create_lm_mg_result Create a marker gene result object for linear modelling approach

Description

This function creates a structured output object named 'glm_mg_result' for storing the marker gene results. The object contains two data frames: top results and full results.

Usage

```
.create_lm_mg_result(top_result, full_result)
```

Arguments

top_result	A data frame containing top results.
full_result	A data frame containing full results.

Value

An S3 object of class 'glm_mg_result' which includes both results data frames.

.get_cluster_vectors Create spatial vectors for clusters

Description

Create spatial vectors for clusters

```
.get_cluster_vectors(
    cluster_info,
    bin_length,
    bin_type,
    bin_param,
    range_list,
    sample_names
)
```

cluster_info	A dataframe/matrix containing the centroid coordinates, cluster label and sample for each cell.The column names must include "x" (x coordinate), "y" (y coordinate), "cluster" (cluster label) and "sample" (sample).
bin_length	A positive integer giving the length of total bins
bin_type	A string indicating which bin shape is to be used for vectorization. One of "square" (default), "rectangle", or "hexagon".
bin_param	A numeric vector indicating the size of the bin. If the bin_type is "square" or "rectangle", this will be a vector of length two giving the numbers of rectangular quadrats in the x and y directions. If the bin_type is "hexagonal", this will be a number giving the side length of hexagons. Positive numbers only.
range_list	A named list of spatial ranges for each sample. Each element should be a list with two components: w_x and w_y , which are numeric vectors of length 2 specifying the x- and y-axis ranges (e.g., from cell or transcript coordinates). The range is calculated with 5 within the window.
sample_names	a vector of strings giving the sample names

Value

a matrix contains the cell count in each grid. Each row refers to a grid, and each column refers to a cluster.

.get_gene_vectors_cm Create spatial vectors for genes from count matrix and cell coordinates

Description

This function will build gene vectors with count matrix and cell locations

```
.get_gene_vectors_cm(
   cluster_info,
   cm_lst,
   bin_type,
   bin_param,
   test_genes,
   range_list
)
```

cluster_info	A dataframe/matrix containing the centroid coordinates, cluster label and sample for each cell.The column names must include "x" (x coordinate), "y" (y coordinate), "cluster" (cluster label) and "sample" (sample).
cm_lst	A list of named matrices containing the count matrix for each sample The name must match the sample column in cluster_info. If this input is provided, the cluster_info must be specified and contain an additional column "cell_id" to link cell location and count matrix. Default is NULL.
bin_type	A string indicating which bin shape is to be used for vectorization. One of "square" (default), "rectangle", or "hexagon".
bin_param	A numeric vector indicating the size of the bin. If the bin_type is "square" or "rectangle", this will be a vector of length two giving the numbers of rectangular quadrats in the x and y directions. If the bin_type is "hexagonal", this will be a number giving the side length of hexagons. Positive numbers only.
test_genes	A vector of strings giving the name of the genes you want to test. This will be used as column names for one of the result matrix gene_mt.
range_list	A named list of spatial ranges for each sample. Each element should be a list with two components: w_x and w_y , which are numeric vectors of length 2 specifying the x- and y-axis ranges (e.g., from cell or transcript coordinates). The range is calculated with 5 within the window.

Value

a matrix contains the transcript count in each grid. Each row refers to a grid, and each column refers to a gene.

.get_gene_vectors_tr Create spatial vectors for genes from transcript coordinates

Description

This function will build gene vectors based on the transcript coordinates of every gene

```
.get_gene_vectors_tr(
   trans_lst,
   test_genes,
   bin_type,
   bin_param,
   bin_length,
   range_list
)
```

trans_lst	If specified, it is a list of named dataframes. Each dataframe refers to one sample and shows the transcript detection coordinates for each gene. Optional parame- ter.
test_genes	A vector of strings giving the name of the genes you want to test. This will be used as column names for one of the result matrix gene_mt.
bin_type	A string indicating which bin shape is to be used for vectorization. One of "square" (default), "rectangle", or "hexagon".
bin_param	A numeric vector indicating the size of the bin. If the bin_type is "square" or "rectangle", this will be a vector of length two giving the numbers of rectangular quadrats in the x and y directions. If the bin_type is "hexagonal", this will be a number giving the side length of hexagons. Positive numbers only.
bin_length	A positive integer giving the length of total bins
range_list	A named list of spatial ranges for each sample. Each element should be a list with two components: w_x and w_y, which are numeric vectors of length 2 specifying the x- and y-axis ranges (e.g., from cell or transcript coordinates). The range is calculated with 5 within the window.

Value

a matrix contains the transcript count in each grid. Each row refers to a grid, and each column refers to a gene.

