# Package 'Giotto'

July 7, 2020

```
Title Spatial single-cell transcriptomics toolbox
Version 0.3.6.9000
Description Toolbox to process, analyze and visualize spatial single-cell expression data
License MIT + file LICENSE
Encoding UTF-8
LazyData true
URL https://rubd.github.io/Giotto/, https://github.com/RubD/Giotto
BugReports https://github.com/RubD/Giotto/issues
RoxygenNote 7.1.1
Depends base (>= 3.5.1),
      utils (>= 3.5.1),
      R (>= 3.5.1)
Imports data.table (>= 1.12.2),
      deldir,
      ggplot2 (>= 3.1.1),
      Matrix,
      magick,
      matrixStats (\geq 0.55.0),
      methods,
      uwot (>= 0.0.0.9010),
      cowplot (>= 0.9.4),
      grDevices,
      graphics,
      RColorBrewer (>= 1.1-2),
      dbscan (>= 1.1-3),
      farver (>= 2.0.3),
      ggalluvial (>= 0.9.1),
      scales (>= 1.0.0),
      ComplexHeatmap (>= 1.20.0),
      qvalue (>= 2.14.1),
      If a (>= 1.12.0),
      igraph (>= 1.2.4.1),
      irlba,
      plotly,
      parallel,
      reticulate (>= 1.14),
```

2 R topics documented:

magrittr,

```
limma,
  ggdendro,
  smfishHmrf,
  devtools,
  reshape2,
  ggraph,
  Rcpp,
  Rfast,
  Rtsne (>= 0.15),
  rlang (>= 0.4.3),
  R.utils,
  fitdistrplus,
Suggests knitr,
  rmarkdown,
  MAST,
  scran (>= 1.10.1),
  png,
  FactoMineR,
  tiff.
  biomaRt.
  trendsceek,
  multinet (>= 3.0.2),
  RTriangle (>= 1.6-0.10)
biocViews
VignetteBuilder knitr
LinkingTo Rcpp,
  RcppArmadillo
Remotes lambdamoses/smfishhmrf-r
R topics documented:
```

addHMRF.......... 

20

average_gene_gene_expression_in_groups
binSpect
calculateHVG
calculateMetaTable
calculateMetaTableCells
calculate_distance_and_weight
cellProximityBarplot
cellProximityEnrichment
cellProximityHeatmap
cellProximityNetwork
cellProximitySpatPlot
cellProximitySpatPlot2D
cellProximitySpatPlot3D
cellProximityVisPlot
cellProximityVisPlot_2D_ggplot
cellProximityVisPlot_2D_plotly
cellProximityVisPlot_3D_plotly
changeGiottoInstructions
changeImageBg
clusterCells
clusterSpatialCorGenes
combCCcom
combineCellProximityGenes
combineCellProximityGenes_per_interaction
combineCPG
combineMetadata
convertEnsemblToGeneSymbol
convert_mgImage_to_array_DT
createCrossSection
createGiottoImage
createGiottoInstructions
createGiottoObject
createGiottoVisiumObject
createHeatmap_DT
createMetagenes
createNearestNetwork
createSpatialDelaunayNetwork
createSpatialEnrich
createSpatialGrid
createSpatialGrid_2D
createSpatialGrid_3D
createSpatialKNNnetwork
createSpatialNetwork
create_average_detection_DT
create_average_DT
create_cell_type_random_cell_IDs
create_crossSection_object
create_delaunayNetwork2D
create_delaunayNetwork3D
create_delaunayNetwork_deldir
create_delaunayNetwork_geometry
create, delaunayNetwork, geometry, 3D

create_delaunayNetwork_RTriangle	79
create_genes_to_use_matrix	79
create_KNNnetwork_dbscan	
create_screeplot	
crossSectionGenePlot	
crossSectionGenePlot3D	
crossSectionPlot	
crossSectionPlot3D	84
decide_cluster_order	
detectSpatialCorGenes	86
detectSpatialPatterns	
dimCellPlot	88
dimCellPlot2D	90
dimGenePlot	93
dimGenePlot2D	94
dimGenePlot3D	96
dimPlot	98
dimPlot2D	100
dimPlot3D	
doHclust	
doHMRF	
doKmeans	
doLeidenCluster	
doLeidenSubCluster	
doLouvainCluster	
doLouvainCluster_community	
doLouvainCluster_multinet	
doLouvainSubCluster	
doLouvainSubCluster_community	
doLouvainSubCluster_multinet	
doRandomWalkCluster	
doSNNCluster	
do_permuttest_random	
DT removeNA	
estimateImageBg	
exportGiottoViewer	
exprCellCellcom	
extractNearestNetwork	
fDataDT	
filterCellProximityGenes	
filterCombinations	
filterCPG	
filterDistributions	
filterGiotto	
findCellProximityGenes	
findCellProximityGenes_per_interaction	
findCPG	
findGiniMarkers	
findGiniMarkers_one_vs_all	
findMarkers_one_vs_all	
findMastMarkers	143

findMastMarkers_one_vs_all	
findNetworkNeighbors	
findScranMarkers	
findScranMarkers_one_vs_all	
find_grid_2D	
find_grid_3D	
$find\_grid\_x$	
find_grid_y	
find_grid_z	
FSV_show	149
get10Xmatrix	150
getClusterSimilarity	150
getDendrogramSplits	151
getDistinctColors	152
getGiottoImage	153
get_os	153
giotto-class	154
heatmSpatialCorGenes	154
hyperGeometricEnrich	156
insertCrossSectionGenePlot3D	157
insertCrossSectionSpatPlot3D	158
jackstrawPlot	
kmeans_binarize	
loadHMRF	161
makeSignMatrixPAGE	162
makeSignMatrixRank	
make_simulated_network	
mergeClusters	164
my_arowMeans	
my_growMeans	
my_rowMeans	165
nnDT to kNN	166
node_clusters	166
normalizeGiotto	167
PAGEEnrich	
pca_giotto	169
pDataDT	170
plotCCcomDotplot	170
plotCCcomHeatmap	171
plotCellProximityGenes	
plotCombineCCcom	
plotCombineCellCellCommunication	
plotCombineCellProximityGenes	
plotCombineCPG	
plotCPG	
plotGiottoImage	
plotHeatmap	
plotICG	
plotInteractionChangedGenes	
plotly_axis_scale_2D	
plotly_axis_scale_3D	
plotly_grid	

plotly_network	
plotMetaDataCellsHeatmap	
plotMetaDataHeatmap	
plotPCA	
plotPCA_2D	
plotPCA_3D	
plotRankSpatvsExpr	
plotRecovery	
plotRecovery_sub	
plotStatDelaunayNetwork	
plotTSNE	
plotTSNE_2D	
plotTSNE_3D	
plotUMAP	. 205
plotUMAP_2D	. 207
plotUMAP_3D	. 209
print.giotto	. 210
rankEnrich	. 210
rankSpatialCorGroups	. 211
rank_binarize	. 212
readExprMatrix	. 212
readGiottoInstructions	. 213
removeCellAnnotation	. 214
removeGeneAnnotation	. 214
replaceGiottoInstructions	
runPCA	
runPCA_factominer	
runPCA_prcomp_irlba	
runtSNE	
runUMAP	
screePlot	
selectPatternGenes	
select_expression_values	
select_spatialNetwork	
set_giotto_python_path	
show,giotto-method	
showClusterDendrogram	
showClusterHeatmap	
showGiottoImageNames	
show Giotto Instructions	
showGrids	
showNetworks	
showPattern	
showPattern2D	
showPattern3D	
showPatternGenes	
showProcessingSteps	
showSaveParameters	
showSpatialCorGenes	
signPCA	
silhouetteRank	
spatCellCellcom	. 439

spatCellPlot
spatCellPlot2D
spatDimCellPlot
spatDimCellPlot2D
spatDimGenePlot
spatDimGenePlot2D
spatDimGenePlot3D
spatDimPlot
spatDimPlot2D
spatDimPlot3D
spatGenePlot
spatGenePlot2D
spatGenePlot3D
spatialAEH
spatialDE
spatNetwDistributions
spatNetwDistributionsDistance
spatNetwDistributionsKneighbors
spatPlot
spatPlot2D
spatPlot3D
spat_fish_func
spat_OR_func
specificCellCellcommunicationScores
split_dendrogram_in_two
standardise_giotto
stitchFieldCoordinates
stitchTileCoordinates
subClusterCells
subsetGiotto
subsetGiottoLocs
trendSceek
updateGiottoImage
viewHMRFresults
viewHMRFresults2D
viewHMRFresults3D
violinPlot
visDimGenePlot
visDimGenePlot_2D_ggplot
visDimGenePlot_3D_plotly
visDimPlot
visDimPlot_2D_ggplot
visDimPlot_2D_plotly
visDimPlot_3D_plotly
visForceLayoutPlot
visGenePlot
visGenePlot_2D_ggplot
visGenePlot_3D_plotly
visPlot
visPlot_2D_ggplot
visPlot_2D_plotly
visPlot_3D_plotly

8 addCellIntMetadata

34	
34	
34	
34	341
33	338
33	
33	
33	
	3

\_ . .

aaaCeнIntMetaaata

#### **Description**

Creates an additional metadata column with information about interacting and non-interacting cell types of the selected cell-cell interaction.

# Usage

```
addCellIntMetadata(
  gobject,
  spatial_network = "spatial_network",
  cluster_column,
  cell_interaction,
  name = "select_int",
  return_gobject = TRUE
)
```

# **Arguments**

```
gobject giotto object

spatial_network

name of spatial network to use

cluster_column column of cell types

cell_interaction

cell-cell interaction to use

name

name for the new metadata column

return_gobject return an updated giotto object
```

#### **Details**

This function will create an additional metadata column which selects interacting cell types for a specific cell-cell interaction. For example, if you want to color interacting astrocytes and oligodendrocytes it will create a new metadata column with the values "select\_astrocytes", "select\_oligodendrocytes", "other\_astrocytes", "other\_oligodendrocytes" and "other". Where "other" is all other cell types found within the selected cell type column.

addCellMetadata 9

#### Value

Giotto object

# **Examples**

```
addCellIntMetadata(gobject)
```

addCellMetadata

addCellMetadata

#### **Description**

adds cell metadata to the giotto object

#### Usage

```
addCellMetadata(
  gobject,
  new_metadata,
  vector_name = NULL,
  by_column = FALSE,
  column_cell_ID = NULL
)
```

# Arguments

```
gobject giotto object

new_metadata new cell metadata to use (data.table, data.frame, ...)

vector_name (optional) custom name if you provide a single vector

by_column merge metadata based on cell_ID column in pDataDT (default = FALSE)

column_cell_ID column name of new metadata to use if by_column = TRUE
```

#### **Details**

You can add additional cell metadata in two manners:

- 1. Provide a data.table or data.frame with cell annotations in the same order as the cell\_ID column in pDataDT(gobject)
- 2. Provide a data.table or data.frame with cell annotations and specificy which column contains the cell IDs, these cell IDs need to match with the cell\_ID column in pDataDT(gobject)

# Value

giotto object

# **Examples**

```
addCellMetadata(gobject)
```

10 addCellStatistics

 ${\tt addCellStatistics}$ 

addCellStatistics

### **Description**

adds cells statistics to the giotto object

### Usage

```
addCellStatistics(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  detection_threshold = 0,
  return_gobject = TRUE
)
```

# **Arguments**

### **Details**

This function will add the following statistics to cell metadata:

- nr\_genes: Denotes in how many genes are detected per cell
- perc\_genes: Denotes what percentage of genes is detected per cell
- total\_expr: Shows the total sum of gene expression per cell

# Value

```
giotto object if return_gobject = TRUE
```

### **Examples**

```
addCellStatistics(gobject)
```

addGeneMetadata 11

addGeneMetadata

addGeneMetadata

### **Description**

adds gene metadata to the giotto object

#### Usage

```
addGeneMetadata(gobject, new_metadata, by_column = F, column_gene_ID = NULL)
```

#### **Arguments**

```
gobject giotto object

new_metadata new metadata to use

by_column merge metadata based on gene_ID column in fDataDT

column_gene_ID column name of new metadata to use if by_column = TRUE
```

#### **Details**

You can add additional gene metadata in two manners: 1. Provide a data.table or data.frame with gene annotations in the same order as the gene\_ID column in fDataDT(gobject) 2. Provide a data.table or data.frame with gene annotations and specificy which column contains the gene IDs, these gene IDs need to match with the gene\_ID column in fDataDT(gobject)

#### Value

giotto object

### **Examples**

```
addGeneMetadata(gobject)
```

addGenesPerc

addGenesPerc

#### **Description**

calculates the total percentage of (normalized) counts for a subset of selected genes

```
addGenesPerc(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  genes = NULL,
  vector_name = "gene_perc",
  return_gobject = TRUE
)
```

12 addGeneStatistics

#### **Arguments**

#### Value

```
giotto object if return_gobject = TRUE, else a vector with
```

# **Examples**

```
addGenesPerc(gobject)
```

addGeneStatistics

addGeneStatistics

### **Description**

adds gene statistics to the giotto object

# Usage

```
addGeneStatistics(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  detection_threshold = 0,
  return_gobject = TRUE
)
```

# **Arguments**

```
gobject giotto object
expression_values
expression values to use
detection_threshold
detection threshold to consider a gene detected
return_gobject boolean: return giotto object (default = TRUE)
```

# **Details**

This function will add the following statistics to gene metadata:

- nr\_cells: Denotes in how many cells the gene is detected
- per\_cells: Denotes in what percentage of cells the gene is detected
- total\_expr: Shows the total sum of gene expression in all cells
- mean\_expr: Average gene expression in all cells
- mean\_expr\_det: Average gene expression in cells with detectable levels of the gene

addGiottoImage 13

#### Value

```
giotto object if return_gobject = TRUE
```

# **Examples**

```
addGeneStatistics(gobject)
```

 ${\tt addGiottoImage}$ 

add Giot to Image

### **Description**

Adds giotto image objects to your giotto object

# Usage

```
addGiottoImage(gobject, images)
```

# **Arguments**

gobject giotto object

images list of giotto image objects, see createGiottoImage

# Value

an updated Giotto object with access to the list of images

### **Examples**

```
addGiottoImage(mg_object)
```

```
add {\tt GiottoImageToSpatPlot}
```

addGiottoImageToSpatPlot

# Description

Add a giotto image to a spatial ggplot object post creation

# Usage

```
addGiottoImageToSpatPlot(spatpl = NULL, gimage = NULL)
```

# **Arguments**

```
spatpl a spatial ggplot object
```

gimage a giotto image, see createGiottoImage

14 addHMRF

#### Value

an updated spatial ggplot object

# **Examples**

addGiottoImageToSpatPlot(mg\_object)

addHMRF

addHMRF

# Description

Add selected results from doHMRF to the giotto object

# Usage

```
addHMRF(gobject, HMRFoutput, k = NULL, betas_to_add = NULL, hmrf_name = NULL)
```

# Arguments

gobject giotto object

HMRF output from doHMRF()

k number of domains

hmrf\_name specify a custom name

# **Details**

Description ...

#### Value

giotto object

# **Examples**

addHMRF(gobject)

addNetworkLayout 15

addNetworkLayout

addNetworkLayout

# Description

Add a network layout for a selected nearest neighbor network

### Usage

```
addNetworkLayout(
  gobject,
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  layout_type = c("drl"),
  options_list = NULL,
  layout_name = "layout",
  return_gobject = TRUE
)
```

# Arguments

# **Details**

This function creates layout coordinates based on the provided kNN or sNN. Currently only the force-directed graph layout "drl", see layout\_with\_drl, is implemented. This provides an alternative to tSNE or UMAP based visualizations.

# Value

giotto object with updated layout for selected NN network

# Examples

```
addNetworkLayout(gobject)
```

16 adjustGiottoMatrix

addStatistics

addStatistics

# Description

adds genes and cells statistics to the giotto object

# Usage

```
addStatistics(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  detection_threshold = 0,
  return_gobject = TRUE
)
```

# **Arguments**

### **Details**

See addGeneStatistics and addCellStatistics

# Value

```
giotto object if return_gobject = TRUE, else a list with results
```

# **Examples**

```
addStatistics(gobject)
```

adjustGiottoMatrix adjustGiottoMatrix

# Description

normalize and/or scale expresion values of Giotto object

all\_plots\_save\_function

#### Usage

```
adjustGiottoMatrix(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  batch_columns = NULL,
  covariate_columns = NULL,
  return_gobject = TRUE,
  update_slot = c("custom")
)
```

### **Arguments**

```
gobject giotto object

expression_values

expression values to use

batch_columns metadata columns that represent different batch (max = 2)

covariate_columns

metadata columns that represent covariates to regress out

return_gobject boolean: return giotto object (default = TRUE)

update_slot expression slot that will be updated (default = custom)
```

#### **Details**

This function implements the removeBatchEffect function to remove known batch effects and to adjust expression values according to provided covariates.

#### Value

giotto object

### **Examples**

```
adjustGiottoMatrix(gobject)
```

```
all_plots_save_function 
 all_plots_save_function
```

#### **Description**

Function to automatically save plots to directory of interest

```
all_plots_save_function(
  gobject,
  plot_object,
  save_dir = NULL,
  save_folder = NULL,
  save_name = NULL,
```

```
default_save_name = "giotto_plot",
    save_format = NULL,
    show_saved_plot = F,
    ncol = 1,
    nrow = 1,
    scale = 1,
    base_width = NULL,
    base_height = NULL,
    base_aspect_ratio = NULL,
    units = NULL,
    dpi = NULL,
    limitsize = TRUE,
    ...
)
```

### **Arguments**

```
gobject
                  giotto object
plot_object
                  object to plot
save_dir
                  directory to save to
save_folder
                  folder in save_dir to save to
                  name of plot
save_name
default_save_name
                  default name to save a plot
save_format
                  format (e.g. png, tiff, pdf, ...)
show_saved_plot
                  load & display the saved plot
                  number of columns
ncol
                  number of rows
nrow
scale
                  scale
base_width
                  width
base_height
                  height
base_aspect_ratio
                  aspect ratio
units
                  units
dpi
                  Plot resolution
                  When TRUE (the default), ggsave will not save images larger than 50x50 inches,
limitsize
                  to prevent the common error of specifying dimensions in pixels.
```

#### See Also

```
{\tt general\_save\_function}
```

# **Examples**

```
all_plots_save_function(gobject)
```

additional parameters to ggplot\_save\_function or general\_save\_function

annotateGiotto 19

annotateGiotto

annotate Giotto

### **Description**

Converts cluster results into provided annotation.

# Usage

```
annotateGiotto(
  gobject,
  annotation_vector = NULL,
  cluster_column = NULL,
  name = "cell_types"
)
```

# Arguments

# Details

You need to specifify which (cluster) column you want to annotate and you need to provide an annotation vector like this:

- 1. identify the cell type of each cluster
- 2. create a vector of these cell types, e.g. cell\_types = c('T-cell', 'B-cell', 'Stromal')
- 3. provide original cluster names to previous vector, e.g. names(cell\_types) = c(2, 1, 3)

# Value

giotto object

### **Examples**

```
annotateGiotto(gobject)
```

```
annotate Spatial Grid \qquad annotate Spatial Grid
```

### **Description**

annotate spatial grid with cell ID and cell metadata (optional)

### Usage

```
annotateSpatialGrid(
  gobject,
  spatial_grid_name = "spatial_grid",
  cluster_columns = NULL
)
```

# Arguments

### Value

annotated spatial grid data.table

### **Examples**

```
annotateSpatialGrid()
```

```
annotateSpatialNetwork
```

annotate Spatial Network

# Description

Annotate spatial network with cell metadata information.

```
annotateSpatialNetwork(
  gobject,
  spatial_network_name = "Delaunay_network",
  cluster_column,
  create_full_network = F
)
```

### **Arguments**

### Value

annotated network in data.table format

### **Examples**

```
annotateSpatialNetwork(gobject)
```

```
annotate\_spatlocs\_with\_spatgrid\_2D \\ annotate\_spatlocs\_with\_spatgrid\_2D
```

### **Description**

annotate spatial locations with 2D spatial grid information

# Usage

```
annotate_spatlocs_with_spatgrid_2D(spatloc, spatgrid)
```

# Arguments

spatloc spatial\_locs slot from giotto object

spatgrid selected spatial\_grid slot from giotto object

# Value

annotated spatial location data.table

# **Examples**

```
annotate_spatlocs_with_spatgrid_2D()
```

# Description

annotate spatial locations with 3D spatial grid information

### Usage

```
annotate_spatlocs_with_spatgrid_3D(spatloc, spatgrid)
```

# Arguments

spatloc spatial\_locs slot from giotto object

spatgrid selected spatial\_grid slot from giotto object

#### Value

annotated spatial location data.table

# **Examples**

```
annotate_spatlocs_with_spatgrid_3D()
```

```
average_gene_gene_expression_in_groups

average_gene_gene_expression_in_groups
```

### **Description**

calculate average expression per cluster

#### Usage

```
average_gene_gene_expression_in_groups(
  gobject,
  cluster_column = "cell_types",
  gene_set_1,
  gene_set_2
)
```

# **Arguments**

```
gobject giotto object to use
cluster_column cluster column with cell type information
gene_set_1 first specific gene set from gene pairs
gene_set_2 second specific gene set from gene pairs
```

binSpect 23

#### **Details**

Details will follow soon.

#### Value

data.table with average expression scores for each cluster

#### **Examples**

```
average_gene_gene_expression_in_groups(gobject)
```

binSpect

binSpect

# Description

Previously: binGetSpatialGenes. BinSpect (Binary Spatial Extraction of genes) is a fast computational method that identifies genes with a spatially coherent expression pattern.

#### Usage

```
binSpect(
  gobject,
  bin_method = c("kmeans", "rank"),
  expression_values = c("normalized", "scaled", "custom"),
  subset_genes = NULL,
  spatial_network_name = "Delaunay_network",
  nstart = 3,
  iter_max = 10,
  percentage_rank = 30,
  do_fisher_test = TRUE,
  calc_hub = FALSE,
  hub_min_int = 3,
  get_av_expr = TRUE,
  get_high_expr = TRUE,
  do_parallel = TRUE,
  cores = NA,
  verbose = T
)
```

# Arguments

24 binSpect

iter\_max kmeans: iter.max parameter

percentage\_rank

percentage of top cells for binarization

do\_fisher\_test perform fisher test

calc\_hub calculate the number of hub cells

hub\_min\_int minimum number of cell-cell interactions for a hub cell

get\_av\_expr calculate the average expression per gene of the high expressing cells

get\_high\_expr calculate the number of high expressing cells per gene

do\_parallel run calculations in parallel with mclapply

cores number of cores to use if do\_parallel = TRUE

verbose be verbose

#### **Details**

We provide two ways to identify spatial genes based on gene expression binarization. Both methods are identicial except for how binarization is performed.

- 1. binarize: Each gene is binarized (0 or 1) in each cell with **kmeans** (k = 2) or based on **rank** percentile
- 2. network: Alll cells are connected through a spatial network based on the physical coordinates
- 3. contingency table: A contingency table is calculated based on all edges of neighboring cells and the binarized expression (0-0, 0-1, 1-0 or 1-1)
- 4. For each gene an odds-ratio (OR) and fisher.test (optional) is calculated

Other statistics are provided (optional):

- Number of cells with high expression (binary = 1)
- Average expression of each gene within high expressing cells
- Number of hub cells, these are high expressing cells that have a user defined number of high expressing neighbors

By selecting a subset of likely spatial genes (e.g. soft thresholding highly variable genes) or using multiple cores can accelerate the speed.

#### Value

data.table with results (see details)

# **Examples**

binSpect(gobject)

calculateHVG 25

calculateHVG

calculateHVG

### **Description**

compute highly variable genes

# Usage

```
calculateHVG(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  method = c("cov_groups", "cov_loess"),
  reverse_log_scale = FALSE,
  logbase = 2,
  expression_threshold = 0,
  nr_expression_groups = 20,
  zscore_threshold = 1.5,
  HVGname = "hvg",
  difference_in_cov = 0.1,
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "HVGplot",
  return_gobject = TRUE
)
```

# **Arguments**

```
gobject
                  giotto object
expression_values
                  expression values to use
method
                  method to calculate highly variable genes
reverse_log_scale
                  reverse log-scale of expression values (default = FALSE)
                  if reverse_log_scale is TRUE, which log base was used?
logbase
expression_threshold
                  expression threshold to consider a gene detected
nr_expression_groups
                  number of expression groups for cov_groups
{\tt zscore\_threshold}
                  zscore to select hvg for cov_groups
HVGname
                  name for highly variable genes in cell metadata
difference_in_cov
                  minimum difference in coefficient of variance required
show_plot
                  show plot
return_plot
                  return ggplot object
```

26 calculateMetaTable

#### **Details**

Currently we provide 2 ways to calculate highly variable genes: 1. high coeff of variance (COV) within groups:

First genes are binned (*nr\_expression\_groups*) into average expression groups and the COV for each gene is converted into a z-score within each bin. Genes with a z-score higher than the threshold (*zscore\_threshold*) are considered highly variable.