.get_lasso_coef help function to get lasso coefficient for every cluster for a given model

Description

help function to get lasso coefficient for every cluster for a given model

```
.get_lasso_coef(
    i_gene,
    gene_mt,
    vec_cluster,
    cluster_names,
    n_fold = 10,
    n_samples,
    sample_names
)
```

BUFFER_FRAC

Arguments

i_gene	Name of the current gene
gene_mt	A matrix contains the transcript count in each grid. Each row refers to a grid, and each column refers to a gene. The column names must be specified and refer to the genes. This can be the output from the function get_vectors.
vec_cluster	A matrix of the spatial vectors for clusters.
cluster_names	A vector of strings giving the name of clusters
n_fold	Optional. A positive number giving the number of folds used for cross validation. This parameter will pass to $cv.glmnet$ to calculate a penalty term for every gene.
n_samples	A positive number giving the number samples
sample_names	A vector specifying the names for the sample

Value

a list of two matrices with the following components

coef_df	A matrix giving the lasso coefficient of each cluster
lambda.1se	the lambda.1se value of best fitted model

BUFFER_FRAC	Default padding parameter for function get_vectors()
-------------	--

Description

A numeric value giving the padding offset proportion when defining the range for x and y coordinates

Usage

BUFFER_FRAC

Format

An object of class numeric of length 1.

compute_permp

Description

This function will run permutation framework to compute a p-value for the correlation between the vectorised genes and clusters each cluster for one sample.

Usage

```
compute_permp(
    x,
    cluster_info,
    perm.size,
    bin_type,
    bin_param,
    test_genes,
    correlation_method = "pearson",
    n_cores = 1,
    correction_method = "BH",
    use_cm = FALSE
)
```

Arguments

x	a named list (of transcript detection coordinates) or named SingleCellExperi- ment or named SpatialExperiment or named SpatialFeatureExperiment object. If a named list is provided, the list element is a dataframe containing the tran- script detection coordinates and column names must include "feature_name" (gene name), "x" (x coordinate), "y" (y coordinate). The list name must match samples in cluster_info.
cluster_info	A dataframe/matrix containing the centroid coordinates and cluster label for each cell. The column names should include "x" (x coordinate), "y" (y coordinate), and "cluster" (cluster label).
perm.size	A positive number specifying permutation times
bin_type	A string indicating which bin shape is to be used for vectorization. One of "square" (default), "rectangle", or "hexagon".
bin_param	A numeric vector indicating the size of the bin. If the bin_type is "square" or "rectangle", this will be a vector of length two giving the numbers of rectangular quadrats in the x and y directions. If the bin_type is "hexagonal", this will be a number giving the side length of hexagons. Positive numbers only.
test_genes	A vector of strings giving the name of the genes you want to test correlation for. gene_mt.
correlation_method	
	A parameter pass to cor indicating which correlation coefficient is to be com- puted. One of "pearson" (default), "kendall", or "spearman": can be abbreviated.

compute_permp

n_cores	A positive number specifying number of cores used for parallelizing permuta- tion testing. Default is one core (sequential processing).
correction_met	hod
	A character string pass to p.adjust specifying the correction method for multiple testing .
use_cm	A boolean value that specifies whether to create spatial vectors for genes using the count matrix and cell coordinates instead of the transcript coordinates when both types of information are available. The default setting is FALSE.

Details

To get a permutation p-value for the correlation between a gene and a cluster, this function will permute the cluster label for each cell randomly, and calculate correlation between the genes and permuted clusters. This process will be repeated for perm.size times, and permutation p-value is calculated as the probability of permuted correlations larger than the observation correlation.

Value

An object of class 'cor_mg_result'. To access specific components of the returned object:

- Use get_cor to retrieve the matrix of observed correlation coefficients.
- Use get_perm_p to access the matrix of raw permutation p-values.
- Use get_perm_adjp to obtain the matrix of adjusted permutation p-values.

```
library(SpatialExperiment)
library(BumpyMatrix)
set.seed(100)
# simulate coordinates for clusters
df_clA <- data.frame(x = rnorm(n=10, mean=20, sd=5),</pre>
                    y = rnorm(n=10, mean=20, sd=5), cluster="A")
df_clB <- data.frame(x = rnorm(n=10, mean=100, sd=5),</pre>
                    y = rnorm(n=10, mean=100, sd=5), cluster="B")
clusters <- rbind(df_clA, df_clB)</pre>
clusters$sample="sample1"
# simulate coordinates for genes
trans_info <- data.frame(rbind(cbind(x = rnorm(n=10, mean=20, sd=5),
                                     y = rnorm(n=10, mean=20, sd=5),
                                     feature_name="gene_A1"),
                     cbind(x = rnorm(n=10, mean=20, sd=5),
                                     y = rnorm(n=10, mean=20, sd=5),
                                     feature_name="gene_A2"),
                     cbind(x = rnorm(n=10, mean=100, sd=5),
                                     y = rnorm(n=10, mean=100, sd=5),
                                     feature_name="gene_B1"),
                     cbind(x = rnorm(n=10, mean=100, sd=5),
                                     y = rnorm(n=10, mean=100, sd=5),
                                     feature_name="gene_B2")))
trans_info$x<-as.numeric(trans_info$x)</pre>
```

```
trans_info$y<-as.numeric(trans_info$y)</pre>
trans_info$cell = rep(paste("cell",1:20, sep=""), times=2)
mol <- BumpyMatrix::splitAsBumpyMatrix(</pre>
     trans_info[, c("x", "y")],
     row = trans_info$feature_name, col = trans_info$cell )
spe_sample1 <- SpatialExperiment(</pre>
        assays = list(molecules = mol),sample_id ="sample1" )
set.seed(100)
corr_res <- compute_permp(x=spe_sample1,</pre>
              cluster_info=clusters,
              perm.size=10,
              bin_type="square",
              bin_param=c(2,2),
              test_genes=unique(trans_info$feature_name),
              correlation_method = "pearson",
              n_cores=1,
              correction_method="BH")
# raw permutation p-value
perm_p <- get_perm_p(corr_res)</pre>
# adjusted permutation p-value
adjusted_perm_p <- get_perm_adjp(corr_res)</pre>
# observed correlation
obs_corr <- get_cor(corr_res)</pre>
```

create_genesets Convert the coordinates of set of genes into vectors.

Description

Convert the coordinates of set of genes into vectors.

```
create_genesets(
    x,
    name_lst,
    cluster_info,
    sample_names,
    bin_type,
    bin_param,
    use_cm = FALSE,
    n_cores = 1
)
```

X	a named list (of transcript detection coordinates) or SingleCellExperiment or SpatialExperiment or SpatialFeatureExperiment object. If a named list is pro- vided, every list element is a dataframe containing the transcript detection co- ordinates and column names must include "feature_name" (nagative control name), "x" (x coordinate) and "y" (y coordinate). The list names must match samples in cluster_info.
name_lst	A named list of strings giving the name of features that are treated as back- ground.
cluster_info	A dataframe/matrix containing the centroid coordinates, cluster and sample label for each cell.The column names must include "x" (x coordinate), "y" (y coordinate), "cluster" (cluster label) and "sample" (sample).
sample_names	a vector of strings giving the sample names
bin_type	A string indicating which bin shape is to be used for vectorization. One of "square" (default), "rectangle", or "hexagon".
bin_param	A numeric vector indicating the size of the bin. If the bin_type is "square" or "rectangle", this will be a vector of length two giving the numbers of rectangular quadrats in the x and y directions. If the bin_type is "hexagonal", this will be a number giving the side length of hexagons. Positive numbers only.
use_cm	A boolean value that specifies whether to create spatial vectors for genes using the count matrix and cell coordinates instead of the transcript coordinates when both types of information are available. The default setting is FALSE.
n_cores	A positive number specifying number of cores used for parallelizing permuta- tion testing. Default is one core (sequential processing).

Value

a list of two matrices with the following components

gene_mt	contains the transcript count in each grid. Each row refers to a grid, and each
	column refers to a gene.
aluatan mt	contains the number of calls in a specific cluster in each grid. Each new refere

cluster_mt contains the number of cells in a specific cluster in each grid. Each row refers to a grid, and each column refers to a cluster.

The row order of gene_mt matches the row order of cluster_mt.

A matrix contains the sum count in each grid. Each row refers to a grid, each column refers to a set in name_lst. The column name will match the names in name_lst.

```
feature_name="B"),
                          cbind(x = runif(10, min=10, max=24),
                                y = runif(10, min=10, max=24),
                                feature_name="C")))
trans$x = as.numeric(trans$x)
trans$y = as.numeric(trans$y)
trans$cell = sample(c("cell1","cell2","cell2"),replace=TRUE,
                         size=nrow(trans))
# create SpatialExperiment object
trans_mol <- BumpyMatrix::splitAsBumpyMatrix(</pre>
    trans[, c("x", "y")],
    row = trans$feature_name, col = trans$cell )
rep1_spe<- SpatialExperiment(</pre>
     assays = list(molecules = trans_mol),sample_id ="sample1" )
geneset_res <- create_genesets(x=rep1_spe, sample=c("sample1"),</pre>
                              name_lst=list(dummy_A=c("A","C"),
                                              dummy_B=c("A","B","C")),
                              bin_type="square",
                              bin_param=c(2,2),cluster_info=NULL)
```

get_cor

Get observed correlation cor_mg_result

Description

Accessor function to retrieve the observed correlation from an 'cor_mg_result' object.

Usage

get_cor(obj)

Arguments

obj

An 'cor_mg_result' object.

Value

A matrix contains the observation statistic for every gene and every cluster. Each row refers to a gene, and each column refers to a cluster

```
y = rnorm(n=10, mean=100, sd=5), cluster="B")
clusters <- rbind(df_clA, df_clB)</pre>
clusters$sample="sample1"
# simulate coordinates for genes
trans_info <- data.frame(rbind(cbind(x = rnorm(n=10, mean=20, sd=5),</pre>
                                     y = rnorm(n=10, mean=20, sd=5),
                                     feature_name="gene_A1"),
                     cbind(x = rnorm(n=10, mean=20, sd=5),
                                     y = rnorm(n=10, mean=20, sd=5),
                                     feature_name="gene_A2"),
                     cbind(x = rnorm(n=10, mean=100, sd=5),
                                     y = rnorm(n=10, mean=100, sd=5),
                                     feature_name="gene_B1"),
                     cbind(x = rnorm(n=10, mean=100, sd=5),
                                     y = rnorm(n=10, mean=100, sd=5),
                                      feature_name="gene_B2")))
trans_info$x<-as.numeric(trans_info$x)</pre>
trans_info$y<-as.numeric(trans_info$y)</pre>
trans_info$cell = rep(paste("cell",1:20, sep=""), times=2)
mol <- BumpyMatrix::splitAsBumpyMatrix(</pre>
     trans_info[, c("x", "y")],
     row = trans_info$feature_name, col = trans_info$cell )
spe_sample1 <- SpatialExperiment(</pre>
        assays = list(molecules = mol),sample_id ="sample1" )
set.seed(100)
corr_res <- compute_permp(x=spe_sample1,</pre>
             cluster_info=clusters,
             perm.size=10,
             bin_type="square",
             bin_param=c(2,2),
             test_genes=unique(trans_info$feature_name),
             correlation_method = "pearson",
             n_cores=1,
             correction_method="BH")
# observed correlation
obs_corr <- get_cor(corr_res)</pre>
```