#### 2. high COV based on loess regression prediction:

A predicted COV is calculated for each gene using loess regression (COV~log(mean expression)) Genes that show a higher than predicted COV (difference\_in\_cov) are considered highly variable.

#### Value

giotto object highly variable genes appended to gene metadata (fDataDT)

### **Examples**

```
# 1. create giotto object
expr_path = system.file("extdata", "seqfish_field_expr.txt", package = 'Giotto')
loc_path = system.file("extdata", "seqfish_field_locs.txt", package = 'Giotto')
VC_small <- createGiottoObject(raw_exprs = expr_path, spatial_locs = loc_path)
# 2. normalize giotto
VC_small <- normalizeGiotto(gobject = VC_small, scalefactor = 6000)
VC_small <- addStatistics(gobject = VC_small)
# 3. highly variable genes detection
VC_small <- calculateHVG(gobject = VC_small)</pre>
```

calculateMetaTable

calculateMetaTable

#### **Description**

calculates the average gene expression for one or more (combined) annotation columns.

```
calculateMetaTable(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  metadata_cols = NULL,
  selected_genes = NULL
)
```

calculateMetaTableCells 27

### **Arguments**

```
gobject giotto object
expression_values
expression values to use
metadata_cols annotation columns found in pDataDT(gobject)
selected_genes subset of genes to use
```

### Value

data.table with average expression values for each gene per (combined) annotation

### **Examples**

```
calculateMetaTable(gobject)
```

```
{\tt calculateMetaTableCells}
```

calculateMetaTableCells

# Description

calculates the average metadata values for one or more (combined) annotation columns.

### Usage

```
calculateMetaTableCells(
  gobject,
  value_cols = NULL,
  metadata_cols = NULL,
  spat_enr_names = NULL
)
```

### **Arguments**

```
gobject giotto object
value_cols metadata or enrichment value columns to use
metadata_cols annotation columns found in pDataDT(gobject)
spat_enr_names which spatial enrichment results to include
```

# Value

data.table with average metadata values per (combined) annotation

#### **Examples**

```
calculateMetaTableCells(gobject)
```

28 cellProximityBarplot

```
calculate\_distance\_and\_weight \\ calculate\_distance\_and\_weight
```

### **Description**

```
calculate_distance_and_weight
```

# Usage

```
calculate_distance_and_weight(
  networkDT = NULL,
  sdimx = "sdimx",
  sdimy = "sdimy",
  sdimz = "sdimz",
  d2_or_d3 = c(2, 3)
)
```

# Arguments

```
networkDT spatial network as data.table
sdimx spatial dimension x
sdimy spatial dimension y
sdimz spatial dimension z
d2_or_d3 number of dimensions
```

 ${\tt cellProximityBarplot} \quad \textit{cellProximityBarplot}$ 

# Description

Create barplot from cell-cell proximity scores

```
cellProximityBarplot(
  gobject,
  CPscore,
  min_orig_ints = 5,
  min_sim_ints = 5,
  p_val = 0.05,
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "cellProximityBarplot"
)
```

#### **Arguments**

```
gobject
                  giotto object
CPscore
                  CPscore, output from cellProximityEnrichment()
                  filter on minimum original cell-cell interactions
min_orig_ints
                  filter on minimum simulated cell-cell interactions
min_sim_ints
                  p-value
p_val
show_plot
                  show plot
return_plot
                  return ggplot object
save_plot
                  directly save the plot [boolean]
                  list of saving parameters from all_plots_save_function
save_param
default_save_name
                  default save name for saving, don't change, change save_name in save_param
```

### **Details**

This function creates a barplot that shows the spatial proximity enrichment or depletion of cell type pairs.

#### Value

ggplot barplot

### **Examples**

```
cellProximityBarplot(CPscore)
```

```
cellProximityEnrichment
```

cellProximityEnrichment

### **Description**

Compute cell-cell interaction enrichment (observed vs expected)

#### **Arguments**

#### **Details**

Spatial proximity enrichment or depletion between pairs of cell types is calculated by calculating the observed over the expected frequency of cell-cell proximity interactions. The expected frequency is the average frequency calculated from a number of spatial network simulations. Each individual simulation is obtained by reshuffling the cell type labels of each node (cell) in the spatial network.

#### Value

List of cell Proximity scores (CPscores) in data.table format. The first data.table (raw\_sim\_table) shows the raw observations of both the original and simulated networks. The second data.table (enrichm\_res) shows the enrichment results.

#### **Examples**

```
cellProximityEnrichment(gobject)
```

```
cellProximityHeatmap cellProximityHeatmap
```

# **Description**

Create heatmap from cell-cell proximity scores

```
cellProximityHeatmap(
  gobject,
  CPscore,
  scale = T,
  order_cell_types = T,
  color_breaks = NULL,
  color_names = NULL,
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "cellProximityHeatmap")
```

cellProximityNetwork 31

#### **Arguments**

```
giotto object
gobject
CPscore
                  CPscore, output from cellProximityEnrichment()
scale
                  scale cell-cell proximity interaction scores
order_cell_types
                  order cell types based on enrichment correlation
                  numerical vector of length 3 to represent min, mean and maximum
color_breaks
                  character color vector of length 3
color_names
show_plot
                  show plot
return_plot
                  return ggplot object
                  directly save the plot [boolean]
save_plot
                  list of saving parameters from all_plots_save_function
save_param
default_save_name
                  default save name for saving, don't change, change save_name in save_param
```

#### **Details**

This function creates a heatmap that shows the spatial proximity enrichment or depletion of cell type pairs.

#### Value

ggplot heatmap

### **Examples**

```
cellProximityHeatmap(CPscore)
```

```
cellProximityNetwork cellProximityNetwork
```

# **Description**

Create network from cell-cell proximity scores

```
cellProximityNetwork(
  gobject,
  CPscore,
  remove_self_edges = FALSE,
  self_loop_strength = 0.1,
  color_depletion = "lightgreen",
  color_enrichment = "red",
  rescale_edge_weights = TRUE,
  edge_weight_range_depletion = c(0.1, 1),
  edge_weight_range_enrichment = c(1, 5),
  layout = c("Fruchterman", "DrL", "Kamada-Kawai"),
```

```
only_show_enrichment_edges = F,
edge_width_range = c(0.1, 2),
node_size = 4,
node_text_size = 6,
show_plot = NA,
return_plot = NA,
save_plot = NA,
save_plot = NA,
save_param = list(),
default_save_name = "cellProximityNetwork")
```

#### **Arguments**

```
gobject
                  giotto object
CPscore
                  CPscore, output from cellProximityEnrichment()
remove_self_edges
                  remove enrichment/depletion edges with itself
self_loop_strength
                  size of self-loops
color_depletion
                  color for depleted cell-cell interactions
color_enrichment
                  color for enriched cell-cell interactions
rescale_edge_weights
                  rescale edge weights (boolean)
edge_weight_range_depletion
                  numerical vector of length 2 to rescale depleted edge weights
edge_weight_range_enrichment
                  numerical vector of length 2 to rescale enriched edge weights
layout
                  layout algorithm to use to draw nodes and edges
only_show_enrichment_edges
                  show only the enriched pairwise scores
edge_width_range
                  range of edge width
                  size of nodes
node_size
node_text_size size of node labels
show_plot
                  show plot
return_plot
                  return ggplot object
save_plot
                  directly save the plot [boolean]
                  list of saving parameters from all_plots_save_function
save_param
default_save_name
                  default save name for saving, don't change, change save_name in save_param
```

#### **Details**

This function creates a network that shows the spatial proximity enrichment or depletion of cell type pairs.

cellProximitySpatPlot 33

#### Value

```
igraph plot
```

#### **Examples**

```
cellProximityNetwork(CPscore)
```

```
cellProximitySpatPlot cellProximitySpatPlot
```

### **Description**

Visualize 2D cell-cell interactions according to spatial coordinates in ggplot mode

### Usage

```
cellProximitySpatPlot(gobject, ...)
```

# **Arguments**

```
gobject
                  giotto object
                  additional parameters
interaction_name
                  cell-cell interaction name
cluster_column cluster column with cell clusters
                  x-axis dimension name (default = 'sdimx')
sdimx
sdimy
                  y-axis dimension name (default = 'sdimy')
cell_color
                  color for cells (see details)
cell_color_code
                  named vector with colors
color_as_factor
                  convert color column to factor
show_other_cells
                  decide if show cells not in network
show_network
                  show underlying spatial network
network_color
                  color of spatial network
spatial_network_name
                  name of spatial network to use
                  show spatial grid
show_grid
                  color of spatial grid
grid_color
spatial_grid_name
                  name of spatial grid to use
coord_fix_ratio
                  fix ratio between x and y-axis
show_legend
                  show legend
point_size_select
                  size of selected points
```

```
point_select_border_col
                  border color of selected points
point_select_border_stroke
                  stroke size of selected points
point_size_other
                  size of other points
point_other_border_col
                  border color of other points
\verb"point_other_border_stroke"
                  stroke size of other points
show_plot
                  show plots
return_plot
                  return ggplot object
save_plot
                  directly save the plot [boolean]
save_param
                  list of saving parameters from all_plots_save_function
default_save_name
                  default save name for saving, don't change, change save_name in save_param
```

#### **Details**

Description of parameters.

#### Value

ggplot

### See Also

```
cellProximitySpatPlot2D and cellProximitySpatPlot3D for 3D
```

### **Examples**

```
cellProximitySpatPlot(gobject)
```

```
cellProximitySpatPlot2D
```

cellProximitySpatPlot2D

# Description

Visualize 2D cell-cell interactions according to spatial coordinates in ggplot mode

```
cellProximitySpatPlot2D(
  gobject,
  interaction_name = NULL,
  cluster_column = NULL,
  sdimx = "sdimx",
  sdimy = "sdimy",
  cell_color = NULL,
```

```
cell_color_code = NULL,
  color_as_factor = T,
  show_other_cells = F,
  show_network = F,
  show_other_network = F,
 network_color = NULL,
  spatial_network_name = "Delaunay_network",
  show\_grid = F,
  grid_color = NULL,
  spatial_grid_name = "spatial_grid",
  coord_fix_ratio = 1,
  show_legend = T,
 point_size_select = 2,
 point_select_border_col = "black",
 point_select_border_stroke = 0.05,
  point_size_other = 1,
  point_alpha_other = 0.3,
 point_other_border_col = "lightgrey",
 point_other_border_stroke = 0.01,
  show_plot = NA,
 return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "cellProximitySpatPlot2D"
)
```

#### **Arguments**

```
giotto object
gobject
interaction_name
                  cell-cell interaction name
cluster_column cluster column with cell clusters
sdimx
                  x-axis dimension name (default = 'sdimx')
                  y-axis dimension name (default = 'sdimy')
sdimy
cell_color
                  color for cells (see details)
cell_color_code
                  named vector with colors
color_as_factor
                  convert color column to factor
show_other_cells
                  decide if show cells not in network
                  show spatial network of selected cells
show_network
show_other_network
                  show spatial network of not selected cells
network_color
                  color of spatial network
spatial_network_name
                  name of spatial network to use
show_grid
                  show spatial grid
grid_color
                  color of spatial grid
```

```
spatial_grid_name
                  name of spatial grid to use
coord_fix_ratio
                  fix ratio between x and y-axis
show_legend
                  show legend
point_size_select
                  size of selected points
point_select_border_col
                  border color of selected points
point_select_border_stroke
                  stroke size of selected points
point_size_other
                  size of other points
point_alpha_other
                  opacity of other points
{\tt point\_other\_border\_col}
                  border color of other points
point_other_border_stroke
                  stroke size of other points
show_plot
                  show plots
return_plot
                  return ggplot object
save_plot
                  directly save the plot [boolean]
save_param
                  list of saving parameters from all_plots_save_function
default_save_name
                  default save name for saving, don't change, change save_name in save_param
```

#### **Details**

Description of parameters.

### Value

ggplot

# **Examples**

cellProximitySpatPlot2D(gobject)

```
cellProximitySpatPlot3D
```

cell Proximity Spat Plot 2D

# Description

Visualize 3D cell-cell interactions according to spatial coordinates in plotly mode

#### Usage

```
cellProximitySpatPlot3D(
  gobject,
  interaction_name = NULL,
  cluster_column = NULL,
  sdimx = "sdimx",
  sdimy = "sdimy",
  sdimz = "sdimz",
  cell_color = NULL,
  cell_color_code = NULL,
  color_as_factor = T,
  show_other_cells = T,
  show_network = T,
  show_other_network = F,
  network_color = NULL,
  spatial_network_name = "Delaunay_network",
  show\_grid = F,
  grid_color = NULL,
  spatial_grid_name = "spatial_grid",
  show_legend = T,
  point_size_select = 4,
  point_size_other = 2,
  point_alpha_other = 0.5,
  axis_scale = c("cube", "real", "custom"),
  custom_ratio = NULL,
  x_ticks = NULL,
  y_ticks = NULL,
  z_ticks = NULL,
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "cellProximitySpatPlot3D",
)
```

```
gobject
                  giotto object
interaction_name
                  cell-cell interaction name
cluster_column cluster column with cell clusters
sdimx
                  x-axis dimension name (default = 'sdimx')
                  y-axis dimension name (default = 'sdimy')
sdimy
                  z-axis dimension name (default = 'sdimz')
sdimz
cell_color
                  color for cells (see details)
cell_color_code
                  named vector with colors
color_as_factor
                  convert color column to factor
```

```
show_other_cells
                  decide if show cells not in network
                  show spatial network of selected cells
show_network
show_other_network
                  show spatial network of not selected cells
network_color
                  color of spatial network
spatial_network_name
                  name of spatial network to use
show_grid
                  show spatial grid
grid_color
                  color of spatial grid
spatial_grid_name
                  name of spatial grid to use
show_legend
                  show legend
point_size_select
                  size of selected points
point_size_other
                  size of other points
point_alpha_other
                  opacity of other points
axis_scale
                  scale of axis
custom_ratio
                  custom ratio of axes
x_ticks
                  ticks on x-axis
                  ticks on y-axis
y_ticks
                  ticks on z-axis
z_ticks
                  show plots
show_plot
return_plot
                  return plotly object
                  directly save the plot [boolean]
save_plot
                  list of saving parameters from all_plots_save_function
save_param
default_save_name
                  default save name for saving, don't change, change save_name in save_param
                  additional parameters
. . .
```

#### **Details**

Description of parameters.

### Value

plotly

# **Examples**

cellProximitySpatPlot3D(gobject)

cellProximityVisPlot 39

cellProximityVisPlot cellProximityVisPlot

# Description

Visualize cell-cell interactions according to spatial coordinates

# Usage

```
cellProximityVisPlot(
  gobject,
  interaction_name = NULL,
  cluster_column = NULL,
  sdimx = NULL,
  sdimy = NULL,
  sdimz = NULL,
  cell_color = NULL,
  cell_color_code = NULL,
  color_as_factor = T,
  show_other_cells = F,
  show_network = F,
  show_other_network = F,
  network_color = NULL,
  spatial_network_name = "Delaunay_network",
  show_grid = F,
  grid_color = NULL,
  spatial_grid_name = "spatial_grid",
  coord_fix_ratio = 1,
  show_legend = T,
  point_size_select = 2,
  point_select_border_col = "black",
  point_select_border_stroke = 0.05,
  point_size_other = 1,
  point_alpha_other = 0.3,
  point_other_border_col = "lightgrey",
  point_other_border_stroke = 0.01,
  axis_scale = c("cube", "real", "custom"),
  custom_ratio = NULL,
  x_ticks = NULL,
  y_ticks = NULL,
  z_{ticks} = NULL,
  plot_method = c("ggplot", "plotly"),
)
```

sdimx x-axis dimension name (default = 'sdimx')
sdimy y-axis dimension name (default = 'sdimy')
sdimz z-axis dimension name (default = 'sdimz')

cell\_color color for cells (see details)

cell\_color\_code

named vector with colors

color\_as\_factor

convert color column to factor

show\_network show underlying spatial network

network\_color color of spatial network

spatial\_network\_name

name of spatial network to use

show\_grid show spatial grid grid\_color color of spatial grid

spatial\_grid\_name

name of spatial grid to use

coord\_fix\_ratio

fix ratio between x and y-axis

show\_legend show legend

point\_size\_select

size of selected points

point\_select\_border\_col

border color of selected points

point\_select\_border\_stroke

stroke size of selected points

point\_size\_other

size of other points

point\_other\_border\_col

border color of other points

point\_other\_border\_stroke

stroke size of other points

### **Details**

Description of parameters.

# Value

ggplot or plotly

### **Examples**

cellProximityVisPlot(gobject)

#### **Description**

Visualize 2D cell-cell interactions according to spatial coordinates in ggplot mode

#### Usage

```
cellProximityVisPlot_2D_ggplot(
  gobject,
  interaction_name = NULL,
  cluster_column = NULL,
  sdimx = NULL,
  sdimy = NULL,
  cell_color = NULL,
  cell_color_code = NULL,
  color_as_factor = T,
  show_other_cells = F,
  show_network = F,
  show_other_network = F,
  network_color = NULL,
  spatial_network_name = "Delaunay_network",
  show\_grid = F,
  grid_color = NULL,
  spatial_grid_name = "spatial_grid",
  coord_fix_ratio = 1,
  show_legend = T,
  point_size_select = 2,
  point_select_border_col = "black",
  point_select_border_stroke = 0.05,
  point_size_other = 1,
  point_alpha_other = 0.3,
  point_other_border_col = "lightgrey",
  point_other_border_stroke = 0.01,
)
```

```
color_as_factor
```

convert color column to factor

show\_other\_cells

decide if show cells not in network

show\_network show underlying spatial network

network\_color color of spatial network

 $spatial\_network\_name$ 

name of spatial network to use

show\_grid show spatial grid

grid\_color color of spatial grid

spatial\_grid\_name

name of spatial grid to use

coord\_fix\_ratio

fix ratio between x and y-axis

show\_legend show legend

point\_size\_select

size of selected points

point\_select\_border\_col

border color of selected points

point\_select\_border\_stroke

stroke size of selected points

point\_size\_other

size of other points

point\_other\_border\_col

border color of other points

point\_other\_border\_stroke

stroke size of other points

#### **Details**

Description of parameters.

# Value

ggplot

# **Examples**

cellProximityVisPlot\_2D\_ggplot(gobject)

```
cell Proximity VisPlot\_2D\_plotly \\ cell Proximity VisPlot\_2D\_plotly
```

# Description

Visualize 2D cell-cell interactions according to spatial coordinates in plotly mode

### Usage

```
cellProximityVisPlot_2D_plotly(
  gobject,
  interaction_name = NULL,
  cluster_column = NULL,
  sdimx = NULL,
  sdimy = NULL,
  cell_color = NULL,
  cell_color_code = NULL,
  color_as_factor = T,
  show_other_cells = F,
  show_network = F,
  show_other_network = F,
  network_color = NULL,
  spatial_network_name = "Delaunay_network",
  show\_grid = F,
  grid_color = NULL,
  spatial_grid_name = "spatial_grid",
  show_legend = T,
  point_size_select = 2,
  point_size_other = 1,
  point_alpha_other = 0.3,
  axis_scale = c("cube", "real", "custom"),
  custom_ratio = NULL,
  x_ticks = NULL,
  y_ticks = NULL,
)
```

```
color_as_factor
                  convert color column to factor
show_other_cells
                  decide if show cells not in network
                  show underlying spatial network
show_network
network_color
                  color of spatial network
spatial_network_name
                  name of spatial network to use
show_grid
                  show spatial grid
                  color of spatial grid
grid_color
spatial_grid_name
                  name of spatial grid to use
show_legend
                  show legend
point_size_select
                  size of selected points
coord_fix_ratio
                  fix ratio between x and y-axis
```

### **Details**

Description of parameters.

#### Value

plotly

#### **Examples**

```
cell Proximity VisPlot\_2D\_plotly (gobject)
```

### **Description**

Visualize 3D cell-cell interactions according to spatial coordinates in plotly mode

### Usage

```
cellProximityVisPlot_3D_plotly(
  gobject,
  interaction_name = NULL,
  cluster_column = NULL,
  sdimx = NULL,
  sdimy = NULL,
  sdimz = NULL,
  cell_color = NULL,
  cell_color_code = NULL,
```

```
color_as_factor = T,
show_other_cells = F,
show_network = F,
show_other_network = F,
network_color = NULL,
spatial_network_name = "Delaunay_network",
show\_grid = F,
grid_color = NULL,
spatial_grid_name = "spatial_grid",
show_legend = T,
point_size_select = 2,
point_size_other = 1,
point_alpha_other = 0.5,
axis_scale = c("cube", "real", "custom"),
custom_ratio = NULL,
x_ticks = NULL,
y_ticks = NULL,
z_{ticks} = NULL,
```

```
gobject
                  giotto object
interaction_name
                  cell-cell interaction name
cluster_column cluster column with cell clusters
sdimx
                  x-axis dimension name (default = 'sdimx')
                  y-axis dimension name (default = 'sdimy')
sdimy
                  z-axis dimension name (default = 'sdimz')
sdimz
cell color
                  color for cells (see details)
cell_color_code
                  named vector with colors
color_as_factor
                  convert color column to factor
show_other_cells
                  decide if show cells not in network
                  show underlying spatial network
show_network
                  color of spatial network
network_color
spatial_network_name
                  name of spatial network to use
                  show spatial grid
show_grid
grid_color
                  color of spatial grid
spatial_grid_name
                  name of spatial grid to use
show_legend
                  show legend
point_size_select
                  size of selected points
coord_fix_ratio
                  fix ratio between x and y-axis
```

### **Details**

Description of parameters.

### Value

plotly

### **Examples**

```
cellProximityVisPlot_3D_plotly(gobject)
```

 ${\tt changeGiottoInstructions}$ 

change Giot to Instructions

# **Description**

Function to change one or more instructions from giotto object

# Usage

```
changeGiottoInstructions(
  gobject,
  params = NULL,
  new_values = NULL,
  return_gobject = TRUE
)
```

# Arguments

```
gobject giotto object

params parameter(s) to change

new_values new value(s) for parameter(s)

return_gobject (boolean) return giotto object
```

#### Value

giotto object with one or more changed instructions

# **Examples**

changeGiottoInstructions()

changeImageBg 47

changeImageBg	changeImageBg	
---------------	---------------	--

### **Description**

Function to change the background color of a magick image plot to another color

### Usage

```
changeImageBg(
  mg_object,
  bg_color,
  perc_range = 10,
  new_color = "#FFFFFF",
  new_name = NULL
)
```

### **Arguments**

```
mg_object magick image or giotto image object
bg_color estimated current background color
perc_range range around estimated background color to include (percentage)
new_color new background color
new_name change name of Giotto image
```

### Value

magick image or giotto image object with updated background color

#### **Examples**

```
changeImageBg(mg_object)
```

clusterCells clusterCells

# Description

cluster cells using a variety of different methods

# Usage

```
clusterCells(
  gobject,
  cluster_method = c("leiden", "louvain_community", "louvain_multinet", "randomwalk",
        "sNNclust", "kmeans", "hierarchical"),
  name = "cluster_name",
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
```

48 clusterCells

```
pyth_leid_resolution = 1,
     pyth_leid_weight_col = "weight",
     pyth_leid_part_type = c("RBConfigurationVertexPartition",
        "ModularityVertexPartition"),
     pyth_leid_init_memb = NULL,
     pyth_leid_iterations = 1000,
     pyth_louv_resolution = 1,
     pyth_louv_weight_col = NULL,
     python_louv_random = F,
     python_path = NULL,
     louvain_gamma = 1,
     louvain\_omega = 1,
     walk\_steps = 4,
     walk_clusters = 10,
     walk_weights = NA,
     sNNclust_k = 20,
     sNNclust_eps = 4,
     sNNclust_minPts = 16,
     borderPoints = TRUE,
     expression_values = c("normalized", "scaled", "custom"),
     genes_to_use = NULL,
     dim_reduction_to_use = c("cells", "pca", "umap", "tsne"),
     dim_reduction_name = "pca",
     dimensions_to_use = 1:10,
     distance_method = c("original", "pearson", "spearman", "euclidean", "maximum",
        "manhattan", "canberra", "binary", "minkowski"),
     km_centers = 10,
     km_iter_max = 100,
     km_nstart = 1000,
     km_algorithm = "Hartigan-Wong",
     hc_agglomeration_method = c("ward.D2", "ward.D", "single", "complete", "average",
        "mcquitty", "median", "centroid"),
     hc_k = 10,
     hc_h = NULL
     return_gobject = TRUE,
     set\_seed = T,
     seed_number = 1234
   )
Arguments
   gobject
                   giotto object
   cluster_method community cluster method to use
                   name for new clustering result
   name
   nn_network_to_use
                   type of NN network to use (kNN vs sNN)
   network_name
                   name of NN network to use
   pyth_leid_resolution
                   resolution for leiden
   pyth_leid_weight_col
                   column to use for weights
```

clusterCells 49

pyth\_leid\_part\_type

partition type to use

pyth\_leid\_init\_memb

initial membership

pyth\_leid\_iterations

number of iterations

pyth\_louv\_resolution

resolution for louvain

pyth\_louv\_weight\_col

python louvain param: weight column

python\_louv\_random

python louvain param: random

python\_path specify specific path to python if required

louvain\_gamma louvain param: gamma or resolution

louvain\_omega louvain param: omega

walk\_steps randomwalk: number of steps
walk\_clusters randomwalk: number of clusters
walk\_weights randomwalk: weight column
sNNclust\_k SNNclust: k neighbors to use

sNNclust\_eps SNNclust: epsilon

sNNclust\_minPts

SNNclust: min points

borderPoints SNNclust: border points

expression\_values

expression values to use

$$\label{eq:genes_to_use} \begin{split} & \text{genes\_to\_use} & = NULL, \\ & \text{dim\_reduction\_to\_use} \end{split}$$

dimension reduction to use

dim\_reduction\_name

name of reduction 'pca',

dimensions\_to\_use

dimensions to use

 $distance\_method$ 

distance method

km\_centers kmeans centers km\_iter\_max kmeans iterations

km\_nstart kmeans random starting points

km\_algorithm kmeans algorithm

hc\_agglomeration\_method

hierarchical clustering method

hc\_k hierachical number of clusters

hc\_h hierarchical tree cutoff

return\_gobject | boolean: return giotto object (default = TRUE)

set\_seed set seed

seed\_number number for seed

#### **Details**

Wrapper for the different clustering methods.

#### Value

giotto object with new clusters appended to cell metadata

#### See Also

 $\label{lem:cluster_doLouvainCluster_multinet} do Louvain Cluster\_community, do Louvain Cluster\_multinet, do Louvain Cluster, do Random Walk Cluster, do SNN Cluster, do Kmeans, do H clust Cluster, do Louvain Cluster, do Louva$ 

### **Examples**

```
clusterCells(gobject)
```

clusterSpatialCorGenes

cluster Spatial Cor Genes

### **Description**

Cluster based on spatially correlated genes

### Usage

```
clusterSpatialCorGenes(
  spatCorObject,
  name = "spat_clus",
  hclust_method = "ward.D",
  k = 10,
  return_obj = TRUE
)
```

### **Arguments**

spatCorObject spatial correlation object

name name for spatial clustering results
hclust\_method method for hierarchical clustering
k number of clusters to extract

return\_obj return spatial correlation object (spatCorObject)

#### Value

spatCorObject or cluster results

#### **Examples**

```
clusterSpatialCorGenes(gobject)
```

combCCcom 51

combCCcom combCCcom

# Description

Combine spatial and expression based cell-cell communication data.tables

# Usage

```
combCCcom(
  spatialCC,
  exprCC,
  min_lig_nr = 3,
  min_rec_nr = 3,
  min_padj_value = 1,
  min_log2fc = 0,
  min_av_diff = 0
)
```

### **Arguments**

```
spatialCC spatial cell-cell communication scores
exprCC expression cell-cell communication scores
min_lig_nr minimum number of ligand cells
min_rec_nr minimum number of receptor cells
min_padj_value minimum adjusted p-value
min_log2fc minimum log2 fold-change
min_av_diff minimum average expression difference
```

### Value

combined data.table with spatial and expression communication data

### **Examples**

```
combCCcom(gobject)
```

```
combine \textit{CellProximityGenes} \\ combine \textit{CellProximityGenes}
```

# Description

Combine CPG scores in a pairwise manner.

#### Usage

```
combineCellProximityGenes(
  cpgObject,
  selected_ints = NULL,
  selected_genes = NULL,
  specific_genes_1 = NULL,
  specific_genes_2 = NULL,
  min_cells = 5,
  min_int_cells = 3,
  min_fdr = 0.05,
  min_spat_diff = 0,
  min_log2_fc = 0.5,
  do_parallel = TRUE,
  cores = NA,
  verbose = T
)
```

### **Arguments**

```
cpgObject
                  cell proximity gene score object
                  subset of selected cell-cell interactions (optional)
selected_ints
selected_genes subset of selected genes (optional)
specific_genes_1
                  specific geneset combo (need to position match specific_genes_2)
specific_genes_2
                  specific geneset combo (need to position match specific_genes_1)
min_cells
                  minimum number of target cell type
min_int_cells
                  minimum number of interacting cell type
min_fdr
                  minimum adjusted p-value
                  minimum absolute spatial expression difference
min_spat_diff
                  minimum absolute log2 fold-change
min_log2_fc
                  run calculations in parallel with mclapply
do_parallel
                  number of cores to use if do_parallel = TRUE
cores
verbose
                  verbose
```

### Value

cpgObject that contains the filtered differential gene scores

# Examples

```
{\tt combineCellProximityGenes(gobject)}
```

 $combine \verb|CellProximityGenes_per_interaction| \\ combine CellProximity Genes\_per\_interaction|$ 

# **Description**

Combine CPG scores per interaction

# Usage

```
combineCellProximityGenes_per_interaction(
  cpgObject,
  sel_int,
  selected_genes = NULL,
  specific_genes_1 = NULL,
  specific_genes_2 = NULL,
  min_cells = 5,
  min_int_cells = 3,
  min_fdr = 0.05,
  min_spat_diff = 0,
  min_log2_fc = 0.5
)
```

#### **Examples**

 ${\tt combineCellProximityGenes\_per\_interaction()}$ 

combineCPG

combineCPG

# Description

Combine CPG scores in a pairwise manner.

# Usage

```
combineCPG(
  cpgObject,
  selected_ints = NULL,
  selected_genes = NULL,
  specific_genes_1 = NULL,
  specific_genes_2 = NULL,
  min_cells = 5,
  min_int_cells = 3,
  min_fdr = 0.05,
  min_spat_diff = 0,
  min_log2_fc = 0.5,
  do_parallel = TRUE,
  cores = NA,
  verbose = T
)
```

54 combineMetadata

#### **Arguments**

```
cpgObject
                  cell proximity gene score object
                  subset of selected cell-cell interactions (optional)
selected_ints
selected_genes subset of selected genes (optional)
specific_genes_1
                  specific geneset combo (need to position match specific_genes_2)
specific_genes_2
                  specific geneset combo (need to position match specific_genes_1)
min_cells
                  minimum number of target cell type
min_int_cells
                  minimum number of interacting cell type
                  minimum adjusted p-value
min_fdr
min_spat_diff
                  minimum absolute spatial expression difference
min_log2_fc
                  minimum absolute log2 fold-change
                  run calculations in parallel with mclapply
do_parallel
                  number of cores to use if do_parallel = TRUE
cores
verbose
                  verbose
```

#### Value

cpgObject that contains the filtered differential gene scores

#### **Examples**

```
combineCPG(gobject)
```

combineMetadata combineMetadata

#### **Description**

This function combines the cell metadata with spatial locations and enrichment results from createSpatialEnrich

#### Usage

```
combineMetadata(gobject, spat_enr_names = NULL)
```

### **Arguments**

```
gobject Giotto object
spat_enr_names names of spatial enrichment results to include
```

### Value

Extended cell metadata in data.table format.

# **Examples**

```
combineMetadata(gobject)
```

convertEnsemblToGeneSymbol

convert Ensembl To Gene Symbol

# Description

This function convert ensembl gene IDs from a matrix to official gene symbols

### Usage

```
convertEnsemblToGeneSymbol(matrix, species = c("mouse", "human"))
```

# Arguments

matrix an expression matrix with ensembl gene IDs as rownames

species species to use for gene symbol conversion

#### **Details**

This function requires that the biomaRt library is installed

#### Value

expression matrix with gene symbols as rownames

# **Examples**

```
convertEnsemblToGeneSymbol(matrix)
```

### **Description**

converts a magick image object to a data.table

### Usage

```
convert_mgImage_to_array_DT(mg_object)
```

# Arguments

mg\_object magick image or Giotto image object

### Value

data.table with image pixel information

56 createCrossSection

createCrossSection

createCrossSection

#### **Description**

Create a virtual 2D cross section.

#### Usage

```
createCrossSection(
  gobject.
 name = "cross_section",
  spatial_network_name = "Delaunay_network",
  thickness_unit = c("cell", "natural"),
  slice_thickness = 2,
  cell_distance_estimate_method = "mean",
  extend_ratio = 0.2,
 method = c("equation", "3 points", "point and norm vector",
    "point and two plane vectors"),
  equation = NULL,
  point1 = NULL,
 point2 = NULL,
  point3 = NULL,
 normVector = NULL,
 planeVector1 = NULL,
 planeVector2 = NULL,
 mesh\_grid\_n = 20,
  return_gobject = TRUE
)
```

### **Arguments**

```
gobject
                   giotto object
name
                   name of cress section object. (default = cross_section)
spatial_network_name
                   name of spatial network object. (default = Delaunay_network)
thickness_unit unit of the virtual section thickness. If "cell", average size of the observed
                   cells is used as length unit. If "natural", the unit of cell location coordinates
                   is used.(default = cell)
slice_thickness
                   thickness of slice
{\tt cell\_distance\_estimate\_method}
                   method to estimate average distance between neighboring cells. (default = mean)
                   deciding the span of the cross section meshgrid, as a ratio of extension compared
extend_ratio
                   to the borders of the vitural tissue section. (default = 0.2)
method
                   method to define the cross section plane. If equation, the plane is defined by
                   a four element numerical vector (equation) in the form of c(A,B,C,D), corre-
```

sponding to a plane with equation Ax+By+Cz=D. If 3 points, the plane is define by the coordinates of 3 points, as given by point1, point2, and point3. If point

createGiottoImage 57

	and norm vector, the plane is defined by the coordinates of one point (point1) in the plane and the coordinates of one norm vector (normVector) to the plane. If point and two plane vector, the plane is defined by the coordinates of one point (point1) in the plane and the coordinates of two vectors (planeVector1, planeVector2) in the plane. (default = equation)
equation	equation required by method "equation".equations needs to be a numerical vector of length 4, in the form of $c(A,B,C,D)$ , which defines plane $Ax+By+Cz=D$ .
point1	coordinates of the first point required by method "3 points", "point and norm vector", and "point and two plane vectors".
point2	coordinates of the second point required by method "3 points"
point3	coordinates of the third point required by method "3 points"
normVector	coordinates of the norm vector required by method "point and norm vector"
planeVector1	coordinates of the first plane vector required by method "point and two plane vectors"
planeVector2	coordinates of the second plane vector required by method "point and two plane vectors"
mesh_grid_n	numer of meshgrid lines to generate along both directions for the cross section plane.
return_gobject	boolean: return giotto object (default = TRUE)

### **Details**

Creates a virtual 2D cross section object for a given spatial network object. The users need to provide the definition of the cross section plane (see method).

#### Value

giotto object with updated spatial network slot

createGiottoImage

# Description

Creates a giotto image that can be added to a Giotto object and/or used to add an image to the spatial plotting functions

### Usage

```
createGiottoImage(
  gobject = NULL,
  spatial_locs = NULL,
  mg_object,
  name = "image",
  xmax_adj = 0,
  xmin_adj = 0,
  ymax_adj = 0,
  ymin_adj = 0
)
```

58 createGiottoInstructions

# Arguments

```
gobject
                  giotto object
spatial_locs
                  spatial locations (alternative if giobject = NULL)
mg_object
                  magick image object
name
                  name for the image
                  adjustment of the maximum x-value to align the image
xmax_adj
xmin_adj
                  adjustment of the minimum x-value to align the image
                  adjustment of the maximum y-value to align the image
ymax_adj
                  adjustment of the minimum y-value to align the image
ymin_adj
```

#### Value

a giotto image object

### **Examples**

```
createGiottoImage(mg_object)
```

createGiottoInstructions

createGiottoInstructions

#### **Description**

Function to set global instructions for giotto functions

#### Usage

```
createGiottoInstructions(
  python_path = NULL,
  show_plot = NULL,
  return_plot = NULL,
  save_plot = NULL,
  save_dir = NULL,
  plot_format = NULL,
  dpi = NULL,
  units = NULL,
  height = NULL,
  width = NULL
```

```
python_path path to python binary to use
show_plot print plot to console, default = TRUE
return_plot return plot as object, default = TRUE
save_plot automatically save plot, dafault = FALSE
save_dir path to directory where to save plots
```

createGiottoObject 59

```
plot_format format of plots (defaults to png)
dpi resolution for raster images
units units of format (defaults to in)
height height of plots
width width of plots
```

#### Value

named vector with giotto instructions

#### See Also

More online information can be found here  $https://rubd.github.io/Giotto\_site/articles/instructions\_and\_plotting.html$ 

### **Examples**

```
createGiottoInstructions()
```

createGiottoObject
create Giotto object

### **Description**

Function to create a giotto object

### Usage

```
createGiottoObject(
  raw_exprs,
  spatial_locs = NULL,
  norm_expr = NULL,
  norm_scaled_expr = NULL,
  custom_expr = NULL,
  cell_metadata = NULL,
  gene_metadata = NULL,
  spatial_network = NULL,
  spatial_network_name = NULL,
  spatial_grid = NULL,
  spatial_grid_name = NULL,
  spatial_enrichment = NULL,
  spatial_enrichment_name = NULL,
  dimension_reduction = NULL,
  nn_network = NULL,
  images = NULL,
  offset_file = NULL,
  instructions = NULL,
  cores = NA
)
```

60 createGiottoObject

#### Arguments

raw\_exprs matrix with raw expression counts [required]

spatial\_locs data.table or data.frame with coordinates for cell centroids

norm\_expr normalized expression values

norm\_scaled\_expr

scaled expression values

custom\_expr custom expression values cell\_metadata cell annotation metadata gene\_metadata gene annotation metadata

spatial\_network

list of spatial network(s)

spatial\_network\_name

list of spatial network name(s)

spatial\_grid list of spatial grid(s)

spatial\_grid\_name

list of spatial grid name(s)

spatial\_enrichment

list of spatial enrichment score(s) for each spatial region

spatial\_enrichment\_name

list of spatial enrichment name(s)

dimension\_reduction

list of dimension reduction(s)

nn\_network list of nearest neighbor network(s)

images list of images

offset\_file file used to stitch fields together (optional)

instructions list of instructions or output result from createGiottoInstructions cores how many cores or threads to use to read data if paths are provided

#### **Details**

[**Requirements**] To create a giotto object you need to provide at least a matrix with genes as row names and cells as column names. This matrix can be provided as a base matrix, sparse Matrix, data.frame, data.table or as a path to any of those. To include spatial information about cells (or regions) you need to provide a matrix, data.table or data.frame (or path to them) with coordinates for all spatial dimensions. This can be 2D (x and y) or 3D (x, y, x). The row order for the cell coordinates should be the same as the column order for the provided expression data.

[Instructions] Additionally an instruction file, generated manually or with createGiottoInstructions can be provided to instructions, if not a default instruction file will be created for the Giotto object.

[Multiple fields] In case a dataset consists of multiple fields, like seqFISH+ for example, an offset file can be provided to stitch the different fields together. stitchFieldCoordinates can be used to generate such an offset file.

[**Processed data**] Processed count data, such as normalized data, can be provided using one of the different expression slots (norm\_expr, norm\_scaled\_expr, custom\_expr).

[Metadata] Cell and gene metadata can be provided using the cell and gene metadata slots. This data can also be added afterwards using the addGeneMetadata or addCellMetadata functions.

[Other information] Additional information can be provided through the appropriate slots:

- · spatial networks
- · spatial girds
- spatial enrichments
- · dimensions reductions
- nearest neighbours networks
- · images

#### Value

giotto object

### **Examples**

```
createGiottoObject(raw_exprs, spatial_locs)
```

createGiottoVisiumObject

createGiottoVisiumObject

# Description

creates Giotto object directly from a 10X visium folder

### Usage

```
createGiottoVisiumObject(
  visium_dir = NULL,
  expr_data = c("raw", "filter"),
  gene_column_index = 1,
  png_name = NULL,
  xmax_adj = 0,
  xmin_adj = 0,
  ymax_adj = 0,
  ymin_adj = 0,
  instructions = NULL,
  cores = NA
)
```

```
visium_dir
                  path to the 10X visium directory [required]
expr_data
                  raw or filtered data (see details)
gene_column_index
                  which column index to select (see details)
                  select name of png to use (see details)
png_name
xmax_adj
                  adjustment of the maximum x-value to align the image
                  adjustment of the minimum x-value to align the image
xmin_adj
                  adjustment of the maximum y-value to align the image
ymax_adj
                  adjustment of the minimum y-value to align the image
ymin_adj
instructions
                  list of instructions or output result from createGiottoInstructions
                  how many cores or threads to use to read data if paths are provided
cores
```

62 createHeatmap\_DT

#### **Details**

expr\_data: raw will take expression data from raw\_feature\_bc\_matrix and filter from filtered\_feature\_bc\_matrix

- gene\_column\_index: which gene identifiers (names) to use if there are multiple columns (e.g. ensemble and gene symbol)
- png\_name: by default the first png will be selected, provide the png name to override this (e.g. myimage.png)

#### Value

giotto object

#### **Examples**

```
createGiottoVisiumObject(visium_dir)
```

createHeatmap\_DT

 $createHeatmap\_DT$ 

# Description

creates order for clusters

# Usage

```
createHeatmap_DT(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  genes,
  cluster_column = NULL,
  cluster_order = c("size", "correlation", "custom"),
  cluster_custom_order = NULL,
  cluster_cor_method = "pearson",
  cluster_hclust_method = "ward.D",
  gene_order = c("correlation", "custom"),
  gene_custom_order = NULL,
  gene_cor_method = "pearson",
  gene_hclust_method = "complete"
)
```

```
gobject giotto object
expression_values
expression values to use
genes genes to use
cluster_column name of column to use for clusters
cluster_order method to determine cluster order
cluster_custom_order
custom order for clusters
```

createMetagenes 63

### **Details**

Creates input data.tables for plotHeatmap function.

#### Value

list

### **Examples**

```
createHeatmap_DT(gobject)
```

createMetagenes

createMetagenes

### **Description**

This function creates an average metagene for gene clusters.

### Usage

```
createMetagenes(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  gene_clusters,
  name = "metagene",
  return_gobject = TRUE
)
```

```
gobject Giotto object
expression_values
expression values to use
gene_clusters numerical vector with genes as names
name name of the metagene results
return_gobject return giotto object
```

64 createNearestNetwork

#### **Details**

```
An example for the 'gene_clusters' could be like this: cluster_vector = c(1, 1, 2, 2); names(cluster_vector) = c('geneA', 'geneB', 'geneC', 'geneD')
```

#### Value

giotto object

## **Examples**

```
createMetagenes(gobject)
```

createNearestNetwork createNearestNetwork

# Description

create a nearest neighbour (NN) network

# Usage

```
createNearestNetwork(
  gobject,
  type = c("sNN", "kNN"),
  dim_reduction_to_use = "pca",
  dim_reduction_name = "pca",
  dimensions_to_use = 1:10,
  genes_to_use = NULL,
  expression_values = c("normalized", "scaled", "custom"),
  name = "sNN.pca",
  return_gobject = TRUE,
  k = 30,
  minimum_shared = 5,
  top_shared = 3,
  verbose = T,
  ...
)
```

createNearestNetwork 65

name arbitrary name for NN network

return\_gobject boolean: return giotto object (default = TRUE)

k number of k neighbors to use minimum\_shared minimum shared neighbors

top\_shared keep at ...
verbose be verbose

... additional parameters for kNN and sNN functions from dbscan

#### **Details**

This function creates a k-nearest neighbour (kNN) or shared nearest neighbour (sNN) network based on the provided dimension reduction space. To run it directly on the gene expression matrix set  $dim\_reduction\_to\_use = NULL$ .

See also kNN and sNN for more information about how the networks are created.

#### Output for kNN:

• from: cell\_ID for source cell

• to: cell\_ID for target cell

• distance: distance between cells

• weight: weight = 1/(1 + distance)

### Output for sNN:

• from: cell\_ID for source cell

• to: cell\_ID for target cell

• distance: distance between cells

• weight: 1/(1 + distance)

• shared: number of shared neighbours

• rank: ranking of pairwise cell neighbours

For sNN networks two additional parameters can be set:

- minimum\_shared: minimum number of shared neighbours needed
- top\_shared: keep this number of the top shared neighbours, irrespective of minimum\_shared setting

#### Value

giotto object with updated NN network

#### **Examples**

createNearestNetwork(gobject)

```
create Spatial Delaunay Network \\ create Spatial Delaunay Network
```

# Description

Create a spatial Delaunay network based on cell centroid physical distances.

# Usage

```
createSpatialDelaunayNetwork(
  gobject,
  method = c("deldir", "delaunayn_geometry", "RTriangle"),
  dimensions = "all",
  name = "Delaunay_network",
  maximum_distance = "auto",
  minimum_k = 0,
  options = "Pp",
  Y = TRUE,
  j = TRUE,
  S = 0,
  verbose = T,
  return_gobject = TRUE,
  ...
)
```

gobject	giotto object	
dimensions	which spatial dimensions to use (default = all)	
name	name for spatial network (default = 'delaunay_network')	
maximum_distance		
	distance cuttof for Delaunay neighbors to consider. If "auto", "upper wisker" value of the distance vector between neighbors is used; see the boxplotgraphics documentation for more details.(default = "auto")	
minimum_k	minimum number of neighbours if maximum_distance != NULL	
options	(geometry) String containing extra control options for the underlying Qhull command; see the Qhull documentation (/doc/qhull/html/qdelaun.html) for the available options. (default = 'Pp', do not report precision problems)	
Υ	(RTriangle) If TRUE prohibits the insertion of Steiner points on the mesh boundary.	
j	(RTriangle) If TRUE jettisons vertices that are not part of the final triangulation from the output.	
S	(RTriangle) Specifies the maximum number of added Steiner points.	
verbose	verbose	
return_gobject	boolean: return giotto object (default = TRUE)	
	Other parameters of the triangulate function	

createSpatialEnrich 67

#### **Details**

Creates a spatial Delaunay network as explained in delaunayn (default), deldir, or triangulate.

#### Value

giotto object with updated spatial network slot

### **Examples**

```
createSpatialDelaunayNetwork(gobject)
```

 $create Spatial Enrich \qquad \textit{create Spatial Enrich}$ 

## **Description**

Function to calculate gene signature enrichment scores per spatial position using a hypergeometric test.

### Usage

```
createSpatialEnrich(
  gobject,
  enrich_method = c("PAGE", "rank", "hypergeometric"),
  sign_matrix,
  expression_values = c("normalized", "scaled", "custom"),
  reverse_log_scale = TRUE,
  logbase = 2,
  p_value = FALSE,
  n_genes = 100,
  n_times = 1000,
  top_percentage = 5,
  output_enrichment = c("original", "zscore"),
  name = "PAGE",
  return_gobject = TRUE
)
```

```
gobject
                  Giotto object
                  method for gene signature enrichment calculation
enrich_method
                  Matrix of signature genes for each cell type / process
sign_matrix
expression_values
                  expression values to use
reverse_log_scale
                  reverse expression values from log scale
logbase
                  log base to use if reverse_log_scale = TRUE
p_value
                  calculate p-value (default = FALSE)
                  (page) number of genes of permutation iterations to calculate p-value
n_genes
```

68 createSpatialGrid

#### **Details**

For details see the individual functions:

```
PAGE: PAGEEnrichPAGE: rankEnrichPAGE: hyperGeometricEnrich
```

### Value

Giotto object or enrichment results if return\_gobject = FALSE

### **Examples**

```
createSpatialEnrich(gobject)
```

#### **Description**

Create a spatial grid.

### Usage

```
createSpatialGrid(
  gobject,
  sdimx_stepsize = NULL,
  sdimy_stepsize = NULL,
  sdimz_stepsize = NULL,
  minimum_padding = 1,
  name = "spatial_grid",
  return_gobject = TRUE
)
```

```
gobject giotto object
sdimx_stepsize stepsize along the x-axis
sdimy_stepsize stepsize along the y-axis
sdimz_stepsize stepsize along the z-axis
```

createSpatialGrid\_2D 69

```
minimum_padding
minimum padding on the edges

name
name for spatial grid (default = 'spatial_grid')

return_gobject boolean: return giotto object (default = TRUE)
```

#### **Details**

Creates a spatial grid with defined x, y (and z) dimensions. The dimension units are based on the provided spatial location units.