get_full_mg Get full lasso result from glm_mg_result

Description

Accessor function to retrieve the 'full_result' dataframe from an 'glm_mg_result' object.

Usage

get_full_mg(obj, coef_cutoff = 0)

obj	An 'glm_mg_result' object.
coef_cutoff	A positive number giving the coefficient cutoff value. Genes whose cluster showing a coefficient value smaller than the cutoff will be removed. Default
	is 0.

Value

A data frame with detailed information for each gene and the most relevant cluster label.

- gene Gene name
- cluster The name of the significant cluster after
- glm_coef The coefficient of the selected cluster in the generalised linear model.
- pearson Pearson correlation between the gene vector and the selected cluster vector.
- max_gg_corr A number showing the maximum pearson correlation for this gene vector and all other gene vectors in the input gene_mt
- max_gc_corr A number showing the maximum pearson correlation for this gene vector and every cluster vectors in the input cluster_mt

```
library(SpatialExperiment)
set.seed(100)
# simulate coordinates for clusters
df_clA <- data.frame(x = rnorm(n=100, mean=20, sd=5),
                  y = rnorm(n=100, mean=20, sd=5), cluster="A")
df_clB <- data.frame(x = rnorm(n=100, mean=100, sd=5),</pre>
                y = rnorm(n=100, mean=100, sd=5), cluster="B")
clusters <- rbind(df_clA, df_clB)</pre>
clusters$sample<-"sample1"
# simulate coordinates for genes
trans_info <- data.frame(rbind(cbind(x = rnorm(n=100, mean=20, sd=5),</pre>
                                 y = rnorm(n=100, mean=20, sd=5),
                                  feature_name="gene_A1"),
                            cbind(x = rnorm(n=100, mean=20, sd=5))
                                  y = rnorm(n=100, mean=20, sd=5),
                                  feature_name="gene_A2"),
                            cbind(x = rnorm(n=100, mean=100, sd=5))
                                  y = rnorm(n=100, mean=100, sd=5),
                                  feature_name="gene_B1"),
                            cbind(x = rnorm(n=100, mean=100, sd=5),
                                  y = rnorm(n=100, mean=100, sd=5),
                                  feature_name="gene_B2")))
trans_info$x<-as.numeric(trans_info$x)</pre>
trans_info$y<-as.numeric(trans_info$y)</pre>
trans_info$cell<-sample(c("cell1","cell2","cell2"),replace=TRUE,</pre>
                         size=nrow(trans_info))
```

```
trans_mol <- BumpyMatrix::splitAsBumpyMatrix(</pre>
    trans_info[, c("x", "y")],
    row = trans_info$feature_name, col = trans_info$cell )
spe<- SpatialExperiment(</pre>
     assays = list(molecules = trans_mol), sample_id ="sample1" )
vecs_lst <- get_vectors(x=spe,sample_names=c("sample1"),</pre>
                     cluster_info = clusters,
                     bin_type = "square",
                     bin_param = c(20, 20),
                     test_genes =c("gene_A1","gene_A2","gene_B1","gene_B2"))
lasso_res <- lasso_markers(gene_mt=vecs_lst$gene_mt,</pre>
                         cluster_mt = vecs_lst$cluster_mt,
                         sample_names=c("sample1"),
                         keep_positive=TRUE,
                         background=NULL)
# the full result
full_result <- get_full_mg(lasso_res, coef_cutoff=0.05)</pre>
```

get_perm_adjp

Get permutation adjusted p value from cor_mg_result

Description

Accessor function to retrieve the permutation adjusted p-value from an 'cor_mg_result' object.

Usage

```
get_perm_adjp(obj)
```

Arguments

obj An 'cor_mg_result' object.