### Value

giotto object with updated spatial grid slot

### **Examples**

```
createSpatialGrid(gobject)
```

```
createSpatialGrid_2D createSpatialGrid_2D
```

#### **Description**

create a spatial grid for 2D spatial data.

## Usage

```
createSpatialGrid_2D(
  gobject,
  sdimx_stepsize = NULL,
  sdimy_stepsize = NULL,
  minimum_padding = 1,
  name = "spatial_grid",
  return_gobject = TRUE
)
```

### **Arguments**

### **Details**

Creates a spatial grid with defined x, y (and z) dimensions. The dimension units are based on the provided spatial location units.

#### Value

giotto object with updated spatial grid slot

#### **Examples**

```
createSpatialGrid_2D(gobject)
```

```
createSpatialGrid_3D createSpatialGrid_3D
```

#### **Description**

Create a spatial grid for 3D spatial data.

### Usage

```
createSpatialGrid_3D(
  gobject,
  sdimx_stepsize = NULL,
  sdimy_stepsize = NULL,
  sdimz_stepsize = NULL,
  minimum_padding = 1,
  name = "spatial_grid",
  return_gobject = TRUE
)
```

# Arguments

## **Details**

Creates a spatial grid with defined x, y (and z) dimensions. The dimension units are based on the provided spatial location units.

#### Value

giotto object with updated spatial grid slot

# Examples

```
createSpatialGrid_3D(gobject)
```

```
createSpatialKNNnetwork
```

*createSpatialKNNnetwork* 

### **Description**

Create a spatial knn network.

#### Usage

```
createSpatialKNNnetwork(
  gobject,
  method = "dbscan",
  dimensions = "all",
  name = "knn_network",
  k = 4,
  maximum_distance = NULL,
  minimum_k = 0,
  verbose = F,
  return_gobject = TRUE,
  ...
)
```

#### **Arguments**

```
gobject giotto object
```

method method to create kNN network

dimensions which spatial dimensions to use (default = all)

name name for spatial network (default = 'spatial\_network')k number of nearest neighbors based on physical distance

maximum\_distance

distance cuttof for nearest neighbors to consider for kNN network

minimum\_k minimum nearest neigbhours if maximum\_distance != NULL

verbose verbose

return\_gobject | boolean: return giotto object (default = TRUE)

### Value

giotto object with updated spatial network slot

**dimensions:** default = 'all' which takes all possible dimensions. Alternatively you can provide a character vector that specififies the spatial dimensions to use, e.g. c("sdimx', "sdimy") or a numerical vector, e.g. 2:3

 $maximum\_distance$ : to create a network based on maximum distance only, you also need to set k to a very high value, e.g. k = 100

#### **Examples**

```
createSpatialKNNnetwork(gobject)
```

72 createSpatialNetwork

```
create Spatial Network \\ create Spatial Network
```

# Description

Create a spatial network based on cell centroid physical distances.

# Usage

```
createSpatialNetwork(
  gobject,
  name = NULL,
  dimensions = "all",
  method = c("Delaunay", "kNN"),
delaunay_method = c("deldir", "delaunayn_geometry", "RTriangle"),
maximum_distance_delaunay = "auto",
  options = "Pp",
  Y = TRUE,
  j = TRUE,
  S = 0,
  minimum_k = 0,
  knn_method = "dbscan",
  k = 4,
  maximum_distance_knn = NULL,
  verbose = F,
  return_gobject = TRUE,
)
```

gobject	giotto object	
name	<pre>name for spatial network (default = 'spatial_network')</pre>	
dimensions	which spatial dimensions to use (default = all)	
method	which method to use to create a spatial network. (default = Delaunay)	
delaunay_method		
	Delaunay method to use	
maximum_distance_delaunay		
	distance cuttof for nearest neighbors to consider for Delaunay network	
options	(geometry) String containing extra control options for the underlying Qhull command; see the Qhull documentation (/doc/qhull/html/qdelaun.html) for the available options. (default = 'Pp', do not report precision problems)	
Υ	(RTriangle) If TRUE prohibits the insertion of Steiner points on the mesh boundary.	
j	(RTriangle) If TRUE jettisons vertices that are not part of the final triangulation from the output.	
S	(RTriangle) Specifies the maximum number of added Steiner points.	
minimum_k	minimum nearest neigbhours if maximum_distance != NULL	

```
knn_method method to create kNN network
k number of nearest neighbors based on physical distance
maximum_distance_knn
distance cuttof for nearest neighbors to consider for kNN network
verbose verbose
return_gobject boolean: return giotto object (default = TRUE)
```

#### **Details**

Creates a spatial network connecting single-cells based on their physical distance to each other. For Delaunay method, neighbors will be decided by delaunay triangulation and a maximum distance criteria. For kNN method, number of neighbors can be determined by k, or maximum distance from each cell with or without setting a minimum k for each cell.

**dimensions:** default = 'all' which takes all possible dimensions. Alternatively you can provide a character vector that specififies the spatial dimensions to use, e.g. c("sdimx', "sdimy") or a numerical vector, e.g. 2:3

#### Value

giotto object with updated spatial network slot

#### **Examples**

```
createSpatialNetwork(gobject)
```

#### **Description**

calculates average gene detection for a cell metadata factor (e.g. cluster)

### Usage

```
create_average_detection_DT(
  gobject,
  meta_data_name,
  expression_values = c("normalized", "scaled", "custom"),
  detection_threshold = 0
)
```

# Arguments

#### Value

data.table with average gene epression values for each factor

### Description

calculates average gene expression for a cell metadata factor (e.g. cluster)

### Usage

```
create_average_DT(
  gobject,
  meta_data_name,
  expression_values = c("normalized", "scaled", "custom")
)
```

### **Arguments**

```
gobject giotto object

meta_data_name name of metadata column to use
expression_values

which expression values to use
```

#### Value

data.table with average gene epression values for each factor

### **Description**

creates randomized cell ids within a selection of cell types

### Usage

```
create_cell_type_random_cell_IDs(
  gobject,
  cluster_column = "cell_types",
  needed_cell_types
)
```

#### **Arguments**

```
gobject giotto object to use

cluster_column cluster column with cell type information

needed_cell_types

vector of cell type names for which a random id will be found
```

#### Details

Details will follow.

#### Value

list of randomly sampled cell ids with same cell type composition

#### **Examples**

```
create_cell_type_random_cell_IDs(gobject)
```

### Description

create a crossSection object

#### Usage

```
create_crossSection_object(
  name = NULL,
  method = NULL,
  thickness_unit = NULL,
  slice_thickness = NULL,
  plane_equation = NULL,
  mesh_grid_n = NULL,
  mesh_obj = NULL,
  cell_subset = NULL,
  cell_subset_spatial_locations = NULL,
  cell_subset_projection_locations = NULL,
  cell_subset_projection_PCA = NULL,
  cell_subset_projection_coords = NULL)
)
```

### **Arguments**

name name of cress section object. (default = cross\_sectino)
method method to define the cross section plane.

thickness\_unit unit of the virtual section thickness. If "cell", average size of the observed cells is used as length unit. If "natural", the unit of cell location coordinates

is used.(default = cell)

```
slice_thickness
                  thickness of slice
plane_equation a numerical vector of length 4, in the form of c(A,B,C,D), which defines plane
                  Ax+By+Cz=D.
mesh_grid_n
                  numer of meshgrid lines to generate along both directions for the cross section
mesh_obj
                  object that stores the cross section meshgrid information.
cell_subset
                  cells selected by the cross section
cell_subset_spatial_locations
                  locations of cells selected by the cross section
cell\_subset\_projection\_locations
                  3D projection coordinates of selected cells onto the cross section plane
cell_subset_projection_PCA
                  pca of projection coordinates
cell_subset_projection_coords
                  2D PCA coordinates of selected cells in the cross section plane
cell_distance_estimate_method
                  method to estimate average distance between neighboring cells. (default = mean)
                  deciding the span of the cross section meshgrid, as a ratio of extension compared
extend_ratio
                  to the borders of the vitural tissue section. (default = 0.2)
```

create\_delaunayNetwork2D

create\_delaunayNetwork2D

### **Description**

Create a spatial Delaunay network.

#### Usage

```
create_delaunayNetwork2D(
  gobject,
  method = c("delaunayn_geometry", "RTriangle", "deldir"),
  sdimx = "sdimx",
  sdimy = "sdimy",
  name = "delaunay_network",
  maximum_distance = "auto",
  minimum_k = 0,
  options = "Pp",
  Y = TRUE,
  j = TRUE,
  S = 0,
  verbose = T,
  return_gobject = TRUE,
  ...
)
```

#### **Examples**

create\_delaunayNetwork2D(gobject)

```
create_delaunayNetwork3D
```

create\_delaunayNetwork3D

### Description

Create a spatial Delaunay network.

### Usage

```
create_delaunayNetwork3D(
  gobject,
  method = "delaunayn_geometry",
  sdimx = "sdimx",
  sdimy = "sdimy",
  sdimz = "sdimz",
  name = "delaunay_network_3D",
  maximum_distance = "auto",
  minimum_k = 0,
  options = "Pp",
  return_gobject = TRUE,
  ...
)
```

### **Examples**

 ${\tt create\_delaunayNetwork3D(gobject)}$ 

```
create\_delaunayNetwork\_deldir \\ create\_delaunayNetwork\_deldir
```

# Description

Create a spatial Delaunay network.

### Usage

```
create_delaunayNetwork_deldir(
   spatial_locations,
   sdimx = "sdimx",
   sdimy = "sdimy",
   ...
)
```

```
create_delaunayNetwork_deldir(gobject)
```

```
create\_delaunayNetwork\_geometry \\ create\_delaunayNetwork\_geometry
```

# Description

Create a spatial Delaunay network.

### Usage

```
create_delaunayNetwork_geometry(
  spatial_locations,
  sdimx = "sdimx",
  sdimy = "sdimy",
  options = "Pp",
   ...
)
```

### **Examples**

```
create_delaunayNetwork_geometry(gobject)
```

### Description

Create a spatial Delaunay network.

### Usage

```
create_delaunayNetwork_geometry_3D(
  spatial_locations,
  sdimx = "sdimx",
  sdimy = "sdimy",
  sdimz = "sdimz",
  options = options,
  ...
)
```

```
{\tt create\_delaunayNetwork\_geometry\_3D(gobject)}
```

```
create\_delaunay Network\_RTriangle \\ create\_delaunay Network\_RTriangle
```

### Description

Create a spatial Delaunay network.

# Usage

```
create_delaunayNetwork_RTriangle(
   spatial_locations,
   sdimx = "sdimx",
   sdimy = "sdimy",
   Y = TRUE,
   j = TRUE,
   S = 0,
   ...
)
```

### **Examples**

create\_delaunayNetwork\_RTriangle(gobject)

### Description

subsets matrix based on vector of genes or hvg column

### Usage

```
create_genes_to_use_matrix(gobject, sel_matrix, genes_to_use, verbose = TRUE)
```

### Arguments

```
gobject giotto object
sel_matrix selected expression matrix
genes_to_use genes to use, character or vector of genes
verbose verbosity
```

#### Value

subsetted matrix based on selected genes

80 create\_screeplot

```
create\_KNNnetwork\_dbscan \\ create\_KNNnetwork\_dbscan
```

# Description

Create a spatial knn network.

### Usage

```
create_KNNnetwork_dbscan(
  spatial_locations,
  sdimx = "sdimx",
  sdimy = "sdimy",
  sdimz = "sdimz",
  k = 4,
  ...
)
```

### **Examples**

```
create_KNNnetwork_dbscan(gobject)
```

create\_screeplot

create\_screeplot

### **Description**

create screeplot with ggplot

# Usage

```
create_screeplot(pca_obj, ncp = 20, ylim = c(0, 20))
```

# Arguments

pca\_obj pca dimension reduction object

ncp number of principal components to calculate

ylim y-axis limits on scree plot

#### Value

ggplot

crossSectionGenePlot 81

```
{\tt crossSectionGenePlot} \quad {\it crossSectionGenePlot}
```

### Description

Visualize cells and gene expression in a virtual cross section according to spatial coordinates

### Usage

```
crossSectionGenePlot(
  gobject = NULL,
  crossSection_obj = NULL,
  name = NULL,
  spatial_network_name = "Delaunay_network",
  default_save_name = "crossSectionGenePlot",
  ...
)
```

### **Arguments**

```
gobject giotto object

crossSection_obj

crossSection object

name name of virtual cross section to use

spatial_network_name

name of spatial network to use

default_save_name

default save name for saving, don't change, change save_name in save_param

parameters for spatGenePlot2D
```

### **Details**

Description of parameters.

#### Value

ggplot

#### See Also

```
spatGenePlot3D and spatGenePlot2D
```

82 crossSectionGenePlot3D

```
crossSectionGenePlot3D
```

crossSectionGenePlot3D

### Description

Visualize cells and gene expression in a virtual cross section according to spatial coordinates

### Usage

```
crossSectionGenePlot3D(
  gobject,
  crossSection_obj = NULL,
  name = NULL,
  spatial_network_name = "Delaunay_network",
  other_cell_color = alpha("lightgrey", 0),
  default_save_name = "crossSectionGenePlot3D",
  ...
)
```

### Arguments

```
gobject giotto object

name name of virtual cross section to use

spatial_network_name
    name of spatial network to use

default_save_name
    default save name for saving, don't change, change save_name in save_param

... parameters for spatGenePlot3D
```

#### **Details**

Description of parameters.

# Value

ggplot

```
crossSectionGenePlot3D(gobject)
```

crossSectionPlot 83

# Description

Visualize cells in a virtual cross section according to spatial coordinates

### Usage

```
crossSectionPlot(
  gobject,
  crossSection_obj = NULL,
  name = NULL,
  spatial_network_name = "Delaunay_network",
  default_save_name = "crossSectionPlot",
   ...
)
```

### Arguments

```
gobject giotto object

name name of virtual cross section to use

spatial_network_name
 name of spatial network to use

default_save_name
 default save name for saving, don't change, change save_name in save_param

... parameters for spatPlot2D
```

# **Details**

Description of parameters.

### Value

ggplot

#### See Also

crossSectionPlot

84 crossSectionPlot3D

crossSectionPlot3D

### Description

Visualize cells in a virtual cross section according to spatial coordinates

### Usage

```
crossSectionPlot3D(
  gobject,
  crossSection_obj = NULL,
  name = NULL,
  spatial_network_name = "Delaunay_network",
  show_other_cells = T,
  other_cell_color = alpha("lightgrey", 0),
  default_save_name = "crossSection3D",
  ...
)
```

### Arguments

### Details

Description of parameters.

### Value

ggplot

```
{\tt crossSectionPlot3D(gobject)}
```

decide\_cluster\_order 85

```
decide_cluster_order
decide_cluster_order
```

### Description

creates order for clusters

### Usage

```
decide_cluster_order(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  genes,
  cluster_column = NULL,
  cluster_order = c("size", "correlation", "custom"),
  cluster_custom_order = NULL,
  cor_method = "pearson",
  hclust_method = "ward.D"
)
```

### **Arguments**

```
gobject giotto object
expression_values
expression values to use
genes genes to use
cluster_column name of column to use for clusters
cluster_order method to determine cluster order
cluster_custom_order
custom order for clusters

cor_method method for correlation
hclust_method method for hierarchical clustering
```

### **Details**

Calculates order for clusters.

#### Value

custom

```
decide_cluster_order(gobject)
```

detectSpatialCorGenes detectSpatialCorGenes

### **Description**

Detect genes that are spatially correlated

### Usage

```
detectSpatialCorGenes(
  gobject,
  method = c("grid", "network"),
  expression_values = c("normalized", "scaled", "custom"),
  subset_genes = NULL,
  spatial_network_name = "Delaunay_network",
  network_smoothing = NULL,
  spatial_grid_name = "spatial_grid",
  min_cells_per_grid = 4,
  cor_method = c("pearson", "kendall", "spearman")
)
```

### **Arguments**

```
gobject
                  giotto object
method
                  method to use for spatial averaging
expression_values
                  gene expression values to use
subset_genes
                  subset of genes to use
spatial_network_name
                  name of spatial network to use
network_smoothing
                  smoothing factor beteen 0 and 1 (default: automatic)
spatial_grid_name
                  name of spatial grid to use
min_cells_per_grid
                  minimum number of cells to consider a grid
cor_method
                  correlation method
```

### **Details**

For method = network, it expects a fully connected spatial network. You can make sure to create a fully connected network by setting minimal\_k > 0 in the createSpatialNetwork function.

- 1. grid-averaging: average gene expression values within a predefined spatial grid
- 2. network-averaging: smoothens the gene expression matrix by averaging the expression within one cell by using the neighbours within the predefined spatial network. b is a smoothening factor that defaults to 1 1/k, where k is the median number of k-neighbors in the selected spatial network. Setting b = 0 means no smoothing and b = 1 means no contribution from its own expression.

The spatCorObject can be further explored with showSpatialCorGenes()

detectSpatialPatterns 87

#### Value

```
returns a spatial correlation object: "spatCorObject"
```

#### See Also

```
showSpatialCorGenes
```

### **Examples**

```
detectSpatialCorGenes(gobject)
```

```
detectSpatialPatterns detectSpatialPatterns
```

### **Description**

Identify spatial patterns through PCA on average expression in a spatial grid.

# Usage

```
detectSpatialPatterns(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  spatial_grid_name = "spatial_grid",
  min_cells_per_grid = 4,
  scale_unit = F,
  ncp = 100,
  show_plot = T,
  PC_zscore = 1.5
)
```

### Arguments

```
gobject
                  giotto object
expression_values
                  expression values to use
spatial_grid_name
                  name of spatial grid to use (default = 'spatial_grid')
min_cells_per_grid
                  minimum number of cells in a grid to be considered
                  scale features
scale_unit
                  number of principal components to calculate
ncp
show_plot
                  show plots
PC_zscore
                  minimum z-score of variance explained by a PC
```

88 dimCellPlot

#### **Details**

Steps to identify spatial patterns:

- 1. average gene expression for cells within a grid, see createSpatialGrid
- 2. perform PCA on the average grid expression profiles
- 3. convert variance of principlal components (PCs) to z-scores and select PCs based on a z-score threshold

#### Value

```
spatial pattern object 'spatPatObj'
```

#### **Examples**

```
detectSpatialPatterns(gobject)
```

dimCellPlot

dimCellPlot

### **Description**

Visualize cells according to dimension reduction coordinates

### Usage

```
dimCellPlot(...)
```

# Arguments

```
Arguments passed on to dimCellPlot2D
gobject giotto object
dim_reduction_to_use dimension reduction to use
dim_reduction_name dimension reduction name
dim1_to_use dimension to use on x-axis
dim2_to_use dimension to use on y-axis
spat_enr_names names of spatial enrichment results to include
cell_annotation_values numeric cell annotation columns
show_NN_network show underlying NN network
nn_network_to_use type of NN network to use (kNN vs sNN)
network_name name of NN network to use, if show_NN_network = TRUE
cell_color_gradient vector with 3 colors for numeric data
gradient_midpoint midpoint for color gradient
gradient_limits vector with lower and upper limits
select_cell_groups select subset of cells/clusters based on cell_color param-
select_cells select subset of cells based on cell IDs
show_other_cells display not selected cells
other_cell_color color of not selected cells
```

dimCellPlot 89

```
other_point_size size of not selected cells
show_cluster_center plot center of selected clusters
show_center_label plot label of selected clusters
center_point_size size of center points
label_size size of labels
label_fontface font of labels
edge_alpha column to use for alpha of the edges
point_shape point with border or not (border or no_border)
point_size size of point (cell)
point_alpha transparancy of dim. reduction points
point_border_col color of border around points
point_border_stroke stroke size of border around points
show_legend show legend
legend_text size of legend text
legend_symbol_size size of legend symbols
background_color color of plot background
axis_text size of axis text
axis_title size of axis title
show_plot show plot
return_plot return ggplot object
save_plot directly save the plot [boolean]
save_param list of saving parameters, see showSaveParameters
default_save_name default save name for saving, don't change, change save_name
    in save_param
```

### Details

Description of parameters. For 3D plots see dimCellPlot2D

#### Value

ggplot

#### See Also

Other dimension reduction cell annotation visualizations: dimCellPlot2D()

```
dimCellPlot(gobject)
```

90 dimCellPlot2D

dimCellPlot2D

dimCellPlot2D

#### **Description**

Visualize cells according to dimension reduction coordinates

### Usage

```
dimCellPlot2D(
  gobject,
  dim_reduction_to_use = "umap",
  dim_reduction_name = "umap",
  dim1_to_use = 1,
  dim2\_to\_use = 2,
  spat_enr_names = NULL,
  cell_annotation_values = NULL,
  show_NN_network = F,
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  cell_color_gradient = c("blue", "white", "red"),
  gradient_midpoint = NULL,
  gradient_limits = NULL,
  select_cell_groups = NULL,
  select_cells = NULL,
  show_other_cells = T,
  other_cell_color = "lightgrey",
  other_point_size = 0.5,
  show_cluster_center = F,
  show\_center\_label = T,
  center_point_size = 4,
  center_point_border_col = "black",
  center_point_border_stroke = 0.1,
  label_size = 4,
  label_fontface = "bold",
  edge_alpha = NULL,
  point_shape = c("border", "no_border"),
  point_size = 1,
  point_alpha = 1,
  point_border_col = "black",
  point_border_stroke = 0.1,
  show_legend = T,
  legend_text = 8,
  legend_symbol_size = 1,
  background_color = "white",
  axis_text = 8,
  axis_title = 8,
  cow_n_col = 2,
  cow_rel_h = 1,
  cow_rel_w = 1,
  cow_align = "h",
```

dimCellPlot2D 91

```
show_plot = NA,
      return_plot = NA,
      save_plot = NA,
      save_param = list(),
      default_save_name = "dimCellPlot2D"
Arguments
                     giotto object
    gobject
    dim_reduction_to_use
                      dimension reduction to use
    dim_reduction_name
                      dimension reduction name
                      dimension to use on x-axis
    dim1_to_use
                      dimension to use on y-axis
    dim2_to_use
    spat_enr_names names of spatial enrichment results to include
    cell_annotation_values
                      numeric cell annotation columns
    show_NN_network
                      show underlying NN network
    nn_network_to_use
                      type of NN network to use (kNN vs sNN)
    network_name
                      name of NN network to use, if show_NN_network = TRUE
    cell_color_gradient
                      vector with 3 colors for numeric data
    gradient_midpoint
                      midpoint for color gradient
    gradient_limits
                      vector with lower and upper limits
    select_cell_groups
                      select subset of cells/clusters based on cell_color parameter
                      select subset of cells based on cell IDs
    select_cells
    show\_other\_cells
                      display not selected cells
    other_cell_color
                      color of not selected cells
    other_point_size
                      size of not selected cells
    show_cluster_center
                      plot center of selected clusters
    show_center_label
                      plot label of selected clusters
    center_point_size
                      size of center points
    label_size
                      size of labels
    label_fontface font of labels
    edge_alpha
                      column to use for alpha of the edges
```

92 dimCellPlot2D

```
point with border or not (border or no_border)
point_shape
point_size
                  size of point (cell)
point_alpha
                  transparancy of dim. reduction points
point_border_col
                  color of border around points
point_border_stroke
                  stroke size of border around points
show_legend
                  show legend
legend_text
                  size of legend text
legend_symbol_size
                  size of legend symbols
background_color
                  color of plot background
axis_text
                  size of axis text
axis_title
                  size of axis title
show_plot
                  show plot
return_plot
                  return ggplot object
save_plot
                  directly save the plot [boolean]
                  list of saving parameters, see showSaveParameters
save_param
default_save_name
                  default save name for saving, don't change, change save_name in save_param
cell_color
                  color for cells (see details)
color_as_factor
                  convert color column to factor
cell_color_code
                  named vector with colors
                  title for plot, defaults to cell_color parameter
title
```

#### **Details**

Description of parameters. For 3D plots see dimPlot3D

### Value

ggplot

# See Also

Other dimension reduction cell annotation visualizations: dimCellPlot()

```
dimCellPlot2D(gobject)
```

dimGenePlot 93

dimGenePlot

dimGenePlot

in save\_param

#### **Description**

Visualize gene expression according to dimension reduction coordinates

#### Usage

```
dimGenePlot(...)
```

#### **Arguments**

```
Arguments passed on to dimGenePlot2D
gobject giotto object
expression_values gene expression values to use
genes genes to show
dim_reduction_to_use dimension reduction to use
dim_reduction_name dimension reduction name
dim1_to_use dimension to use on x-axis
dim2_to_use dimension to use on y-axis
show_NN_network show underlying NN network
nn_network_to_use type of NN network to use (kNN vs sNN)
network_name name of NN network to use, if show NN network = TRUE
edge_alpha column to use for alpha of the edges
scale_alpha_with_expression scale expression with ggplot alpha parameter
point_shape point with border or not (border or no_border)
point_size size of point (cell)
point_alpha transparancy of points
cell_color_gradient vector with 3 colors for numeric data
gradient_midpoint midpoint for color gradient
gradient_limits vector with lower and upper limits
point_border_col color of border around points
point_border_stroke stroke size of border around points
show_legend show legend
legend_text size of legend text
background_color color of plot background
axis_text size of axis text
axis_title size of axis title
cow_n_col cowplot param: how many columns
cow_rel_h cowplot param: relative height
cow_rel_w cowplot param: relative width
cow_align cowplot param: how to align
show_plot show plots
return_plot return ggplot object
save_plot directly save the plot [boolean]
save_param list of saving parameters, see showSaveParameters
default_save_name default save name for saving, don't change, change save_name
```

94 dimGenePlot2D

#### **Details**

Description of parameters.

#### Value

ggplot

#### See Also

```
dimGenePlot3D
```

Other dimension reduction gene expression visualizations: dimGenePlot2D(), dimGenePlot3D()

#### **Examples**

```
dimGenePlot(gobject)
```

dimGenePlot2D

dimGenePlot2D

### **Description**

Visualize gene expression according to dimension reduction coordinates

### Usage

```
dimGenePlot2D(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  genes = NULL,
  dim_reduction_to_use = "umap",
  dim_reduction_name = "umap",
  dim1_to_use = 1,
  dim2_to_use = 2,
  show_NN_network = F,
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  network_color = "lightgray",
  edge_alpha = NULL,
  scale_alpha_with_expression = FALSE,
  point_shape = c("border", "no_border"),
  point_size = 1,
  point_alpha = 1,
  cell_color_gradient = c("blue", "white", "red"),
  gradient_midpoint = NULL,
  gradient_limits = NULL,
  point_border_col = "black",
  point_border_stroke = 0.1,
  show_legend = T,
  legend_text = 8,
  background_color = "white",
  axis_text = 8,
```

dimGenePlot2D 95

```
axis_title = 8,
      cow_n_col = 2,
      cow_rel_h = 1,
      cow_rel_w = 1,
      cow_align = "h",
      show_plot = NA,
      return_plot = NA,
      save_plot = NA,
      save_param = list(),
      default_save_name = "dimGenePlot2D"
    )
Arguments
    gobject
                     giotto object
    expression_values
                     gene expression values to use
                     genes to show
    genes
    dim_reduction_to_use
                     dimension reduction to use
    dim_reduction_name
                     dimension reduction name
    dim1_to_use
                     dimension to use on x-axis
    dim2_to_use
                     dimension to use on y-axis
    show_NN_network
                     show underlying NN network
    nn_network_to_use
                     type of NN network to use (kNN vs sNN)
                     name of NN network to use, if show_NN_network = TRUE
    network_name
    edge_alpha
                     column to use for alpha of the edges
    scale_alpha_with_expression
                     scale expression with ggplot alpha parameter
                     point with border or not (border or no_border)
    point_shape
                     size of point (cell)
    point_size
    point_alpha
                     transparancy of points
    cell_color_gradient
                     vector with 3 colors for numeric data
    gradient_midpoint
                     midpoint for color gradient
    gradient_limits
                     vector with lower and upper limits
    point_border_col
                     color of border around points
    point_border_stroke
                     stroke size of border around points
    show_legend
                     show legend
    legend_text
                     size of legend text
```

96 dimGenePlot3D

color of plot background axis\_text size of axis text size of axis title axis\_title cow\_n\_col cowplot param: how many columns cow\_rel\_h cowplot param: relative height cowplot param: relative width cow\_rel\_w cow\_align cowplot param: how to align show\_plot show plots return\_plot return ggplot object

save\_plot directly save the plot [boolean]

save\_param list of saving parameters, see showSaveParameters

default\_save\_name

background\_color

default save name for saving, don't change, change save\_name in save\_param

... additional parameters for cowplot::save\_plot()

#### **Details**

Description of parameters.

# Value

ggplot

#### See Also

```
dimGenePlot3D
```

Other dimension reduction gene expression visualizations: dimGenePlot3D(), dimGenePlot()

# **Examples**

dimGenePlot2D(gobject)

dimGenePlot3D dimGenePlot3D

### Description

Visualize cells and gene expression according to dimension reduction coordinates

dimGenePlot3D 97

#### Usage

```
dimGenePlot3D(
     gobject,
     expression_values = c("normalized", "scaled", "custom"),
     genes = NULL,
     dim_reduction_to_use = "umap",
     dim_reduction_name = "umap",
     dim1_to_use = 1,
     dim2\_to\_use = 2,
     dim3_to_use = 3,
      show_NN_network = F,
     nn_network_to_use = "sNN",
     network_name = "sNN.pca",
     network_color = "lightgray",
     cluster_column = NULL,
      select_cell_groups = NULL,
      select_cells = NULL,
      show_other_cells = T,
     other_cell_color = "lightgrey",
     other_point_size = 1,
     edge_alpha = NULL,
     point_size = 2,
      genes_high_color = NULL,
     genes_mid_color = "white",
     genes_low_color = "blue",
      show_legend = T,
     show_plot = NA,
     return_plot = NA,
      save_plot = NA,
      save_param = list(),
     default_save_name = "dimGenePlot3D"
   )
Arguments
   gobject
                    giotto object
   expression_values
                    gene expression values to use
                    genes to show
   genes
   dim_reduction_to_use
                    dimension reduction to use
   dim_reduction_name
                    dimension reduction name
   dim1_to_use
                    dimension to use on x-axis
   dim2_to_use
                    dimension to use on y-axis
   dim3_to_use
                    dimension to use on z-axis
   show_NN_network
                    show underlying NN network
   nn_network_to_use
```

type of NN network to use (kNN vs sNN)

98 dimPlot

```
name of NN network to use, if show_NN_network = TRUE
network_name
                  column to use for alpha of the edges
edge_alpha
                  size of point (cell)
point_size
show_legend
                  show legend
show_plot
                  show plots
return_plot
                  return ggplot object
save_plot
                  directly save the plot [boolean]
                  list of saving parameters, see showSaveParameters
save_param
default_save_name
                  default save name for saving, don't change, change save_name in save_param
                  additional parameters for cowplot::save_plot()
```

#### **Details**

Description of parameters.

#### Value

ggplot

#### See Also

Other dimension reduction gene expression visualizations: dimGenePlot2D(), dimGenePlot()

#### **Examples**

```
dimGenePlot3D(gobject)
```

dimPlot

dimPlot

### **Description**

Visualize cells according to dimension reduction coordinates

#### Usage

```
dimPlot(...)
```

### **Arguments**

arguments passed on to dimPlot2D
gobject giotto object
group\_by\_subset subset the group\_by factor column
dim\_reduction\_to\_use dimension reduction to use
dim\_reduction\_name dimension reduction name
dim1\_to\_use dimension to use on x-axis
dim2\_to\_use dimension to use on y-axis

dimPlot 99

```
spat_enr_names names of spatial enrichment results to include
show_NN_network show underlying NN network
nn_network_to_use type of NN network to use (kNN vs sNN)
network_name name of NN network to use, if show_NN_network = TRUE
cell_color color for cells (see details)
color_as_factor convert color column to factor
cell_color_code named vector with colors
cell_color_gradient vector with 3 colors for numeric data
gradient_midpoint midpoint for color gradient
gradient_limits vector with lower and upper limits
select_cell_groups select subset of cells/clusters based on cell_color param-
select_cells select subset of cells based on cell IDs
show_other_cells display not selected cells
other_cell_color color of not selected cells
other_point_size size of not selected cells
show_cluster_center plot center of selected clusters
show_center_label plot label of selected clusters
center_point_size size of center points
label_size size of labels
label_fontface font of labels
edge_alpha column to use for alpha of the edges
point_shape point with border or not (border or no_border)
point_size size of point (cell)
point_alpha transparancy of point
point_border_col color of border around points
point_border_stroke stroke size of border around points
title title for plot, defaults to cell_color parameter
show_legend show legend
legend_text size of legend text
legend_symbol_size size of legend symbols
background_color color of plot background
axis_text size of axis text
axis_title size of axis title
cow_n_col cowplot param: how many columns
cow_rel_h cowplot param: relative height
cow_rel_w cowplot param: relative width
cow_align cowplot param: how to align
show_plot show plot
return_plot return ggplot object
save_plot directly save the plot [boolean]
save_param list of saving parameters, see showSaveParameters
default_save_name default save name for saving, don't change, change save_name
    in save_param
```

#### **Details**

Description of parameters, see dimPlot2D. For 3D plots see dimPlot3D

100 dimPlot2D

#### Value

ggplot

#### See Also

```
Other reduced dimension visualizations: dimPlot2D(), dimPlot3D(), plotPCA_2D(), plotPCA_3D(), plotPCA(), plotTSNE_2D(), plotTSNE_3D(), plotTSNE(), plotUMAP_2D(), plotUMAP_3D(), plotUMAP()
```

### **Examples**

```
dimPlot(gobject)
```

dimPlot2D

dimPlot2D

### Description

Visualize cells according to dimension reduction coordinates

### Usage

```
dimPlot2D(
  gobject,
  group_by = NULL,
  group_by_subset = NULL,
  dim_reduction_to_use = "umap",
  dim_reduction_name = "umap",
  dim1_to_use = 1,
  dim2_to_use = 2,
  spat_enr_names = NULL,
  show_NN_network = F,
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  cell_color = NULL,
  color_as_factor = T,
  cell_color_code = NULL,
  cell_color_gradient = c("blue", "white", "red"),
  gradient_midpoint = NULL,
  gradient_limits = NULL,
  select_cell_groups = NULL,
  select_cells = NULL,
  show_other_cells = T,
  other_cell_color = "lightgrey",
  other_point_size = 0.5,
  show_cluster_center = F,
  show_center_label = T,
  center_point_size = 4,
  center_point_border_col = "black",
  center_point_border_stroke = 0.1,
  label_size = 4,
  label_fontface = "bold",
```

dimPlot2D 101

```
edge_alpha = NULL,
      point_shape = c("border", "no_border"),
      point_size = 1,
      point_alpha = 1,
      point_border_col = "black",
      point_border_stroke = 0.1,
      title = NULL,
      show_legend = T,
      legend_text = 8,
      legend_symbol_size = 1,
      background_color = "white",
      axis_text = 8,
      axis_title = 8,
      cow_n_col = 2,
      cow_rel_h = 1,
      cow_rel_w = 1,
      cow_align = "h",
      show_plot = NA,
      return_plot = NA,
      save_plot = NA,
      save_param = list(),
      default_save_name = "dimPlot2D"
    )
Arguments
    gobject
                    giotto object
    group_by_subset
                    subset the group_by factor column
    dim_reduction_to_use
                    dimension reduction to use
   dim_reduction_name
                    dimension reduction name
   dim1_to_use
                    dimension to use on x-axis
    dim2_to_use
                    dimension to use on y-axis
    spat_enr_names names of spatial enrichment results to include
    show_NN_network
                    show underlying NN network
   nn_network_to_use
                    type of NN network to use (kNN vs sNN)
   network_name
                    name of NN network to use, if show_NN_network = TRUE
    cell_color
                    color for cells (see details)
    color_as_factor
                    convert color column to factor
    cell_color_code
                    named vector with colors
    cell_color_gradient
                     vector with 3 colors for numeric data
    gradient_midpoint
                    midpoint for color gradient
```

102 dimPlot2D

```
gradient_limits
                  vector with lower and upper limits
select_cell_groups
                  select subset of cells/clusters based on cell_color parameter
select_cells
                  select subset of cells based on cell IDs
show_other_cells
                  display not selected cells
other_cell_color
                  color of not selected cells
other_point_size
                  size of not selected cells
show_cluster_center
                  plot center of selected clusters
show_center_label
                  plot label of selected clusters
center_point_size
                  size of center points
label_size
                  size of labels
label_fontface font of labels
edge_alpha
                  column to use for alpha of the edges
                  point with border or not (border or no_border)
point_shape
point_size
                  size of point (cell)
point_alpha
                  transparancy of point
point_border_col
                  color of border around points
point_border_stroke
                  stroke size of border around points
title
                  title for plot, defaults to cell_color parameter
show_legend
                  show legend
legend_text
                  size of legend text
legend_symbol_size
                  size of legend symbols
background_color
                  color of plot background
axis_text
                  size of axis text
axis_title
                  size of axis title
                  cowplot param: how many columns
cow_n_col
                  cowplot param: relative height
cow_rel_h
cow_rel_w
                  cowplot param: relative width
                  cowplot param: how to align
cow_align
show_plot
                  show plot
return_plot
                  return ggplot object
save_plot
                  directly save the plot [boolean]
save_param
                  list of saving parameters, see showSaveParameters
default_save_name
                  default save name for saving, don't change, change save_name in save_param
                  create multiple plots based on cell annotation column
groub_by
```

dimPlot3D 103

#### **Details**

Description of parameters. For 3D plots see dimPlot3D

#### Value

ggplot

#### See Also

```
Other reduced dimension visualizations: dimPlot3D(), dimPlot(), plotPCA_2D(), plotPCA_3D(), plotPCA(), plotTSNE_2D(), plotTSNE_3D(), plotUMAP_2D(), plotUMAP_3D(), plotUMAP()
```

#### **Examples**

```
dimPlot2D(gobject)
```

dimPlot3D

dimPlot3D

### **Description**

Visualize cells according to dimension reduction coordinates

### Usage

```
dimPlot3D(
  gobject,
  dim_reduction_to_use = "umap",
  dim_reduction_name = "umap",
  dim1_to_use = 1,
  dim2\_to\_use = 2,
  dim3_to_use = 3,
  select_cell_groups = NULL,
  select_cells = NULL,
  show_other_cells = T,
  other_cell_color = "lightgrey",
  other_point_size = 2,
  show_NN_network = F,
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  color_as_factor = T,
  cell_color = NULL,
  cell_color_code = NULL,
  show_cluster_center = F,
  show_center_label = T,
  center_point_size = 4,
  label_size = 4,
  edge_alpha = NULL,
  point_size = 3,
  show_plot = NA,
  return_plot = NA,
```

104 dimPlot3D

 $save_plot = NA$ ,

```
save_param = list(),
      default_save_name = "dim3D"
Arguments
    gobject
                      giotto object
    dim_reduction_to_use
                      dimension reduction to use
    dim_reduction_name
                      dimension reduction name
    dim1_to_use
                      dimension to use on x-axis
    dim2_to_use
                      dimension to use on y-axis
    dim3_to_use
                      dimension to use on z-axis
    select_cell_groups
                      select subset of cells/clusters based on cell_color parameter
                      select subset of cells based on cell IDs
    select_cells
    show_other_cells
                      display not selected cells
    other_cell_color
                      color of not selected cells
    other_point_size
                      size of not selected cells
    show_NN_network
                      show underlying NN network
    nn_network_to_use
                      type of NN network to use (kNN vs sNN)
    network_name
                      name of NN network to use, if show_NN_network = TRUE
    color_as_factor
                      convert color column to factor
    cell_color
                      color for cells (see details)
    cell_color_code
                      named vector with colors
    show_cluster_center
                      plot center of selected clusters
    show_center_label
                      plot label of selected clusters
    center_point_size
                      size of center points
    label_size
                      size of labels
    edge_alpha
                      column to use for alpha of the edges
    point_size
                      size of point (cell)
                      show plot
    show_plot
    return_plot
                      return ggplot object
                      directly save the plot [boolean]
    save_plot
                      list of saving parameters, see showSaveParameters
    save_param
    default_save_name
                      default save name for saving, don't change, change save_name in save_param
    show_legend
                      show legend
```

doHclust 105

#### **Details**

Description of parameters.

#### Value

plotly

#### See Also

```
Other reduced dimension visualizations: dimPlot2D(), dimPlot(), plotPCA_2D(), plotPCA_3D(), plotPCA(), plotTSNE_2D(), plotTSNE_3D(), plotUMAP_2D(), plotUMAP_3D(), plotUMAP()
```

### **Examples**

```
dimPlot3D(gobject)
```

doHclust

doHclust

#### **Description**

cluster cells using hierarchical clustering algorithm

### Usage

```
doHclust(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  genes_to_use = NULL,
  dim_reduction_to_use = c("cells", "pca", "umap", "tsne"),
  dim_reduction_name = "pca",
  dimensions_to_use = 1:10,
  distance_method = c("pearson", "spearman", "original", "euclidean", "maximum",
  "manhattan", "canberra", "binary", "minkowski"),
agglomeration_method = c("ward.D2", "ward.D", "single", "complete", "average",
    "mcquitty", "median", "centroid"),
  k = 10,
  h = NULL,
  name = "hclust",
  return_gobject = TRUE,
  set\_seed = T,
  seed_number = 1234
)
```

#### **Arguments**

```
gobject giotto object
expression_values
expression values to use
genes_to_use subset of genes to use
```

106 doHMRF

```
dim_reduction_to_use
                 dimension reduction to use
dim_reduction_name
                 dimensions reduction name
dimensions_to_use
                 dimensions to use
distance_method
                 distance method
agglomeration_method
                 agglomeration method for hclust
k
                 number of final clusters
                 cut hierarchical tree at height = h
h
                 name for hierarchical clustering
name
return_gobject boolean: return giotto object (default = TRUE)
                 set seed
set_seed
seed_number
                 number for seed
```

#### **Details**

Description on how to use Kmeans clustering method.

#### Value

giotto object with new clusters appended to cell metadata

### See Also

hclust

### **Examples**

doHclust(gobject)

doHMRF

doHMRF

### Description

Run HMRF

# Usage

```
doHMRF(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  spatial_network_name = "Delaunay_network",
  spatial_genes = NULL,
  spatial_dimensions = c("sdimx", "sdimy", "sdimz"),
  dim_reduction_to_use = NULL,
  dim_reduction_name = "pca",
```

doHMRF107

```
dimensions_to_use = 1:10,
  name = "test",
  k = 10,
  betas = c(0, 2, 50),
  tolerance = 1e-10,
  zscore = c("none", "rowcol", "colrow"),
  numinit = 100,
  python_path = NULL,
  output_folder = NULL,
  overwrite_output = TRUE
)
gobject
                giotto object
```

### **Arguments**

```
expression_values
                 expression values to use
spatial_network_name
                 name of spatial network to use for HMRF
spatial_genes
                 spatial genes to use for HMRF
spatial_dimensions
                 select spatial dimensions to use, default is all possible dimensions
dim_reduction_to_use
                 use another dimension reduction set as input
dim_reduction_name
                 name of dimension reduction set to use
dimensions_to_use
                 number of dimensions to use as input
                 name of HMRF run
name
                 number of HMRF domains
k
betas
                 betas to test for
tolerance
                 tolerance
zscore
                 zscore
numinit
                 number of initializations
                 python path to use
python_path
output_folder
                 output folder to save results
overwrite_output
```

overwrite output folder

#### Details

Description of HMRF parameters ...

### Value

Creates a directory with results that can be viewed with viewHMRFresults

```
doHMRF(gobject)
```

108 doKmeans

doKmeans

doKmeans

### **Description**

cluster cells using kmeans algorithm

### Usage

```
doKmeans(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  genes_to_use = NULL,
  dim_reduction_to_use = c("cells", "pca", "umap", "tsne"),
  dim_reduction_name = "pca",
  dimensions_to_use = 1:10,
  distance_method = c("original", "pearson", "spearman", "euclidean", "maximum",
    "manhattan", "canberra", "binary", "minkowski"),
  centers = 10,
  iter_max = 100,
  nstart = 1000,
  algorithm = "Hartigan-Wong",
  name = "kmeans",
  return_gobject = TRUE,
  set_seed = T,
  seed_number = 1234
)
```

#### **Arguments**

```
gobject
                 giotto object
expression_values
                 expression values to use
                 subset of genes to use
genes_to_use
dim_reduction_to_use
                 dimension reduction to use
dim_reduction_name
                 dimensions reduction name
dimensions_to_use
                 dimensions to use
distance_method
                 distance method
                 number of final clusters
centers
                 kmeans maximum iterations
iter_max
nstart
                 kmeans nstart
                 kmeans algorithm
algorithm
name
                 name for kmeans clustering
return_gobject boolean: return giotto object (default = TRUE)
set_seed
                 set seed
seed_number
                 number for seed
```

doLeidenCluster 109

### **Details**

Description on how to use Kmeans clustering method.

### Value

giotto object with new clusters appended to cell metadata

#### See Also

kmeans

### **Examples**

```
doKmeans(gobject)
```

doLeidenCluster

doLeidenCluster

# **Description**

cluster cells using a NN-network and the Leiden community detection algorithm

# Usage

```
doLeidenCluster(
  gobject,
  name = "leiden_clus",
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  python_path = NULL,
  resolution = 1,
  weight_col = "weight",
  partition_type = c("RBConfigurationVertexPartition", "ModularityVertexPartition"),
  init_membership = NULL,
  n_iterations = 1000,
  return_gobject = TRUE,
  set_seed = T,
  seed_number = 1234
)
```

110 doLeidenSubCluster

```
partition_type The type of partition to use for optimisation.
init_membership
```

initial membership of cells for the partition

n\_iterations number of interations to run the Leiden algorithm. If the number of iterations

is negative, the Leiden algorithm is run until an iteration in which there was no

improvement.

return\_gobject boolean: return giotto object (default = TRUE)

set\_seed set seed

seed\_number number for seed

#### **Details**

This function is a wrapper for the Leiden algorithm implemented in python, which can detect communities in graphs of millions of nodes (cells), as long as they can fit in memory. See the https://github.com/vtraag/leidenalgleidenalg github page or the https://leidenalg.readthedocs.io/en/stable/index.htmlreadthedocs page for more information.

Partition types available and information:

- RBConfigurationVertexPartition: Implements Reichardt and Bornholdt's Potts model with a configuration null model. This quality function is well-defined only for positive edge weights. This quality function uses a linear resolution parameter.
- Modularity Vertex Partition: Implements modularity. This quality function is well-defined only for positive edge weights. It does *not* use the resolution parameter

Set  $weight\_col = NULL$  to give equal weight (=1) to each edge.

#### Value

giotto object with new clusters appended to cell metadata

### **Examples**

```
doLeidenCluster(gobject)
```

doLeidenSubCluster

doLeidenSubCluster

### **Description**

Further subcluster cells using a NN-network and the Leiden algorithm

# Usage

doLeidenSubCluster 111

```
hvg_min_perc_cells = 5,
hvg_mean_expr_det = 1,
use_all_genes_as_hvg = FALSE,
min_nr_of_hvg = 5,
pca_param = list(expression_values = "normalized", scale_unit = T),
nn_param = list(dimensions_to_use = 1:20),
k_neighbors = 10,
resolution = 0.5,
n_iterations = 500,
python_path = NULL,
nn_network_to_use = "sNN",
network_name = "sNN.pca",
return_gobject = TRUE,
verbose = T
)
```

# **Arguments**

```
giotto object
gobject
                  name for new clustering result
name
cluster_column cluster column to subcluster
selected_clusters
                  only do subclustering on these clusters
hvg_param
                  parameters for calculateHVG
hvg_min_perc_cells
                  threshold for detection in min percentage of cells
hvg_mean_expr_det
                  threshold for mean expression level in cells with detection
use_all_genes_as_hvg
                  forces all genes to be HVG and to be used as input for PCA
                  minimum number of HVG, or all genes will be used as input for PCA
min_nr_of_hvg
                  parameters for runPCA
pca_param
                  parameters for parameters for createNearestNetwork
nn_param
k_neighbors
                  number of k for createNearestNetwork
resolution
                  resolution of Leiden clustering
n_iterations
                  number of interations to run the Leiden algorithm.
python_path
                  specify specific path to python if required
nn_network_to_use
                  type of NN network to use (kNN vs sNN)
                  name of NN network to use
network_name
return_gobject boolean: return giotto object (default = TRUE)
verbose
                  verbose
```

# **Details**

This function performs subclustering using the Leiden algorithm on selected clusters. The systematic steps are:

• 1. subset Giotto object

112 doLouvainCluster

- 2. identify highly variable genes
- 3. run PCA
- 4. create nearest neighbouring network
- 5. do Leiden clustering

### Value

giotto object with new subclusters appended to cell metadata

### See Also

doLeidenCluster

### **Examples**

```
doLeidenSubCluster(gobject)
```

doLouvainCluster

doLouvainCluster

# **Description**

cluster cells using a NN-network and the Louvain algorithm.