Value

A matrix contains the adjusted permutation p-value. Each row refers to a gene, and each column refers to a cluster.

```
# simulate coordinates for genes
trans_info <- data.frame(rbind(cbind(x = rnorm(n=10, mean=20, sd=5),</pre>
                                     y = rnorm(n=10, mean=20, sd=5),
                                      feature_name="gene_A1"),
                     cbind(x = rnorm(n=10, mean=20, sd=5),
                                     y = rnorm(n=10, mean=20, sd=5),
                                      feature_name="gene_A2"),
                     cbind(x = rnorm(n=10, mean=100, sd=5),
                                     y = rnorm(n=10, mean=100, sd=5),
                                      feature_name="gene_B1"),
                     cbind(x = rnorm(n=10, mean=100, sd=5),
                                     y = rnorm(n=10, mean=100, sd=5),
                                      feature_name="gene_B2")))
trans_info$x<-as.numeric(trans_info$x)</pre>
trans_info$y<-as.numeric(trans_info$y)</pre>
trans_info$cell = rep(paste("cell",1:20, sep=""), times=2)
mol <- BumpyMatrix::splitAsBumpyMatrix(</pre>
     trans_info[, c("x", "y")],
     row = trans_info$feature_name, col = trans_info$cell )
spe_sample1 <- SpatialExperiment(</pre>
        assays = list(molecules = mol),sample_id ="sample1" )
set.seed(100)
corr_res <- compute_permp(x=spe_sample1,</pre>
             cluster_info=clusters,
             perm.size=10,
             bin_type="square",
             bin_param=c(2,2),
             test_genes=unique(trans_info$feature_name),
             correlation_method = "pearson",
             n_cores=1,
             correction_method="BH")
# adjusted permutation p-value
adjusted_perm_p <- get_perm_adjp(corr_res)</pre>
```

get_perm_p

Get permutation p value from cor_mg_result

Description

Accessor function to retrieve the raw permutation p-value from an 'cor_mg_result' object.

Usage

get_perm_p(obj)

Arguments

obj An 'cor_mg_result' object.

get_perm_p

Value

A matrix contains the raw permutation p-value. Each row refers to a gene, and each column refers to a cluster.

```
library(SpatialExperiment)
library(BumpyMatrix)
set.seed(100)
# simulate coordinates for clusters
df_clA <- data.frame(x = rnorm(n=10, mean=20, sd=5),</pre>
                     y = rnorm(n=10, mean=20, sd=5), cluster="A")
df_clB <- data.frame(x = rnorm(n=10, mean=100, sd=5),</pre>
                     y = rnorm(n=10, mean=100, sd=5), cluster="B")
clusters <- rbind(df_clA, df_clB)</pre>
clusters$sample<-"sample1"
# simulate coordinates for genes
trans_info <- data.frame(rbind(cbind(x = rnorm(n=10, mean=20, sd=5),</pre>
                                      y = rnorm(n=10, mean=20, sd=5),
                                      feature_name="gene_A1"),
                     cbind(x = rnorm(n=10, mean=20, sd=5),
                                      y = rnorm(n=10, mean=20, sd=5),
                                      feature_name="gene_A2"),
                     cbind(x = rnorm(n=10, mean=100, sd=5),
                                      y = rnorm(n=10, mean=100, sd=5),
                                      feature_name="gene_B1"),
                     cbind(x = rnorm(n=10, mean=100, sd=5),
                                      y = rnorm(n=10, mean=100, sd=5),
                                      feature_name="gene_B2")))
trans_info$x<-as.numeric(trans_info$x)</pre>
trans_info$y<-as.numeric(trans_info$y)</pre>
trans_info$cell = rep(paste("cell",1:20, sep=""), times=2)
mol <- BumpyMatrix::splitAsBumpyMatrix(</pre>
     trans_info[, c("x", "y")],
     row = trans_info$feature_name, col = trans_info$cell )
spe_sample1 <- SpatialExperiment(</pre>
        assays = list(molecules = mol),sample_id ="sample1" )
set.seed(100)
corr_res <- compute_permp(x=spe_sample1,</pre>
             cluster_info=clusters,
             perm.size=10,
             bin_type="square",
             bin_param=c(2,2),
             test_genes=unique(trans_info$feature_name),
             correlation_method = "pearson",
             n_cores=1,
             correction_method="BH")
# raw permutation p-value
```

get_top_mg

Description

Accessor function to retrieve the 'top_result' dataframe from an 'glm_mg_result' object.

Usage

```
get_top_mg(obj, coef_cutoff = 0.05)
```

Arguments

obj	An 'glm_mg_result' object.
coef_cutoff	A positive number giving the coefficient cutoff value. Genes whose top cluster showing a coefficient value smaller than the cutoff will be marked as non-marker
	genes ("NoSig"). Default is 0.05.

Value

A data frame with detailed information for each gene and the most relevant cluster label.