# Usage

```
doLouvainCluster(
  gobject,
  version = c("community", "multinet"),
  name = "louvain_clus",
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  python_path = NULL,
  resolution = 1,
  weight_col = NULL,
  gamma = 1,
  omega = 1,
  louv_random = F,
  return_gobject = TRUE,
  set_seed = F,
  seed_number = 1234,
)
```

```
gobject giotto object

version implemented version of Louvain clustering to use

name name for cluster

nn_network_to_use

type of NN network to use (kNN vs sNN)
```

network\_name name of NN network to use [community] specify specific path to python if required python\_path resolution [community] resolution weight\_col weight column name [multinet] Resolution parameter for modularity in the generalized louvain method. gamma omega [multinet] Inter-layer weight parameter in the generalized louvain method [community] Will randomize the node evaluation order and the community evallouv\_random uation order to get different partitions at each call return\_gobject boolean: return giotto object (default = TRUE) set\_seed set seed seed\_number number for seed

# **Details**

. . .

Louvain clustering using the community or multinet implementation of the louvain clustering algorithm.

#### Value

giotto object with new clusters appended to cell metadata

additional parameters

### See Also

doLouvainCluster\_community and doLouvainCluster\_multinet

# **Examples**

```
doLouvainCluster(gobject)
```

```
\label{lower} do Louvain Cluster\_community \\ do Louvain Cluster\_community
```

### Description

cluster cells using a NN-network and the Louvain algorithm from the community module in Python

# Usage

```
doLouvainCluster_community(
  gobject,
  name = "louvain_clus",
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  python_path = NULL,
  resolution = 1,
  weight_col = NULL,
  louv_random = F,
```

```
return_gobject = TRUE,
 set_seed = F,
  seed_number = 1234
)
```

# **Arguments**

giotto object gobject name for cluster name nn\_network\_to\_use type of NN network to use (kNN vs sNN) name of NN network to use network\_name specify specific path to python if required python\_path resolution resolution weight\_col weight column to use for edges Will randomize the node evaluation order and the community evaluation order louv\_random to get different partitions at each call return\_gobject boolean: return giotto object (default = TRUE)

set\_seed

seed\_number number for seed

set seed

# **Details**

This function is a wrapper for the Louvain algorithm implemented in Python, which can detect communities in graphs of nodes (cells). See the https://python-louvain.readthedocs.io/ en/latest/index.htmlreadthedocs page for more information.

Set  $weight\_col = NULL$  to give equal weight (=1) to each edge.

### Value

giotto object with new clusters appended to cell metadata

# **Examples**

```
doLouvainCluster_community(gobject)
```

```
doLouvainCluster_multinet
                        doLouvainCluster_multinet
```

# **Description**

cluster cells using a NN-network and the Louvain algorithm from the multinet package in R.

doLouvainSubCluster 115

#### Usage

```
doLouvainCluster_multinet(
  gobject,
  name = "louvain_clus",
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  gamma = 1,
  omega = 1,
  return_gobject = TRUE,
  set_seed = F,
  seed_number = 1234
)
```

### **Arguments**

```
gobject giotto object
name name for cluster
```

nn\_network\_to\_use

type of NN network to use (kNN vs sNN)

network\_name name of NN network to use

gamma Resolution parameter for modularity in the generalized louvain method.

omega Inter-layer weight parameter in the generalized louvain method.

return\_gobject boolean: return giotto object (default = TRUE)

set\_seed set seed

seed\_number number for seed

# **Details**

See glouvain\_ml from the multinet package in R for more information.

# Value

giotto object with new clusters appended to cell metadata

# **Examples**

```
doLouvainCluster_multinet(gobject)
```

 ${\tt doLouvainSubCluster} \qquad {\tt doLouvainSubCluster}$ 

# **Description**

subcluster cells using a NN-network and the Louvain algorithm

116 doLouvainSubCluster

#### **Usage**

```
doLouvainSubCluster(
  gobject,
  name = "sub_louvain_clus",
  version = c("community", "multinet"),
  cluster_column = NULL,
  selected_clusters = NULL,
 hvg_param = list(reverse_log_scale = T, difference_in_variance = 1, expression_values
    = "normalized"),
  hvg_min_perc_cells = 5,
  hvg_mean_expr_det = 1,
  use_all_genes_as_hvg = FALSE,
  min_nr_of_hvg = 5,
  pca_param = list(expression_values = "normalized", scale_unit = T),
  nn_param = list(dimensions_to_use = 1:20),
  k_neighbors = 10,
  resolution = 0.5,
  gamma = 1,
  omega = 1,
  python_path = NULL,
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  return_gobject = TRUE,
  verbose = T
)
```

```
gobject
                  giotto object
name
                  name for new clustering result
version
                  version of Louvain algorithm to use
cluster_column cluster column to subcluster
selected_clusters
                  only do subclustering on these clusters
                  parameters for calculateHVG
hvg_param
hvg_min_perc_cells
                  threshold for detection in min percentage of cells
hvg_mean_expr_det
                  threshold for mean expression level in cells with detection
use_all_genes_as_hvg
                  forces all genes to be HVG and to be used as input for PCA
                  minimum number of HVG, or all genes will be used as input for PCA
min_nr_of_hvg
                  parameters for runPCA
pca_param
nn_param
                  parameters for parameters for createNearestNetwork
k_neighbors
                  number of k for createNearestNetwork
resolution
                  resolution for community algorithm
gamma
                  gamma
omega
                  omega
```

### **Details**

This function performs subclustering using the Louvain algorithm on selected clusters. The systematic steps are:

- 1. subset Giotto object
- 2. identify highly variable genes
- 3. run PCA
- 4. create nearest neighbouring network
- 5. do Louvain clustering

#### Value

giotto object with new subclusters appended to cell metadata

### See Also

doLouvainCluster\_multinet and doLouvainCluster\_community

### **Examples**

```
doLouvainSubCluster(gobject)
```

```
\label{lower} do Louvain SubCluster\_community \\ do Louvain SubCluster\_community
```

# Description

subcluster cells using a NN-network and the Louvain community detection algorithm

#### Usage

```
min_nr_of_hvg = 5,
pca_param = list(expression_values = "normalized", scale_unit = T),
nn_param = list(dimensions_to_use = 1:20),
k_neighbors = 10,
resolution = 0.5,
python_path = NULL,
nn_network_to_use = "sNN",
network_name = "sNN.pca",
return_gobject = TRUE,
verbose = T
```

#### **Arguments**

```
gobject
                  giotto object
name
                  name for new clustering result
cluster_column cluster column to subcluster
selected_clusters
                  only do subclustering on these clusters
                  parameters for calculateHVG
hvg_param
hvg_min_perc_cells
                  threshold for detection in min percentage of cells
hvg_mean_expr_det
                  threshold for mean expression level in cells with detection
use_all_genes_as_hvg
                  forces all genes to be HVG and to be used as input for PCA
min_nr_of_hvg
                 minimum number of HVG, or all genes will be used as input for PCA
                  parameters for runPCA
pca_param
nn_param
                  parameters for parameters for createNearestNetwork
k_neighbors
                  number of k for createNearestNetwork
resolution
                  resolution
                  specify specific path to python if required
python_path
nn_network_to_use
                  type of NN network to use (kNN vs sNN)
network_name
                  name of NN network to use
return_gobject boolean: return giotto object (default = TRUE)
verbose
                  verbose
```

### **Details**

This function performs subclustering using the Louvain community algorithm on selected clusters. The systematic steps are:

- 1. subset Giotto object
- 2. identify highly variable genes
- 3. run PCA
- 4. create nearest neighbouring network
- 5. do Louvain community clustering

#### Value

giotto object with new subclusters appended to cell metadata

#### See Also

```
doLouvainCluster_community
```

### **Examples**

```
doLouvainSubCluster_community(gobject)
```

```
{\tt doLouvainSubCluster\_multinet}
```

doLouvainSubCluster\_multinet

# Description

subcluster cells using a NN-network and the Louvain multinet detection algorithm

# Usage

```
doLouvainSubCluster_multinet(
  gobject,
  name = "sub_louvain_mult_clus",
  cluster_column = NULL,
  selected_clusters = NULL,
 hvg_param = list(reverse_log_scale = T, difference_in_variance = 1, expression_values
    = "normalized"),
  hvg_min_perc_cells = 5,
  hvg_mean_expr_det = 1,
  use_all_genes_as_hvg = FALSE,
  min_nr_of_hvg = 5,
  pca_param = list(expression_values = "normalized", scale_unit = T),
  nn_param = list(dimensions_to_use = 1:20),
  k_neighbors = 10,
  gamma = 1,
  omega = 1,
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  return_gobject = TRUE,
  verbose = T
)
```

```
gobject giotto object

name name for new clustering result

cluster_column cluster column to subcluster

selected_clusters

only do subclustering on these clusters
```

hvg\_param parameters for calculateHVG

hvg\_min\_perc\_cells

threshold for detection in min percentage of cells

hvg\_mean\_expr\_det

threshold for mean expression level in cells with detection

use\_all\_genes\_as\_hvg

forces all genes to be HVG and to be used as input for PCA

min\_nr\_of\_hvg minimum number of HVG, or all genes will be used as input for PCA

pca\_param parameters for runPCA

nn\_param parameters for parameters for createNearestNetwork

k\_neighbors number of k for createNearestNetwork

gamma gamma
omega omega
nn\_network\_to\_use

type of NN network to use (kNN vs sNN)

network\_name name of NN network to use

return\_gobject boolean: return giotto object (default = TRUE)

verbose verbose

#### Details

This function performs subclustering using the Louvain multinet algorithm on selected clusters. The systematic steps are:

- 1. subset Giotto object
- 2. identify highly variable genes
- 3. run PCA
- 4. create nearest neighbouring network
- 5. do Louvain multinet clustering

#### Value

giotto object with new subclusters appended to cell metadata

#### See Also

doLouvainCluster\_multinet

# **Examples**

doLouvainSubCluster\_multinet(gobject)

doRandomWalkCluster 121

doRandomWalkCluster

doRandomWalkCluster

# **Description**

Cluster cells using a random walk approach.

# Usage

```
doRandomWalkCluster(
  gobject,
  name = "random_walk_clus",
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  walk_steps = 4,
  walk_clusters = 10,
  walk_weights = NA,
  return_gobject = TRUE,
  set_seed = F,
  seed_number = 1234
)
```

# **Arguments**

```
gobject
                 giotto object
                 name for cluster
name
nn_network_to_use
                 type of NN network to use (kNN vs sNN)
                 name of NN network to use
network_name
                 number of walking steps
walk_steps
walk_clusters
                 number of final clusters
walk_weights
                 cluster column defining the walk weights
return_gobject | boolean: return giotto object (default = TRUE)
set_seed
                 set seed
seed_number
                 number for seed
```

# **Details**

See cluster\_walktrap function from the igraph package in R for more information.

### Value

giotto object with new clusters appended to cell metadata

```
{\tt doRandomWalkCluster(gobject)}
```

122 doSNNCluster

doSNNCluster

doSNNCluster

# **Description**

Cluster cells using a SNN cluster approach.

### Usage

```
doSNNCluster(
  gobject,
  name = "sNN_clus",
  nn_network_to_use = "kNN",
  network_name = "kNN.pca",
  k = 20,
  eps = 4,
  minPts = 16,
  borderPoints = TRUE,
  return_gobject = TRUE,
  set_seed = F,
  seed_number = 1234
)
```

# **Arguments**

gobject giotto object name name for cluster

 $nn\_network\_to\_use$ 

type of NN network to use (only works on kNN)

 $network\_name \qquad name \ of \ kNN \ network \ to \ use$ 

k Neighborhood size for nearest neighbor sparsification to create the shared NN

graph.

eps Two objects are only reachable from each other if they share at least eps nearest

neighbors.

minPts minimum number of points that share at least eps nearest neighbors for a point

to be considered a core points.

borderPoints should borderPoints be assigned to clusters like in DBSCAN?

return\_gobject boolean: return giotto object (default = TRUE)

set\_seed set seed

seed\_number number for seed

# **Details**

See sNNclust from dbscan package

### Value

giotto object with new clusters appended to cell metadata

do\_permuttest\_random 123

# **Examples**

```
doSNNCluster(gobject)
```

```
{\tt do\_permuttest\_random} \quad \textit{do\_permuttest\_random}
```

# Description

calculate random values

Performs permutation test on subsets of a matrix

# Usage

```
{\tt do\_permuttest\_random(}
  expr_values,
  select_ind,
  other_ind,
  name = "perm_1",
  mean_method,
  offset = 0.1
)
do_permuttest(
  expr_values,
  select_ind,
  other_ind,
  n_{perm} = 1000,
  adjust_method = "fdr",
  mean_method,
  offset = 0.1,
  cores = 2
)
```

# **Examples**

```
do_permuttest_random()
do_permuttest_random()
```

DT\_removeNA

DT\_removeNA

# Description

set NA values to 0 in a data.table object

# Usage

```
DT_removeNA(DT)
```

# Arguments

DT

data.table

124 exportGiottoViewer

 $\verb"estimateImageBg"$ 

estimateImageBg

# **Description**

helps to estimate which color is the background color of your plot

# Usage

```
estimateImageBg(mg_object, top_color_range = 1:50)
```

# **Arguments**

```
mg_object magick image or Giotto image object top_color_range top possible background colors to return
```

### Value

vector of pixel color frequencies and an associated barplot

# **Examples**

```
estimateImageBg(mg_object)
```

exportGiottoViewer

*exportGiottoViewer* 

### **Description**

compute highly variable genes

# Usage

```
exportGiottoViewer(
  gobject,
  output_directory = NULL,
  spat_enr_names = NULL,
  factor_annotations = NULL,
  numeric_annotations = NULL,
  dim_reductions,
  dim_reduction_names,
  expression_values = c("scaled", "normalized", "custom"),
  dim_red_rounding = NULL,
  dim_red_rescale = c(-20, 20),
  expression_rounding = 2,
  overwrite_dir = T,
  verbose = T
```

exprCellCellcom 125

### **Arguments**

gobject giotto object output\_directory directory where to save the files spat\_enr\_names spatial enrichment results to include for annotations factor\_annotations giotto cell annotations to view as factor numeric\_annotations giotto cell annotations to view as numeric dim\_reductions high level dimension reductions to view dim\_reduction\_names specific dimension reduction names expression\_values expression values to use in Viewer dim\_red\_rounding numerical indicating how to round the coordinates dim\_red\_rescale numericals to rescale the coordinates expression\_rounding numerical indicating how to round the expression data overwrite files in the directory if it already existed overwrite\_dir verbose be verbose

# **Details**

Giotto Viewer expects the results from Giotto Analyzer in a specific format, which is provided by this function. To include enrichment results from createSpatialEnrich include the provided spatial enrichment name (default PAGE or rank) and add the gene signature names (.e.g cell types) to the numeric annotations parameter.

### Value

writes the necessary output to use in Giotto Viewer

### **Examples**

exportGiottoViewer(gobject)

exprCellCellcom exprCellCellcom

# Description

Cell-Cell communication scores based on expression only

126 exprCellCellcom

#### Usage

### **Arguments**

```
giotto object to use
gobject
cluster_column cluster column with cell type information
                  number of iterations
random_iter
                  first specific gene set from gene pairs
gene_set_1
                  second specific gene set from gene pairs
gene_set_2
log2FC_addendum
                  addendum to add when calculating log2FC
                  which method to adjust p-values
adjust_method
                  adjust multiple hypotheses at the cell or gene level
adjust_target
verbose
                  verbose
```

# **Details**

Statistical framework to identify if pairs of genes (such as ligand-receptor combinations) are expressed at higher levels than expected based on a reshuffled null distribution of gene expression values, without considering the spatial position of cells. More details will follow soon.

### Value

Cell-Cell communication scores for gene pairs based on expression only

```
exprCellCellcom(gobject)
```

extractNearestNetwork 127

```
{\tt extractNearestNetwork} \ \ \textit{extractNearestNetwork}
```

# **Description**

Extracts a NN-network from a Giotto object

# Usage

```
extractNearestNetwork(
  gobject,
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  output = c("igraph", "data.table")
)
```

# Arguments

# Value

igraph or data.table object

# **Examples**

extractNearestNetwork(gobject)

 ${\sf fDataDT}$ 

fDataDT

# **Description**

show gene metadata

# Usage

```
fDataDT(gobject)
```

# **Arguments**

gobject giotto object

# Value

data.table with gene metadata

#### **Examples**

```
pDataDT(gobject)
```

```
filter {\tt CellProximity Genes}
```

filter Cell Proximity Genes

### **Description**

Filter cell proximity gene scores.

# Usage

```
filterCellProximityGenes(
  cpgObject,
  min_cells = 4,
  min_cells_expr = 1,
  min_int_cells = 4,
  min_int_cells_expr = 1,
  min_fdr = 0.1,
  min_spat_diff = 0.2,
  min_log2_fc = 0.2,
  min_zscore = 2,
  zscores_column = c("cell_type", "genes"),
  direction = c("both", "up", "down")
)
```

# Arguments

```
cpg0bject
                  cell proximity gene score object
\min_{cells}
                  minimum number of source cell type
min_cells_expr minimum expression level for source cell type
min_int_cells
                  minimum number of interacting neighbor cell type
min_int_cells_expr
                  minimum expression level for interacting neighbor cell type
                  minimum adjusted p-value
min_fdr
                  minimum absolute spatial expression difference
min_spat_diff
min_log2_fc
                  minimum log2 fold-change
min_zscore
                  minimum z-score change
zscores_column calculate z-scores over cell types or genes
                  differential expression directions to keep
direction
```

# Value

cpgObject that contains the filtered differential gene scores

```
filterCellProximityGenes(gobject)
```

filterCombinations 129

filterCombinations filterCombinations

### **Description**

Shows how many genes and cells are lost with combinations of thresholds.

# Usage

```
filterCombinations(
  gobject,
  expression_values = c("raw", "normalized", "scaled", "custom"),
  expression_thresholds = c(1, 2),
  gene_det_in_min_cells = c(5, 50),
  min_det_genes_per_cell = c(200, 400),
  scale_x_axis = "identity",
  x_axis_offset = 0,
  scale_y_axis = "identity",
  y_axis_offset = 0,
  show_plot = TRUE,
  return_plot = FALSE,
  save_plot = NA,
  save_param = list(),
  default_save_name = "filterCombinations")
```

### **Arguments**

```
gobject
                  giotto object
expression_values
                  expression values to use
expression_thresholds
                  all thresholds to consider a gene expressed
gene_det_in_min_cells
                  minimum number of cells that should express a gene to consider that gene fur-
                  ther
min_det_genes_per_cell
                  minimum number of expressed genes per cell to consider that cell further
                  ggplot transformation for x-axis (e.g. log2)
scale_x_axis
x_axis_offset
                  x-axis offset to be used together with the scaling transformation
                  ggplot transformation for y-axis (e.g. log2)
scale_y_axis
y_axis_offset
                  y-axis offset to be used together with the scaling transformation
                  show plot
show_plot
return_plot
                  return only ggplot object
                  directly save the plot [boolean]
save_plot
                  list of saving parameters from all_plots_save_function
save_param
default_save_name
```

default save name for saving, don't change, change save\_name in save\_param

filterCPG

#### **Details**

Creates a scatterplot that visualizes the number of genes and cells that are lost with a specific combination of a gene and cell threshold given an arbitrary cutoff to call a gene expressed. This function can be used to make an informed decision at the filtering step with filterGiotto.

### Value

list of data.table and ggplot object

### **Examples**

filterCombinations(gobject)

filterCPG

*filterCPG* 

# **Description**

Filter cell proximity gene scores.

# Usage

```
filterCPG(
  cpgObject,
  min_cells = 4,
  min_cells_expr = 1,
  min_int_cells = 4,
  min_int_cells_expr = 1,
  min_fdr = 0.1,
  min_spat_diff = 0.2,
  min_log2_fc = 0.2,
  min_zscore = 2,
  zscores_column = c("cell_type", "genes"),
  direction = c("both", "up", "down")
)
```

```
cpgObject
                 cell proximity gene score object
                 minimum number of source cell type
min_cells
min_cells_expr minimum expression level for source cell type
min_int_cells
                 minimum number of interacting neighbor cell type
min_int_cells_expr
                 minimum expression level for interacting neighbor cell type
min_fdr
                 minimum adjusted p-value
min_spat_diff
                 minimum absolute spatial expression difference
min_log2_fc
                 minimum log2 fold-change
                 minimum z-score change
min_zscore
zscores_column calculate z-scores over cell types or genes
                 differential expression directions to keep
direction
```

filterDistributions 131

#### Value

cpgObject that contains the filtered differential gene scores

#### **Examples**

```
filterCPG(gobject)
```

filterDistributions filterDistributions

# Description

show gene or cell distribution after filtering on expression threshold

### Usage

```
filterDistributions(
  gobject,
  expression_values = c("raw", "normalized", "scaled", "custom"),
  expression_threshold = 1,
  detection = c("genes", "cells"),
  plot_type = c("histogram", "violin"),
  nr_bins = 30,
  fill_color = "lightblue",
  scale_axis = "identity",
  axis_offset = 0,
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "filterDistributions"
)
```

```
giotto object
gobject
expression_values
                  expression values to use
expression_threshold
                  threshold to consider a gene expressed
detection
                  consider genes or cells
plot_type
                  type of plot
nr_bins
                  number of bins for histogram plot
fill_color
                  fill color for plots
scale_axis
                  ggplot transformation for axis (e.g. log2)
axis_offset
                  offset to be used together with the scaling transformation
show_plot
                  show plot
return_plot
                  return ggplot object
```

132 filterGiotto

#### Value

ggplot object

# **Examples**

```
filterDistributions(gobject)
```

filterGiotto

filterGiotto

# **Description**

filter Giotto object based on expression threshold

# Usage

```
filterGiotto(
  gobject,
  expression_values = c("raw", "normalized", "scaled", "custom"),
  expression_threshold = 1,
  gene_det_in_min_cells = 100,
  min_det_genes_per_cell = 100,
  verbose = F
)
```

# **Arguments**

```
gobject giotto object

expression_values

expression values to use

expression_threshold

threshold to consider a gene expressed

gene_det_in_min_cells

minimum # of cells that need to express a gene

min_det_genes_per_cell

minimum # of genes that need to be detected in a cell

verbose

verbose
```

# **Details**

The function filterCombinations can be used to explore the effect of different parameter values.

### Value

```
giotto object
```

### **Examples**

```
filterGiotto(gobject)
```

```
findCellProximityGenes
```

findCellProximityGenes

# **Description**

Identifies genes that are differentially expressed due to proximity to other cell types.

### Usage

```
findCellProximityGenes(
 gobject,
 expression_values = "normalized",
  selected_genes = NULL,
 cluster_column,
  spatial_network_name = "Delaunay_network",
 minimum_unique_cells = 1,
 minimum_unique_int_cells = 1,
 diff_test = c("permutation", "limma", "t.test", "wilcox"),
 mean_method = c("arithmic", "geometric"),
 offset = 0.1,
 adjust_method = c("bonferroni", "BH", "holm", "hochberg", "hommel", "BY", "fdr",
    "none"),
 nr_permutations = 1000,
 exclude_selected_cells_from_test = T,
 do_parallel = TRUE,
  cores = NA
)
```

```
gobject
                  giotto object
expression_values
                  expression values to use
selected_genes subset of selected genes (optional)
cluster_column name of column to use for cell types
spatial_network_name
                  name of spatial network to use
minimum_unique_cells
                  minimum number of target cells required
minimum_unique_int_cells
                  minimum number of interacting cells required
diff_test
                  which differential expression test
mean_method
                  method to use to calculate the mean
offset
                  offset value to use when calculating log2 ratio
```

adjust\_method which method to adjust p-values

nr\_permutations

number of permutations if diff\_test = permutation

exclude\_selected\_cells\_from\_test

exclude interacting cells other cells

do\_parallel run calculations in parallel with mclapply

cores number of cores to use if do\_parallel = TRUE

#### **Details**

Function to calculate if genes are differentially expressed in cell types when they interact (approximated by physical proximity) with other cell types. The results data.table in the cpgObject contains - at least - the following columns:

- genes: All or selected list of tested genes
- sel: average gene expression in the interacting cells from the target cell type
- other: average gene expression in the NOT-interacting cells from the target cell type
- log2fc: log2 fold-change between sel and other
- · diff: spatial expression difference between sel and other
- p.value: associated p-value
- p.adj: adjusted p-value
- cell\_type: target cell type
- int\_cell\_type: interacting cell type
- nr\_select: number of cells for selected target cell type
- int\_nr\_select: number of cells for interacting cell type
- nr\_other: number of other cells of selected target cell type
- int\_nr\_other: number of other cells for interacting cell type
- unif\_int: cell-cell interaction

### Value

cpgObject that contains the differential gene scores

#### **Examples**

findCellProximityGenes(gobject)

```
find {\tt CellProximityGenes\_per\_interaction} \\ find {\tt CellProximityGenes\_per\_interaction}
```

# Description

Identifies genes that are differentially expressed due to proximity to other cell types.