- gene Gene name
- top_cluster The name of the most relevant cluster after thresholding the coefficients.
- glm_coef The coefficient of the selected cluster in the generalised linear model.
- pearson Pearson correlation between the gene vector and the selected cluster vector.
- max_gg_corr A number showing the maximum pearson correlation for this gene vector and all other gene vectors in the input gene_mt
- max_gc_corr A number showing the maximum pearson correlation for this gene vector and every cluster vectors in the input cluster_mt

```
feature_name="gene_A1"),
                            cbind(x = rnorm(n=100, mean=20, sd=5),
                                   y = rnorm(n=100, mean=20, sd=5),
                                   feature_name="gene_A2"),
                            cbind(x = rnorm(n=100, mean=100, sd=5),
                                   y = rnorm(n=100, mean=100, sd=5),
                                   feature_name="gene_B1"),
                            cbind(x = rnorm(n=100, mean=100, sd=5))
                                   y = rnorm(n=100, mean=100, sd=5),
                                   feature_name="gene_B2")))
trans_info$x<-as.numeric(trans_info$x)</pre>
trans_info$y<-as.numeric(trans_info$y)</pre>
trans_info$cell<-sample(c("cell1","cell2","cell2"),replace=TRUE,</pre>
                         size=nrow(trans_info))
trans_mol <- BumpyMatrix::splitAsBumpyMatrix(</pre>
    trans_info[, c("x", "y")],
    row = trans_info$feature_name, col = trans_info$cell )
spe<- SpatialExperiment(</pre>
     assays = list(molecules = trans_mol),sample_id ="sample1" )
vecs_lst <- get_vectors(x=spe,sample_names=c("sample1"),</pre>
                     cluster_info = clusters,
                     bin_type = "square",
                     bin_param = c(20, 20),
                     test_genes =c("gene_A1", "gene_A2", "gene_B1", "gene_B2"))
lasso_res <- lasso_markers(gene_mt=vecs_lst$gene_mt,</pre>
                         cluster_mt = vecs_lst$cluster_mt,
                         sample_names=c("sample1"),
                         keep_positive=TRUE,
                         background=NULL)
# the top result
top_result<- get_top_mg(lasso_res, coef_cutoff=0.05)</pre>
```

get_vectors

Vectorise the spatial coordinates

Description

This function will convert the coordinates into a numeric vector for genes and clusters.

```
get_vectors(
    x,
    cluster_info,
    sample_names,
    bin_type,
    bin_param,
    test_genes,
    use_cm = FALSE,
    n_cores = 1
)
```

X	a named list (of transcript detection coordinates) or SingleCellExperiment or SpatialExperiment or SpatialFeatureExperiment object. If a named list is pro- vided, every list element is a dataframe containing the transcript detection co- ordinates and column names must include "feature_name" (gene name), "x" (x coordinate), "y" (y coordinate). The list names must match samples in clus- ter_info.
cluster_info	A dataframe/matrix containing the centroid coordinates, cluster label and sample for each cell.The column names must include "x" (x coordinate), "y" (y coordinate), "cluster" (cluster label) and "sample" (sample). It is strongly recommended to use syntactically valid names for columns clusters and samples. If invalid names are detected, the function make.names will be employed to generate valid names. A message will also be displayed to indicate this change.
sample_names	a vector of strings giving the sample names. It is strongly recommended to use syntactically valid names for columns clusters and samples. If invalid names are detected, the function make.names will be employed to generate valid names. A message will also be displayed to indicate this change.
bin_type	A string indicating which bin shape is to be used for vectorization. One of "square" (default), "rectangle", or "hexagon".
bin_param	A numeric vector indicating the size of the bin. If the bin_type is "square" or "rectangle", this will be a vector of length two giving the numbers of rectangular quadrats in the x and y directions. If the bin_type is "hexagonal", this will be a number giving the side length of hexagons. Positive numbers only.
test_genes	A vector of strings giving the name of the genes you want to create gene vector. This will be used as column names for one of the result matrix gene_mt.
use_cm	A boolean value that specifies whether to create spatial vectors for genes using the count matrix and cell coordinates instead of the transcript coordinates when both types of information are available. The default setting is FALSE.
n_cores	A positive number specifying number of cores used for parallelizing permuta- tion testing. Default is one core (sequential processing).

Details

This function can be used to generate input for lasso_markers by specifying all the parameters.

Suppose the input data contains n genes, c clusters, and k samples, we want to use $a \times a$ square bin to convert the coordinates of genes and clusters into 1d vectors.

If k = 1, the returned list will contain one matrix for gene vectors (gene_mt) of dimension $a^2 \times n$ and one matrix for cluster vectors (cluster_mt) of dimension $a^2 \times c$.

If k > 1, gene and cluster vectors are constructed for each sample separately and concat together. There will be additional k columns on the returned cluster_mt, which is the one-hot encoding of the sample information.

Moreover, this function can vectorise genes and clusters separately based on the input. If x is NULL, this function will return vectorised clusters based on cluster_info. If cluster_info is NULL, this function will return vectorised genes based on x.

get_vectors

Value

a list of two matrices with the following components

gene_mt	contains the transcript count in each grid. Each row refers to a grid, and each column refers to a gene.
cluster_mt	contains the number of cells in a specific cluster in each grid. Each row refers to a grid, and each column refers to a cluster.

The row order of gene_mt matches the row order of cluster_mt.

```
library(SpatialExperiment)
set.seed(100)
# simulate coordinates for clusters
df_clA = data.frame(x = rnorm(n=100, mean=20, sd=5)),
                 y = rnorm(n=100, mean=20, sd=5), cluster="A")
df_clB = data.frame(x = rnorm(n=100, mean=100, sd=5),
                y = rnorm(n=100, mean=100, sd=5), cluster="B")
clusters = rbind(df_clA, df_clB)
clusters$sample="sample1"
# simulate coordinates for genes
trans_info = data.frame(rbind(cbind(x = rnorm(n=10, mean=20, sd=5),
                                y = rnorm(n=10, mean=20, sd=5),
                                 feature_name="gene_A1"),
                           cbind(x = rnorm(n=10, mean=20, sd=5),
                                 y = rnorm(n=10, mean=20, sd=5),
                                 feature_name="gene_A2"),
                           cbind(x = rnorm(n=10, mean=100, sd=5),
                                 y = rnorm(n=10, mean=100, sd=5),
                                  feature_name="gene_B1"),
                           cbind(x = rnorm(n=10, mean=100, sd=5),
                                 y = rnorm(n=10, mean=100, sd=5),
                                  feature_name="gene_B2")))
trans_info$x=as.numeric(trans_info$x)
trans_info$y=as.numeric(trans_info$y)
trans_info$cell = sample(c("cell1","cell2","cell2"),replace=TRUE,
                        size=nrow(trans_info))
# use named list as input
vecs_lst = get_vectors(x= list("sample1" = trans_info),
                    sample_names=c("sample1"),
                    cluster_info = clusters,
                    bin_type = "square",
                    bin_param = c(5,5),
                    test_genes =c("gene_A1", "gene_A2", "gene_B1", "gene_B2"))
# use SpatialExperiment object as input
trans_mol <- BumpyMatrix::splitAsBumpyMatrix(</pre>
    trans_info[, c("x", "y")],
    row = trans_info$feature_name, col = trans_info$cell )
spe<- SpatialExperiment(</pre>
```

lasso_markers

Find marker genes with spatial coordinates

Description

This function will find the most spatially relevant cluster label for each gene.

Usage

```
lasso_markers(
  gene_mt,
  cluster_mt,
  sample_names,
  keep_positive = TRUE,
  background = NULL,
  n_fold = 10
)
```

Arguments

gene_mt	A matrix contains the transcript count in each grid. Each row refers to a grid, and each column refers to a gene. The column names must be specified and refer to the genes. This can be the output from the function get_vectors.
cluster_mt	A matrix contains the number of cells in a specific cluster in each grid. Each row refers to a grid, and each column refers to a cluster. The column names must be specified and refer to the clusters. Please do not assign integers as column names. This can be the output from the function get_vectors.
sample_names	A vector specifying the names for the samples.
keep_positive	A logical flag indicating whether to return positively correlated clusters or not.
background	Optional. A matrix providing the background information. Each row refers to a grid, and each column refers to one category of background information. Number of rows must equal to the number of rows in gene_mt and cluster_mt. Can be obtained by only providing coordinates matrices cluster_info. to function get_vectors.
n_fold	Optional. A positive number giving the number of folds used for cross valida- tion. This parameter will pass to cv.glmnet to calculate a penalty term for every gene.

lasso_markers

Details

This function will take the converted gene and cluster vectors from function get_vectors, and return the most relevant cluster label for each gene. If there are multiple samples in the dataset, this function will find shared markers across different samples by including additional sample vectors in the input cluster_mt.

This function treats all input cluster vectors as features, and create a penalized linear model for one gene vector with lasso regularization. Clusters with non-zero coefficient will be selected, and these clusters will be used to formulate a generalised linear model for this gene vector.

- If the input keep_positive is TRUE, the clusters with positive coefficient and significant p-value will be saved in the output matrix lasso_full_result. The cluster with a positive coefficient and the minimum p-value will be regarded as the most relevant cluster to this gene and be saved in the output matrix lasso_result.
- If the input keep_positive is FALSE, the clusters with negative coefficient and significant p-value will be saved in the output matrix lasso_full_result. The cluster with a negative coefficient and the minimum p-value will be regarded as the most relevant cluster to this gene and be saved in the output matrix lasso_result.

If there is no clusters with significant p-value, the a string "NoSig" will be returned for this gene.

The parameter background can be used to capture unwanted noise pattern in the dataset. For example, we can include negative control genes as a background cluster in the model. If the most relevant cluster selected by one gene matches the background "clusters", we will return "NoSig" for this gene.

Value

An object of class 'glm_mg_result' To access specific components of the returned object:

- Use get_top_mg to retrieve the top result data frame
- Use get_full_mg to retrieve full result data frame

See Also

get_vectors

Examples

simulate coordinates for genes

```
trans_info <- data.frame(rbind(cbind(x = rnorm(n=100, mean=20, sd=5),</pre>
                                  y = rnorm(n=100, mean=20, sd=5),
                                   feature_name="gene_A1"),
                            cbind(x = rnorm(n=100, mean=20, sd=5),
                                   y = rnorm(n=100, mean=20, sd=5),
                                   feature_name="gene_A2"),
                            cbind(x = rnorm(n=100, mean=100, sd=5),
                                   y = rnorm(n=100, mean=100, sd=5),
                                   feature_name="gene_B1"),
                            cbind(x = rnorm(n=100, mean=100, sd=5),
                                   y = rnorm(n=100, mean=100, sd=5),
                                   feature_name="gene_B2")))
trans_info$x<-as.numeric(trans_info$x)</pre>
trans_info$y<-as.numeric(trans_info$y)</pre>
trans_info$cell<-sample(c("cell1","cell2","cell2"),replace=TRUE,</pre>
                         size=nrow(trans_info))
trans_mol <- BumpyMatrix::splitAsBumpyMatrix(</pre>
    trans_info[, c("x", "y")],
    row = trans_info$feature_name, col = trans_info$cell )
spe<- SpatialExperiment(</pre>
     assays = list(molecules = trans_mol),sample_id ="sample1" )
vecs_lst <- get_vectors(x=spe,sample_names=c("sample1"),</pre>
                     cluster_info = clusters,
                     bin_type = "square",
                     bin_param = c(20, 20),
                     test_genes =c("gene_A1", "gene_A2", "gene_B1", "gene_B2"))
lasso_res <- lasso_markers(gene_mt=vecs_lst$gene_mt,</pre>
                         cluster_mt = vecs_lst$cluster_mt,
                         sample_names=c("sample1"),
                         keep_positive=TRUE,
                         background=NULL)
# the top result
top_result <- get_top_mg(lasso_res, coef_cutoff=0.05)</pre>
# the full result
full_result <- get_full_mg(lasso_res, coef_cutoff=0.05)</pre>
```