### Usage

```
findCellProximityGenes_per_interaction(
  expr_values,
 cell_metadata,
 annot_spatnetwork,
 sel_int,
 cluster_column = NULL,
 minimum_unique_cells = 1,
 minimum_unique_int_cells = 1,
 exclude_selected_cells_from_test = T,
 diff_test = c("permutation", "limma", "t.test", "wilcox"),
 mean_method = c("arithmic", "geometric"),
 offset = 0.1,
 adjust_method = "bonferroni",
 nr_permutations = 100,
 cores = 1
)
```

### **Examples**

 ${\tt findCellProximityGenes\_per\_interaction()}$ 

findCPG

findCPG

# **Description**

Identifies genes that are differentially expressed due to proximity to other cell types.

# Usage

```
findCPG(
  gobject,
  expression_values = "normalized",
  selected_genes = NULL,
  cluster_column,
  spatial_network_name = "Delaunay_network",
  minimum_unique_cells = 1,
  minimum_unique_int_cells = 1,
  diff_test = c("permutation", "limma", "t.test", "wilcox"),
```

136 findCPG

# Arguments

```
gobject
                  giotto object
expression_values
                  expression values to use
selected_genes subset of selected genes (optional)
cluster_column name of column to use for cell types
spatial_network_name
                  name of spatial network to use
minimum_unique_cells
                  minimum number of target cells required
minimum_unique_int_cells
                  minimum number of interacting cells required
diff_test
                  which differential expression test
mean_method
                  method to use to calculate the mean
offset
                  offset value to use when calculating log2 ratio
adjust_method
                  which method to adjust p-values
nr_permutations
                  number of permutations if diff_test = permutation
exclude_selected_cells_from_test
                  exclude interacting cells other cells
                  run calculations in parallel with mclapply
do_parallel
                  number of cores to use if do_parallel = TRUE
cores
```

### **Details**

Function to calculate if genes are differentially expressed in cell types when they interact (approximated by physical proximity) with other cell types. The results data.table in the cpgObject contains - at least - the following columns:

- genes: All or selected list of tested genes
- sel: average gene expression in the interacting cells from the target cell type
- other: average gene expression in the NOT-interacting cells from the target cell type
- log2fc: log2 fold-change between sel and other
- diff: spatial expression difference between sel and other
- p.value: associated p-value
- p.adj: adjusted p-value

findGiniMarkers 137

- cell\_type: target cell type
- int\_cell\_type: interacting cell type
- nr\_select: number of cells for selected target cell type
- int\_nr\_select: number of cells for interacting cell type
- nr\_other: number of other cells of selected target cell type
- int\_nr\_other: number of other cells for interacting cell type
- unif\_int: cell-cell interaction

#### Value

cpgObject that contains the differential gene scores

### **Examples**

```
findCPG(gobject)
```

findGiniMarkers

findGiniMarkers

### **Description**

Identify marker genes for selected clusters based on gini detection and expression scores.

# Usage

```
findGiniMarkers(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  cluster_column,
  subset_clusters = NULL,
  group_1 = NULL,
  group_2 = NULL,
  min_expr_gini_score = 0.2,
  min_det_gini_score = 0.2,
  detection_threshold = 0,
  rank_score = 1,
  min_genes = 5
)
```

```
min_expr_gini_score
```

filter on minimum gini coefficient for expression

min\_det\_gini\_score

filter on minimum gini coefficient for detection

detection\_threshold

detection threshold for gene expression

rank\_score rank scores for both detection and expression to include

min\_genes minimum number of top genes to return

### **Details**

Detection of marker genes using the <a href="https://en.wikipedia.org/wiki/Gini\_coefficientgini">https://en.wikipedia.org/wiki/Gini\_coefficientgini</a> coefficient is based on the following steps/principles per gene:

- 1. calculate average expression per cluster
- 2. calculate detection fraction per cluster
- 3. calculate gini-coefficient for av. expression values over all clusters
- 4. calculate gini-coefficient for detection fractions over all clusters
- 5. convert gini-scores to rank scores
- 6. for each gene create combined score = detection rank x expression rank x expr gini-coefficient x detection gini-coefficient
- 7. for each gene sort on expression and detection rank and combined score

As a results "top gini" genes are genes that are very selectivily expressed in a specific cluster, however not always expressed in all cells of that cluster. In other words highly specific, but not necessarily sensitive at the single-cell level.

To perform differential expression between cluster groups you need to specificy cluster IDs to the parameters *group\_1* and *group\_2*.

# Value

data.table with marker genes

### **Examples**

findGiniMarkers(gobject)

findGiniMarkers\_one\_vs\_all

findGiniMarkers\_one\_vs\_all

# **Description**

Identify marker genes for all clusters in a one vs all manner based on gini detection and expression scores.

### Usage

```
findGiniMarkers_one_vs_all(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  cluster_column,
  subset_clusters = NULL,
  min_expr_gini_score = 0.5,
  min_det_gini_score = 0.5,
  detection_threshold = 0,
  rank_score = 1,
  min_genes = 4,
  verbose = TRUE
)
```

# **Arguments**

```
giotto object
gobject
expression_values
                  gene expression values to use
cluster_column clusters to use
subset_clusters
                  selection of clusters to compare
min_expr_gini_score
                  filter on minimum gini coefficient on expression
min_det_gini_score
                  filter on minimum gini coefficient on detection
detection_threshold
                  detection threshold for gene expression
rank_score
                 rank scores for both detection and expression to include
min_genes
                  minimum number of top genes to return
verbose
                  be verbose
```

# Value

data.table with marker genes

# See Also

findGiniMarkers

```
findGiniMarkers_one_vs_all(gobject)
```

140 findMarkers

findMarkers

findMarkers

# **Description**

Identify marker genes for selected clusters.

# Usage

```
findMarkers(
 gobject,
 expression_values = c("normalized", "scaled", "custom"),
  cluster_column = NULL,
 method = c("scran", "gini", "mast"),
  subset_clusters = NULL,
  group_1 = NULL,
 group_2 = NULL,
 min_expr_gini_score = 0.5,
 min_det_gini_score = 0.5,
 detection_threshold = 0,
 rank_score = 1,
 min\_genes = 4,
 group_1_name = NULL,
  group_2_name = NULL,
 adjust_columns = NULL,
)
```

```
gobject
                  giotto object
expression_values
                  gene expression values to use
cluster_column clusters to use
method
                  method to use to detect differentially expressed genes
subset_clusters
                  selection of clusters to compare
                  group 1 cluster IDs from cluster_column for pairwise comparison
group_1
                  group 2 cluster IDs from cluster_column for pairwise comparison
group_2
min_expr_gini_score
                  gini: filter on minimum gini coefficient for expression
min_det_gini_score
                  gini: filter minimum gini coefficient for detection
{\tt detection\_threshold}
                  gini: detection threshold for gene expression
                  gini: rank scores to include
rank_score
                  minimum number of top genes to return (for gini)
min_genes
                  mast: custom name for group_1 clusters
group_1_name
```

findMarkers\_one\_vs\_all

```
group_2_name mast: custom name for group_2 clusters

adjust_columns mast: column in pDataDT to adjust for (e.g. detection rate)

additional parameters for the findMarkers function in scran or zlm function in MAST
```

141

### **Details**

Wrapper for all individual functions to detect marker genes for clusters.

### Value

data.table with marker genes

#### See Also

findScranMarkers, findGiniMarkers and findMastMarkers

### **Examples**

```
findMarkers(gobject)
```

# Description

Identify marker genes for all clusters in a one vs all manner.

# Usage

```
findMarkers_one_vs_all(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  cluster_column,
  subset_clusters = NULL,
  method = c("scran", "gini", "mast"),
  pval = 0.01,
  logFC = 0.5,
  min_genes = 10,
  min_expr_gini_score = 0.5,
  min_det_gini_score = 0.5,
  detection_threshold = 0,
  rank_score = 1,
  adjust_columns = NULL,
  verbose = TRUE,
)
```

### **Arguments**

gobject giotto object

expression\_values

gene expression values to use

cluster\_column clusters to use

subset\_clusters

selection of clusters to compare

method method to use to detect differentially expressed genes

pval scran & mast: filter on minimal p-value

logFC scan & mast: filter on logFC

min\_genes minimum genes to keep per cluster, overrides pval and logFC

min\_expr\_gini\_score

gini: filter on minimum gini coefficient for expression

min\_det\_gini\_score

gini: filter minimum gini coefficient for detection

detection\_threshold

gini: detection threshold for gene expression

rank\_score gini: rank scores to include

adjust\_columns mast: column in pDataDT to adjust for (e.g. detection rate)

verbose be verbose

... additional parameters for the findMarkers function in scran or zlm function in

MAST

### **Details**

Wrapper for all one vs all functions to detect marker genes for clusters.

### Value

data.table with marker genes

### See Also

findScranMarkers\_one\_vs\_all, findGiniMarkers\_one\_vs\_all and findMastMarkers\_one\_vs\_all

```
findMarkers_one_vs_all(gobject)
```

findMastMarkers 143

findMastMarkers findMastMarkers

### **Description**

Identify marker genes for selected clusters based on the MAST package.

# Usage

```
findMastMarkers(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  cluster_column,
  group_1 = NULL,
  group_1_name = NULL,
  group_2 = NULL,
  group_2_name = NULL,
  adjust_columns = NULL,
  ...
)
```

# **Arguments**

```
gobject
                  giotto object
expression_values
                  gene expression values to use
cluster_column clusters to use
                  group 1 cluster IDs from cluster_column for pairwise comparison
group_1
                  custom name for group_1 clusters
group_1_name
                  group 2 cluster IDs from cluster_column for pairwise comparison
group_2
group_2_name
                  custom name for group_2 clusters
adjust_columns column in pDataDT to adjust for (e.g. detection rate)
                  additional parameters for the zlm function in MAST
. . .
```

# **Details**

This is a minimal convenience wrapper around the zlm from the MAST package to detect differentially expressed genes.

# Value

data.table with marker genes

```
findMastMarkers(gobject)
```

# **Description**

Identify marker genes for all clusters in a one vs all manner based on the MAST package.

# Usage

```
findMastMarkers_one_vs_all(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  cluster_column,
  subset_clusters = NULL,
  adjust_columns = NULL,
  pval = 0.001,
  logFC = 1,
  min_genes = 10,
  verbose = TRUE,
  ...
)
```

# **Arguments**

```
gobject
                  giotto object
expression_values
                  gene expression values to use
cluster_column clusters to use
subset_clusters
                  selection of clusters to compare
adjust_columns column in pDataDT to adjust for (e.g. detection rate)
                  filter on minimal p-value
pval
logFC
                  filter on logFC
min_genes
                  minimum genes to keep per cluster, overrides pval and logFC
                  be verbose
verbose
                  additional parameters for the zlm function in MAST
```

# Value

data.table with marker genes

# See Also

findMastMarkers

```
findMastMarkers_one_vs_all(gobject)
```

findNetworkNeighbors 145

```
{\it findNetworkNeighbors} \quad {\it findNetworkNeighbors}
```

#### **Description**

Find the spatial neighbors for a selected group of cells within the selected spatial network.

# Usage

```
findNetworkNeighbors(
  gobject,
  spatial_network_name,
  source_cell_ids = NULL,
  name = "nb_cells"
)
```

## **Arguments**

# Value

data.table

#### **Examples**

findNetworkNeighbors(gobject)

findScranMarkers

findScranMarkers

# Description

Identify marker genes for all or selected clusters based on scran's implementation of findMarkers.

```
findScranMarkers(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  cluster_column,
  subset_clusters = NULL,
  group_1 = NULL,
  group_2 = NULL,
  ...
)
```

#### **Arguments**

#### **Details**

This is a minimal convenience wrapper around the findMarkers function from the scran package.

To perform differential expression between cluster groups you need to specificy cluster IDs to the parameters *group\_1* and *group\_2*.

#### Value

data.table with marker genes

#### **Examples**

```
findScranMarkers(gobject)
```

### **Description**

Identify marker genes for all clusters in a one vs all manner based on scran's implementation of findMarkers.

```
findScranMarkers_one_vs_all(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  cluster_column,
  subset_clusters = NULL,
  pval = 0.01,
  logFC = 0.5,
  min_genes = 10,
  verbose = TRUE,
  ...
)
```

find\_grid\_2D 147

#### **Arguments**

gobject giotto object

expression\_values

gene expression values to use

cluster\_column clusters to use

subset\_clusters

subset of clusters to use

pval filter on minimal p-value

logFC filter on logFC

min\_genes minimum genes to keep per cluster, overrides pval and logFC

verbose be verbose

... additional parameters for the findMarkers function in scran

#### Value

data.table with marker genes

#### See Also

findScranMarkers

# **Examples**

findScranMarkers\_one\_vs\_all(gobject)

find\_grid\_2D

 $find\_grid\_2D$ 

# Description

find grid location in 2D

# Usage

```
find_grid_2D(grid_DT, x_loc, y_loc)
```

find\_grid\_3D

find\_grid\_3D

# Description

find grid location in 3D

```
find_grid_3D(grid_DT, x_loc, y_loc, z_loc)
```

148 find\_grid\_z

find\_grid\_x

find\_grid\_x

# Description

find grid location on x-axis

# Usage

```
find_grid_x(grid_DT, x_loc)
```

find\_grid\_y

find\_grid\_y

# Description

find grid location on y-axis

# Usage

```
find_grid_y(grid_DT, y_loc)
```

find\_grid\_z

 $find\_grid\_z$ 

# Description

find grid location on z-axis

```
find_grid_z(grid_DT, z_loc)
```

FSV\_show 149

FSV\_show FSV\_show

# Description

Visualize spatial varible genes caculated by spatial\_DE

# Usage

```
FSV_show(
  results,
  ms_results = NULL,
  size = c(4, 2, 1),
  color = c("blue", "green", "red"),
  sig_alpha = 0.5,
  unsig_alpha = 0.5
)
```

# **Arguments**

results results caculated by spatial\_DE

ms\_results ms\_results caculated by spatial\_DE

size indicate different levels of qval

color indicate different SV features

sig\_alpha transparency of significant genes

unsig\_alpha transparency of unsignificant genes

# **Details**

Description of parameters.

# Value

ggplot object

# **Examples**

```
FSV_show(results)
```

150 getClusterSimilarity

get10Xmatrix

get10Xmatrix

#### **Description**

This function creates an expression matrix from a 10X structured folder

## Usage

```
get10Xmatrix(path_to_data, gene_column_index = 1)
```

#### **Arguments**

```
path_to_data path to the 10X folder gene_column_index which column from the features or genes .tsv file to use for row ids
```

#### **Details**

A typical 10X folder is named raw\_feature\_bc\_matrix or raw\_feature\_bc\_matrix and tt has 3 files:

- barcodes.tsv(.gz)
- features.tsv(.gz) or genes.tsv(.gz)
- matrix.mtx(.gz)

By default the first column of the features or genes .tsv file will be used, however if multiple annotations are provided (e.g. ensembl gene ids and gene symbols) the user can select another column.

#### Value

sparse expression matrix from 10X

# **Examples**

```
get10Xmatrix(path_to_data)
```

```
getClusterSimilarity
getClusterSimilarity
```

#### **Description**

Creates data.table with pairwise correlation scores between each cluster.

```
getClusterSimilarity(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  cluster_column,
  cor = c("pearson", "spearman")
)
```

getDendrogramSplits 151

#### **Arguments**

```
gobject giotto object
expression_values
expression values to use
cluster_column name of column to use for clusters
cor correlation score to calculate distance
```

#### **Details**

Creates data.table with pairwise correlation scores between each cluster and the group size (# of cells) for each cluster. This information can be used together with mergeClusters to combine very similar or small clusters into bigger clusters.

#### Value

data.table

#### **Examples**

```
getClusterSimilarity(gobject)
```

```
getDendrogramSplits getDendrogramSplits
```

# Description

Split dendrogram at each node and keep the leave (label) information..

# Usage

```
getDendrogramSplits(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  cluster_column,
  cor = c("pearson", "spearman"),
  distance = "ward.D",
  h = NULL,
  h_color = "red",
  show_dend = TRUE,
  verbose = TRUE
)
```

#### **Arguments**

```
gobject giotto object
expression_values
expression values to use
cluster_column name of column to use for clusters
cor correlation score to calculate distance
```

152 getDistinctColors

distance distance method to use for hierarchical clustering

h height of horizontal lines to plot

h\_color color of horizontal lines

show\_dend show dendrogram

verbose be verbose

#### **Details**

Creates a data.table with three columns and each row represents a node in the dendrogram. For each node the height of the node is given together with the two subdendrograms. This information can be used to determine in a hierarchical manner differentially expressed marker genes at each node.

#### Value

data.table object

### **Examples**

getDendrogramSplits(gobject)

getDistinctColors

# Description

Returns a number of distint colors based on the RGB scale

# Usage

```
getDistinctColors(n)
```

# Arguments

n number of colors wanted

#### Value

number of distinct colors

getGiottoImage 153

 ${\tt getGiottoImage}$ 

getGiottoImage

# Description

get get a giotto image from a giotto object

# Usage

```
getGiottoImage(gobject, image_name)
```

# **Arguments**

gobject

giotto object

image\_name

 $name\ of\ giotto\ image\ show {\tt GiottoImageNames}$ 

#### Value

a giotto image

# **Examples**

```
getGiottoImage(gobject)
```

get\_os

get\_os

# Description

 $return\ the\ type\ of\ operating\ system,\ see\ https://conjugateprior.org/2015/06/identifying-the-os-from-r/$ 

# Usage

```
get_os()
```

## Value

character osx, linux or windows

giotto-class

S4 giotto Class

# Description

Framework of giotto object to store and work with spatial expression data

#### **Slots**

```
raw_exprs raw expression counts
norm_expr normalized expression counts
norm_scaled_expr normalized and scaled expression counts
custom_expr custom normalized counts
spatial_locs spatial location coordinates for cells
cell_metadata metadata for cells
gene_metadata metadata for genes
cell_ID unique cell IDs
gene_ID unique gene IDs
spatial_network spatial network in data.table/data.frame format
spatial_grid spatial grid in data.table/data.frame format
spatial_enrichment slot to save spatial enrichment-like results
dimension_reduction slot to save dimension reduction coordinates
nn_network nearest neighbor network in igraph format
images slot to store giotto images
parameters slot to save parameters that have been used
instructions slot for global function instructions
offset_file offset file used to stitch together image fields
OS_platform Operating System to run Giotto analysis on
```

 $heatmSpatialCorGenes \quad \textit{heatmSpatialCorGenes}$ 

## **Description**

Create heatmap of spatially correlated genes

heatmSpatialCorGenes 155

#### Usage

```
heatmSpatialCorGenes(
  gobject,
  spatCorObject,
  use_clus_name = NULL,
  show_cluster_annot = TRUE,
  show_row_dend = T,
  show_column_dend = F,
  show_row_names = F,
  show_column_names = F,
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "heatmSpatialCorGenes",
  ...
)
```

## **Arguments**

```
gobject
                 giotto object
                 spatial correlation object
spatCorObject
                 name of clusters to visualize (from clusterSpatialCorGenes())
use_clus_name
show_cluster_annot
                 show cluster annotation on top of heatmap
                 show row dendrogram
show_row_dend
show_column_dend
                 show column dendrogram
show_row_names show row names
show_column_names
                 show column names
show_plot
                 show plot
return_plot
                 return ggplot object
                 directly save the plot [boolean]
save_plot
                 list of saving parameters from all_plots_save_function
save_param
default_save_name
                 default save name for saving, don't change, change save_name in save_param
                 additional parameters to the Heatmap function from ComplexHeatmap
. . .
```

#### Value

Heatmap generated by ComplexHeatmap

#### **Examples**

```
heatmSpatialCorGenes(gobject)
```

 $hyperGeometric Enrich \qquad hyperGeometric Enrich$ 

#### **Description**

Function to calculate gene signature enrichment scores per spatial position using a hypergeometric test.

# Usage

```
hyperGeometricEnrich(
  gobject,
  sign_matrix,
  expression_values = c("normalized", "scaled", "custom"),
  reverse_log_scale = TRUE,
  logbase = 2,
  top_percentage = 5,
  output_enrichment = c("original", "zscore")
)
```

#### **Arguments**

```
gobject Giotto object

sign_matrix Matrix of signature genes for each cell type / process

expression_values

expression values to use

reverse_log_scale

reverse expression values from log scale

logbase log base to use if reverse_log_scale = TRUE

top_percentage percentage of cells that will be considered to have gene expression with matrix binarization

output_enrichment

how to return enrichment output
```

## Details

The enrichment score is calculated based on the p-value from the hypergeometric test, -log10(p-value).

#### Value

data.table with enrichment results

#### **Examples**

hyperGeometricEnrich(gobject)

```
insert {\tt CrossSectionGenePlot3D}
```

insertCrossSectionGenePlot3D

#### **Description**

Visualize cells and gene expression in a virtual cross section according to spatial coordinates

#### Usage

```
insertCrossSectionGenePlot3D(
 gobject,
 crossSection_obj = NULL,
 name = NULL,
 spatial_network_name = "Delaunay_network",
 mesh_grid_color = "#1f77b4",
 mesh_grid_width = 3,
 mesh_grid_style = "dot",
 sdimx = "sdimx",
  sdimy = "sdimy",
  sdimz = "sdimz",
  show_other_cells = F,
 axis_scale = c("cube", "real", "custom"),
  custom_ratio = NULL,
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
 default_save_name = "spatGenePlot3D_with_cross_section",
)
```

#### **Arguments**

```
gobject
                  giotto object
                  name of virtual cross section to use
name
spatial_network_name
                  name of spatial network to use
mesh_grid_color
                  color for the meshgrid lines
mesh_grid_width
                  width for the meshgrid lines
mesh_grid_style
                  style for the meshgrid lines
                  x-axis dimension name (default = 'sdimx')
sdimx
                  y-axis dimension name (default = 'sdimy')
sdimy
sdimz
                  z-axis dimension name (default = 'sdimy')
show_other_cells
                  display not selected cells
```

#### **Details**

Description of parameters.

#### Value

ggplot

#### **Examples**

insertCrossSectionGenePlot3D(gobject)

```
insert {\tt CrossSectionSpatPlot3D} \\ insert {\tt CrossSectionSpatPlot3D}
```

## **Description**

Visualize the meshgrid lines of cross section together with cells

```
insertCrossSectionSpatPlot3D(
  gobject,
  crossSection_obj = NULL,
  name = NULL,
  spatial_network_name = "Delaunay_network",
  mesh_grid_color = "#1f77b4",
  mesh_grid_width = 3,
  mesh_grid_style = "dot",
  sdimx = "sdimx",
  sdimy = "sdimy",
  sdimz = "sdimz",
  show_other_cells = F,
  axis_scale = c("cube", "real", "custom"),
  custom_ratio = NULL,
  default_save_name = "spat3D_with_cross_section",
)
```

jackstrawPlot 159

## **Arguments**

```
giotto object
gobject
name
                  name of virtual cross section to use
spatial_network_name
                  name of spatial network to use
mesh_grid_color
                  color for the meshgrid lines
mesh_grid_width
                  width for the meshgrid lines
mesh_grid_style
                  style for the meshgrid lines
sdimx
                  x-axis dimension name (default = 'sdimx')
sdimy
                  y-axis dimension name (default = 'sdimy')
sdimz
                  z-axis dimension name (default = 'sdimy')
show_other_cells
                  display not selected cells
axis_scale
                  axis_scale
custom_ratio
                  custom_ratio
default_save_name
                  default save name for saving, don't change, change save_name in save_param
                  parameters for spatPlot3D
```

## **Details**

Description of parameters.