rep1_clusters Rep1 selected cells

Description

A data frame file containing the coordinates and cluster label for each cell of the selected subset of rep1.

Usage

data(rep1_clusters)

rep1_neg

Format

A data frame with 1705 rows and 6 variables:

anno the provided cell type annotation

cluster cluster label

x x coordinates

 \mathbf{y} y coordiantes

cells cell id

sample sample id

Value

data frame

Source

<https://cf.10xgenomics.com/samples/xenium/1.0.1/Xenium_FFPE_Human_Breast_Cancer_Rep1/ Xenium_FFPE_Human_Breast_Cancer_Rep1_outs.zip>

rep1_neg

Rep1 negative control genes within the selected region.

Description

A SpatialExperiment object containing the coordinates for every negative control detection for rep1_sub

Usage

data(rep1_neg)

Format

A SpatialExperiment object. The molecules assay slot is a BumpyDataFrameMatrix obejct. Can retrieve DataFrame version by calling 'BumpyMatrix::unsplitAsDataFrame(molecules(rep1_neg))'. The molecules slot contains:

x x coordinates

y y coordiantes

feature_name negative control probe name

category negative control category

Value

SpatialExperiment

Source

https://cf.10xgenomics.com/samples/xenium/1.0.1/Xenium_FFPE_Human_Breast_Cancer_Rep1/Xenium_FFPE_Human_Cancer_Rep1_outs.zip

rep1_sub

A small section of Xenium human breast cancer rep1.

Description

A SpatialExperiment object containing the coordinates for every transcript

Usage

data(rep1_sub)

Format

A SpatialExperiment object with 20 genes and 1713 cells. The molecules assay slot is a Bumpy-DataFrameMatrix obejct. Can retrieve DataFrame version by calling 'BumpyMatrix::unsplitAsDataFrame(molecules(rep2_sr The molecules assay contains 79576 rows and 3 variables:

- x x coordinates
- y y coordiantes

feature_name transcript name

Value

SpatialExperiment

Source

<https://cf.10xgenomics.com/samples/xenium/1.0.1/Xenium_FFPE_Human_Breast_Cancer_Rep1/ Xenium_FFPE_Human_Breast_Cancer_Rep1_outs.zip> rep2_clusters Rep2 selected cells

Description

A csv file containing the coordinates and cluster label for each cell of the selected subset of rep2.

Usage

data(rep2_clusters)

Format

A data frame with 1815 rows and and 6 variables:

anno the provided cell type annotation

cluster cluster label

x x coordinates

y y coordiantes

cells cell id

sample sample id

Value

data frame

Source

<https://cf.10xgenomics.com/samples/xenium/1.0.1/Xenium_FFPE_Human_Breast_Cancer_Rep2/Xenium_FFPE_Human_Breast_Cancer_Rep2_outs.zip>

rep2_neg

Rep2 negative control genes within the selected region.

Description

A data frame containing the coordinates for every negative control detection for rep2

Usage

data(rep2_neg)

Format

A SpatialExperiment object. The molecules assay slot is a BumpyDataFrameMatrix obejct. Can retrieve DataFrame version by calling 'BumpyMatrix::unsplitAsDataFrame(molecules(rep2_neg))'. The molecules slot contains:

x x coordinates

y y coordiantes

feature_name negative control probe name .

category negative control category

Value

SpatialExperiment

Source

<https://cf.10xgenomics.com/samples/xenium/1.0.1/Xenium_FFPE_Human_Breast_Cancer_Rep2/Xenium_FFPE_Human_Breast_Cancer_Rep2_outs.zip>

rep2_sub

A small section of Xenium human breast cancer rep2.

Description

A SpatialExperiment object containing the coordinates for every negative control detection for rep2_sub

Usage

data(rep2_sub)

Format

A SpatialExperiment object with 20 genes and 1829 cells. The molecules assay slot is a Bumpy-DataFrameMatrix obejct. Can retrieve DataFrame version by calling 'BumpyMatrix::unsplitAsDataFrame(molecules(rep2_st)). The molecules slot contains:

x x coordinatesy y coordiantesfeature_name transcript name

Value

SpatialExperiment

Source

<https://cf.10xgenomics.com/samples/xenium/1.0.1/Xenium_FFPE_Human_Breast_Cancer_Rep2/ Xenium_FFPE_Human_Breast_Cancer_Rep2_outs.zip>

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