## Value

ggplot

#### **Examples**

insertCrossSectionSpatPlot3D(gobject)

jackstrawPlot jackstrawPlot

# Description

identify significant prinicipal components (PCs)

jackstrawPlot

#### Usage

```
jackstrawPlot(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  reduction = c("cells", "genes"),
  genes_to_use = NULL,
  center = FALSE,
  scale_unit = FALSE,
  ncp = 20,
  ylim = c(0, 1),
  iter = 10,
  threshold = 0.01,
  verbose = T,
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "jackstrawPlot"
)
```

#### **Arguments**

```
gobject
                  giotto object
expression_values
                  expression values to use
                  cells or genes
reduction
                  subset of genes to use for PCA
genes_to_use
                  center data before PCA
center
scale_unit
                  scale features before PCA
ncp
                  number of principal components to calculate
                  y-axis limits on jackstraw plot
ylim
                  number of interations for jackstraw
iter
                  p-value threshold to call a PC significant
threshold
verbose
                  show progress of jackstraw method
                  show plot
show_plot
return_plot
                  return ggplot object
                  directly save the plot [boolean]
save_plot
save_param
                  list of saving parameters from all_plots_save_function()
default_save_name
```

# **Details**

The Jackstraw method uses the permutationPA function. By systematically permuting genes it identifies robust, and thus significant, PCs.

default save name for saving, don't change, change save\_name in save\_param

kmeans\_binarize 161

#### Value

ggplot object for jackstraw method

#### **Examples**

```
jackstrawPlot(gobject)
```

kmeans\_binarize

kmeans\_binarize

# Description

create binarized scores from a vector using kmeans

#### Usage

```
kmeans_binarize(x, nstart = 3, iter.max = 10)
```

loadHMRF

loadHMRF

# Description

load previous HMRF

### Usage

```
loadHMRF(
  name_used = "test",
  output_folder_used,
  k_used = 10,
  betas_used,
  python_path_used
)
```

# Arguments

```
name_used name of HMRF that was run
output_folder_used
output folder that was used
k_used number of HMRF domains that was tested
betas_used betas that were tested
python_path_used
python path that was used
```

### **Details**

Description of HMRF parameters ...

#### Value

reloads a previous ran HMRF from doHRMF

#### **Examples**

loadHMRF(gobject)

makeSignMatrixPAGE

make Sign Matrix PAGE

#### **Description**

Function to convert a list of signature genes (e.g. for cell types or processes) into a binary matrix format that can be used with the PAGE enrichment option. Each cell type or process should have a vector of cell-type or process specific genes. These vectors need to be combined into a list (sign\_list). The names of the cell types or processes that are provided in the list need to be given (sign\_names).

#### Usage

```
makeSignMatrixPAGE(sign_names, sign_list)
```

#### **Arguments**

sign\_names vector with names for each provided gene signature

sign\_list list of genes (signature)

#### Value

matrix

## See Also

**PAGEEnrich** 

#### **Examples**

 ${\tt makeSignMatrixPAGE()}$ 

makeSignMatrixRank 163

makeSignMatrixRank makeSignMatrixRank

#### **Description**

Function to convert a single-cell count matrix and a corresponding single-cell cluster vector into a rank matrix that can be used with the Rank enrichment option.

#### Usage

```
makeSignMatrixRank(sc_matrix, sc_cluster_ids, gobject = NULL)
```

#### **Arguments**

```
sc_matrix matrix of single-cell RNAseq expression data
sc_cluster_ids vector of cluster ids
gobject if giotto object is given then only genes present in both datasets will be considered
```

#### Value

matrix

#### See Also

rankEnrich

# **Examples**

```
makeSignMatrixRank()
```

```
make_simulated_network
```

make\_simulated\_network

## **Description**

Simulate random network.

# Usage

```
make_simulated_network(
  gobject,
  spatial_network_name = "Delaunay_network",
  cluster_column,
  number_of_simulations = 100
)
```

# **Examples**

```
{\tt make\_simulated\_network(gobject)}
```

164 mergeClusters

 ${\tt mergeClusters}$ 

mergeClusters

#### **Description**

Merge selected clusters based on pairwise correlation scores and size of cluster.

## Usage

```
mergeClusters(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  cluster_column,
  cor = c("pearson", "spearman"),
  new_cluster_name = "merged_cluster",
  min_cor_score = 0.8,
  max_group_size = 20,
  force_min_group_size = 10,
  max_sim_clusters = 10,
  return_gobject = TRUE,
  verbose = TRUE
)
```

#### **Arguments**

```
giotto object
gobject
expression_values
                  expression values to use
cluster_column name of column to use for clusters
                  correlation score to calculate distance
cor
new_cluster_name
                  new name for merged clusters
                 min correlation score to merge pairwise clusters
min_cor_score
max_group_size max cluster size that can be merged
force_min_group_size
                  size of clusters that will be merged with their most similar neighbor(s)
max_sim_clusters
                  maximum number of clusters to potentially merge to reach force_min_group_size
return_gobject return giotto object
                  be verbose
verhose
```

#### **Details**

Merge selected clusters based on pairwise correlation scores and size of cluster. To avoid large clusters to merge the max\_group\_size can be lowered. Small clusters can be forcibly merged with their most similar pairwise cluster by adjusting the force\_min\_group\_size parameter. Clusters smaller than this value will be merged independent on the provided min\_cor\_score value. The force\_min\_group\_size might not always be reached if clusters have already been merged before A giotto object is returned by default, if FALSE then the merging vector will be returned.

my\_arowMeans 165

#### Value

Giotto object

#### **Examples**

```
mergeClusters(gobject)
```

my\_arowMeans

my\_arowMeans

# Description

arithmic rowMeans that works for a single column

# Usage

```
my_arowMeans(x)
```

# **Examples**

my\_arowMeans(x)

my\_growMeans

my\_growMeans

# Description

geometric rowMeans that works for a single column

# Usage

```
my\_growMeans(x, offset = 0.1)
```

# **Examples**

```
my\_growMeans(x)
```

my\_rowMeans

my\_rowMeans

# Description

arithmic or geometric rowMeans that works for a single column

# Usage

```
my_rowMeans(x, method = c("arithmic", "geometric"), offset = 0.1)
```

# **Examples**

```
my_rowMeans(x)
```

node\_clusters

nnDT\_to\_kNN

 $nnDT\_to\_kNN$ 

# Description

Convert a nearest network data.table to a kNN object

# Usage

```
nnDT_to_kNN(nnDT)
```

# **Arguments**

nnDT

nearest neighbor network in data.table format

#### Value

kNN object

node\_clusters

node\_clusters

# Description

Merge selected clusters based on pairwise correlation scores and size of cluster.

# Usage

```
node_clusters(hclus_obj, verbose = TRUE)
```

# Arguments

hclus\_obj hclus object verbose be verbose

# Value

list of splitted dendrogram nodes from high to low node height

# **Examples**

```
node_clusters(hclus_obj)
```

normalizeGiotto 167

normalizeGiotto	normalizeGiotto
-----------------	-----------------

### **Description**

fast normalize and/or scale expresion values of Giotto object

#### Usage

```
normalizeGiotto(
  gobject,
  norm_methods = c("standard", "osmFISH"),
  library_size_norm = TRUE,
  scalefactor = 6000,
  log_norm = TRUE,
  log_offset = 1,
  logbase = 2,
  scale_genes = T,
  scale_cells = T,
  scale_order = c("first_genes", "first_cells"),
  verbose = F
)
```

#### **Arguments**

```
gobject
                  giotto object
norm_methods
                  normalization method to use
library_size_norm
                  normalize cells by library size
scalefactor
                  scale factor to use after library size normalization
                  transform values to log-scale
log_norm
log_offset
                  offset value to add to expression matrix, default = 1
logbase
                  log base to use to log normalize expression values
                  z-score genes over all cells
scale_genes
scale_cells
                  z-score cells over all genes
scale_order
                  order to scale genes and cells
verbose
                  be verbose
```

#### **Details**

Currently there are two 'methods' to normalize your raw counts data.

A. The standard method follows the standard protocol which can be adjusted using the provided parameters and follows the following order:

- 1. Data normalization for total library size and scaling by a custom scale-factor.
- 2. Log transformation of data.
- 3. Z-scoring of data by genes and/or cells.

168 PAGEEnrich

- B. The normalization method as provided by the osmFISH paper is also implemented:
  - 1. First normalize genes, for each gene divide the counts by the total gene count and multiply by the total number of genes.
  - 2. Next normalize cells, for each cell divide the normalized gene counts by the total counts per cell and multiply by the total number of cells.

This data will be saved in the Giotto slot for custom expression.

#### Value

giotto object

#### **Examples**

```
normalizeGiotto(gobject)
```

**PAGEEnrich** 

**PAGEEnrich** 

# Description

Function to calculate gene signature enrichment scores per spatial position using PAGE.

#### Usage

```
PAGEEnrich(
  gobject,
  sign_matrix,
  expression_values = c("normalized", "scaled", "custom"),
  reverse_log_scale = TRUE,
  logbase = 2,
  output_enrichment = c("original", "zscore")
)
```

## **Arguments**

```
gobject Giotto object

sign_matrix Matrix of signature genes for each cell type / process

expression_values

expression values to use

reverse_log_scale

reverse expression values from log scale

logbase log base to use if reverse_log_scale = TRUE

output_enrichment

how to return enrichment output
```

pca\_giotto 169

#### **Details**

sign\_matrix: a binary matrix with genes as row names and cell-types as column names. Alternatively a list of signature genes can be provided to makeSignMatrixPAGE, which will create the matrix for you.

The enrichment Z score is calculated by using method (PAGE) from Kim SY et al., BMC bioinformatics, 2005 as  $Z = ((Sm^{\circ}mu) * m^{(1/2)})/delta$ . For each gene in each spot, mu is the fold change values versus the mean expression and delta is the standard deviation. Sm is the mean fold change value of a specific marker gene set and m is the size of a given marker gene set.

#### Value

data.table with enrichment results

#### See Also

```
makeSignMatrixPAGE
```

#### **Examples**

PAGEEnrich(gobject)

pca\_giotto

pca\_giotto

## **Description**

performs PCA based on Rfast

# Usage

```
pca_giotto(mymatrix, center = T, scale = T, k = 50)
```

#### **Arguments**

matrix or object that can be converted to matrix mymatrix

center center data scale scale features

k number of principal components to calculate

#### Value

list of eigenvalues, eigenvectors and pca coordinates

170 plotCCcomDotplot

pDataDT

pDataDT

#### **Description**

show cell metadata

#### Usage

```
pDataDT(gobject)
```

#### **Arguments**

gobject

giotto object

#### Value

data.table with cell metadata

## **Examples**

```
pDataDT(gobject)
```

plotCCcomDotplot

plotCCcomDotplot

# Description

Plots dotplot for ligand-receptor communication scores in cell-cell interactions

```
plotCCcomDotplot(
  gobject,
  comScores,
  selected_LR = NULL,
  selected_cell_LR = NULL,
  show_LR_names = TRUE,
  show_cell_LR_names = TRUE,
  cluster_on = c("PI", "LR_expr", "log2fc"),
  cor_method = c("pearson", "kendall", "spearman"),
 aggl_method = c("ward.D", "ward.D2", "single", "complete", "average", "mcquitty",
    "median", "centroid"),
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "plotCCcomDotplot"
```

plotCCcomHeatmap 171

#### **Arguments**

```
gobject
                  giotto object
                  communinication scores from exprCellCellcom or spatCellCellcom
comScores
selected_LR
                  selected ligand-receptor combinations
selected_cell_LR
                  selected cell-cell combinations for ligand-receptor combinations
                  show ligand-receptor names
show_LR_names
show_cell_LR_names
                  show cell-cell names
                  values to use for clustering of cell-cell and ligand-receptor pairs
cluster_on
                  correlation method used for clustering
cor\_method
aggl_method
                  agglomeration method used by hclust
show_plot
                  show plots
return_plot
                  return plotting object
                  directly save the plot [boolean]
save_plot
                  list of saving parameters from all_plots_save_function
save_param
default_save_name
                  default save name for saving, don't change, change save_name in save_param
show
                  values to show on heatmap
```

#### Value

ggplot

#### **Examples**

```
plotCCcomDotplot(CPGscores)
```

plotCCcomHeatmap plotCCcomHeatmap

## **Description**

Plots heatmap for ligand-receptor communication scores in cell-cell interactions

```
plotCCcomHeatmap(
  gobject,
  comScores,
  selected_LR = NULL,
  selected_cell_LR = NULL,
  show_LR_names = TRUE,
  show_cell_LR_names = TRUE,
  show = c("PI", "LR_expr", "log2fc"),
  cor_method = c("pearson", "kendall", "spearman"),
  aggl_method = c("ward.D", "ward.D2", "single", "complete", "average", "mcquitty",
```

```
"median", "centroid"),
show_plot = NA,
return_plot = NA,
save_plot = NA,
save_param = list(),
default_save_name = "plotCCcomHeatmap")
```

# Arguments

gobject giotto object comScores  $communinication\ scores\ from\ expr\cell\cellcom\ or\ spat\cell\cellcom$ selected ligand-receptor combinations selected\_LR selected\_cell\_LR selected cell-cell combinations for ligand-receptor combinations show\_LR\_names show ligand-receptor names show\_cell\_LR\_names show cell-cell names show values to show on heatmap cor\_method correlation method used for clustering agglomeration method used by hclust  $aggl_method$ show\_plot show plots return\_plot return plotting object save\_plot directly save the plot [boolean] list of saving parameters from all\_plots\_save\_function save\_param default\_save\_name

default save name for saving, don't change, change save\_name in save\_param

#### Value

ggplot

# **Examples**

```
plotCCcomHeatmap(CPGscores)
```

```
plotCellProximityGenes
```

plotCellProximityGenes

# Description

Create visualization for cell proximity gene scores

#### Usage

```
plotCellProximityGenes(
  gobject,
  cpgObject,
  method = c("volcano", "cell_barplot", "cell-cell", "cell_sankey", "heatmap",
    "dotplot"),
  min_cells = 4,
  min_cells_expr = 1,
  min_int_cells = 4,
  min_int_cells_expr = 1,
  min_fdr = 0.1,
  min_spat_diff = 0.2,
  min_log2_fc = 0.2,
  min_zscore = 2,
  zscores_column = c("cell_type", "genes"),
  direction = c("both", "up", "down"),
  cell_color_code = NULL,
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "plotCellProximityGenes"
```

#### **Arguments**

```
gobject
                  giotto object
cpgObject
                  cell proximity gene score object
method
                  plotting method to use
min_cells
                  minimum number of source cell type
min_cells_expr minimum expression level for source cell type
min_int_cells
                  minimum number of interacting neighbor cell type
min_int_cells_expr
                  minimum expression level for interacting neighbor cell type
min_fdr
                  minimum adjusted p-value
min_spat_diff
                  minimum absolute spatial expression difference
                  minimum log2 fold-change
min_log2_fc
min_zscore
                  minimum z-score change
zscores_column calculate z-scores over cell types or genes
direction
                  differential expression directions to keep
cell_color_code
                  vector of colors with cell types as names
show_plot
                  show plots
return_plot
                  return plotting object
save_plot
                  directly save the plot [boolean]
                  list of saving parameters from all_plots_save_function
save_param
default_save_name
```

default save name for saving, don't change, change save\_name in save\_param

174 plotCombineCCcom

#### Value

plot

#### **Examples**

```
plotCellProximityGenes(CPGscores)
```

plotCombineCCcom

plotCombineCCcom

#### **Description**

Create visualization for combined (pairwise) cell proximity gene scores

### Usage

```
plotCombineCCcom(
  gobject,
  combCCcom,
  selected_LR = NULL,
  selected_cell_LR = NULL,
  detail_plot = T,
  simple_plot = F,
  simple_plot_facet = c("interaction", "genes"),
  facet_scales = "fixed",
  facet_ncol = length(selected_LR),
  facet_nrow = length(selected_cell_LR),
  colors = c("#9932CC", "#FF8C00"),
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "plotCombineCCcom"
)
```

#### **Arguments**

```
gobject
                  giotto object
{\sf combCCcom}
                  combined communcation scores, output from combCCcom()
selected_LR
                  selected ligand-receptor pair
selected_cell_LR
                  selected cell-cell interaction pair for ligand-receptor pair
{\tt detail\_plot}
                  show detailed info in both interacting cell types
simple_plot
                  show a simplified plot
simple_plot_facet
                  facet on interactions or genes with simple plot
facet_scales
                  ggplot facet scales paramter
facet_ncol
                  ggplot facet ncol parameter
```

```
facet_nrow ggplot facet nrow parameter

colors vector with two colors to use

show_plot show plots

return_plot return plotting object

save_plot directly save the plot [boolean]

save_param list of saving parameters from all_plots_save_function

default_save_name

default save name for saving, don't change, change save_name in save_param
```

#### Value

ggplot

#### **Examples**

```
plotCombineCCcom(CPGscores)
```

```
plot {\tt Combine Cell Cell Communication} \\ plot {\tt Combine Cell Cell Communication} \\
```

# Description

Create visualization for combined (pairwise) cell proximity gene scores

```
plotCombineCellCellCommunication(
  gobject,
  combCCcom,
  selected_LR = NULL,
  selected_cell_LR = NULL,
  detail_plot = T,
  simple_plot = F,
  simple_plot_facet = c("interaction", "genes"),
  facet_scales = "fixed",
  facet_ncol = length(selected_LR),
  facet_nrow = length(selected_cell_LR),
  colors = c("#9932CC", "#FF8C00"),
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "plotCombineCellCellCommunication"
```

#### **Arguments**

```
giotto object
gobject
combCCcom
                  combined communcation scores, output from combCCcom()
selected_LR
                  selected ligand-receptor pair
selected_cell_LR
                  selected cell-cell interaction pair for ligand-receptor pair
detail_plot
                  show detailed info in both interacting cell types
                  show a simplified plot
simple_plot
simple_plot_facet
                  facet on interactions or genes with simple plot
facet_scales
                  ggplot facet scales paramter
                  ggplot facet ncol parameter
facet_ncol
facet_nrow
                  ggplot facet nrow parameter
colors
                  vector with two colors to use
show_plot
                  show plots
return_plot
                  return plotting object
                  directly save the plot [boolean]
save_plot
save_param
                  list of saving parameters from all_plots_save_function
default_save_name
                  default save name for saving, don't change, change save_name in save_param
```

#### Value

ggplot

#### **Examples**

```
plotCombineCellCellCommunication(CPGscores)
```

```
plot {\tt Combine Cell Proximity Genes} \\ plot {\tt Combine Cell Proximity Genes}
```

#### **Description**

Create visualization for combined (pairwise) cell proximity gene scores

```
plotCombineCellProximityGenes(
  gobject,
  combCpgObject,
  selected_interactions = NULL,
  selected_gene_to_gene = NULL,
  detail_plot = T,
  simple_plot = F,
```

```
simple_plot_facet = c("interaction", "genes"),
facet_scales = "fixed",
facet_ncol = length(selected_gene_to_gene),
facet_nrow = length(selected_interactions),
colors = c("#9932CC", "#FF8C00"),
show_plot = NA,
return_plot = NA,
save_plot = NA,
save_param = list(),
default_save_name = "plotCombineCPG"
)
```

#### **Arguments**

```
gobject
                  giotto object
combCpgObject
                  CPGscores, output from combineCellProximityGenes()
selected_interactions
                  interactions to show
selected_gene_to_gene
                  pairwise gene combinations to show
detail_plot
                  show detailed info in both interacting cell types
simple_plot
                  show a simplified plot
simple_plot_facet
                  facet on interactions or genes with simple plot
facet_scales
                  ggplot facet scales paramter
facet_ncol
                  ggplot facet ncol parameter
facet_nrow
                  ggplot facet nrow parameter
colors
                  vector with two colors to use
show_plot
                  show plots
return_plot
                  return plotting object
                  directly save the plot [boolean]
save_plot
save_param
                  list of saving parameters from all_plots_save_function
default_save_name
                  default save name for saving, don't change, change save_name in save_param
```

#### Value

ggplot

## **Examples**

```
plotCombineCellProximityGenes(CPGscores)
```

178 plotCombineCPG

plotCombineCPG plotCombineCPG

#### **Description**

Create visualization for combined (pairwise) cell proximity gene scores

## Usage

```
plotCombineCPG(
  gobject,
  combCpgObject,
  selected_interactions = NULL,
  selected_gene_to_gene = NULL,
  detail_plot = T,
  simple_plot = F,
  simple_plot_facet = c("interaction", "genes"),
  facet_scales = "fixed",
  facet_ncol = length(selected_gene_to_gene),
  facet_nrow = length(selected_interactions),
  colors = c("#9932CC", "#FF8C00"),
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "plotCombineCPG"
)
```

## **Arguments**

```
gobject
                 giotto object
combCpgObject
                 CPGscores, output from combineCellProximityGenes()
selected_interactions
                 interactions to show
selected_gene_to_gene
                 pairwise gene combinations to show
detail_plot
                 show detailed info in both interacting cell types
simple_plot
                 show a simplified plot
simple_plot_facet
                 facet on interactions or genes with simple plot
facet_scales
                 ggplot facet scales paramter
facet_ncol
                 ggplot facet ncol parameter
facet_nrow
                 ggplot facet nrow parameter
colors
                 vector with two colors to use
show_plot
                 show plots
return_plot
                 return plotting object
                 directly save the plot [boolean]
save_plot
save_param
                 list of saving parameters from all_plots_save_function
default_save_name
```

default save name for saving, don't change, change save\_name in save\_param

plotCPG 179

#### Value

ggplot

#### **Examples**

```
plotCombineCPG(CPGscores)
```

plotCPG

plotCPG

#### **Description**

Create visualization for cell proximity gene scores

## Usage

```
plotCPG(
  gobject,
  cpgObject,
  method = c("volcano", "cell_barplot", "cell-cell", "cell_sankey", "heatmap",
    "dotplot"),
  min_cells = 5,
  min_cells_expr = 1,
  min_int_cells = 3,
  min_int_cells_expr = 1,
  min_fdr = 0.05,
  min_spat_diff = 0.2,
  min_log2_fc = 0.2,
  min_zscore = 2,
  zscores_column = c("cell_type", "genes"),
  direction = c("both", "up", "down"),
  cell_color_code = NULL,
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "plotCPG"
)
```

## Arguments

```
gobject giotto object
cell proximity gene score object
method plotting method to use
min_cells minimum number of source cell type
min_cells_expr minimum expression level for source cell type
min_int_cells minimum number of interacting neighbor cell type
min_int_cells_expr
minimum expression level for interacting neighbor cell type
```

180 plotGiottoImage

min\_fdr minimum adjusted p-value

min\_spat\_diff minimum absolute spatial expression difference

min\_log2\_fc minimum log2 fold-change min\_zscore minimum z-score change

zscores\_column calculate z-scores over cell types or genes direction differential expression directions to keep

cell\_color\_code

vector of colors with cell types as names

show\_plot show plots

return\_plot return plotting object

save\_plot directly save the plot [boolean]

save\_param list of saving parameters from all\_plots\_save\_function

default\_save\_name

default save name for saving, don't change, change save\_name in save\_param

#### Value

plot

# **Examples**

plotCPG(CPGscores)

plotGiottoImage plotGiottoImage

## **Description**

get plot a giotto image from a giotto object

#### Usage

```
plotGiottoImage(gobject, image_name)
```

# **Arguments**

gobject giotto object

image\_name name of giotto image showGiottoImageNames

# Value

plot

#### **Examples**

```
plotGiottoImage(gobject)
```

plotHeatmap 181

plotHeatmap

plotHeatmap

#### **Description**

Creates heatmap for genes and clusters.

#### Usage

```
plotHeatmap(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  genes,
  cluster_column = NULL,
  cluster_order = c("size", "correlation", "custom"),
  cluster_custom_order = NULL,
  cluster_color_code = NULL,
  cluster_cor_method = "pearson",
  cluster_hclust_method = "ward.D"
  gene_order = c("correlation", "custom"),
  gene_custom_order = NULL,
  gene_cor_method = "pearson",
  gene_hclust_method = "complete",
  show_values = c("rescaled", "z-scaled", "original"),
  size_vertical_lines = 1.1,
  gradient_colors = c("blue", "yellow", "red"),
  gene_label_selection = NULL,
  axis_text_y_size = NULL,
  legend_nrows = 1,
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "plotHeatmap"
)
```

```
gobject giotto object
expression_values
expression values to use
genes genes to use
cluster_column name of column to use for clusters
cluster_order method to determine cluster order
cluster_custom_order
custom order for clusters
cluster_color_code
color code for clusters
cluster_cor_method
method for cluster correlation
```

182 plotHeatmap

```
cluster_hclust_method
                  method for hierarchical clustering of clusters
                  method to determine gene order
gene_order
gene_custom_order
                  custom order for genes
{\tt gene\_cor\_method}
                  method for gene correlation
gene_hclust_method
                  method for hierarchical clustering of genes
show_values
                  which values to show on heatmap
size_vertical_lines
                  sizes for vertical lines
gradient_colors
                  colors for heatmap gradient
gene_label_selection
                  subset of genes to show on y-axis
axis_text_y_size
                  size for y-axis text
legend_nrows
                  number of rows for the cluster legend
show_plot
                  show plot
return_plot
                  return ggplot object
                  directly save the plot [boolean]
save_plot
save_param
                  list of saving parameters, see showSaveParameters
default_save_name
                  default save name
```

### Details

If you want to display many genes there are 2 ways to proceed:

- 1. set axis\_text\_y\_size to a really small value and show all genes
- 2. provide a subset of genes to display to gene\_label\_selection

#### Value

ggplot

```
## Not run:
plotHeatmap(gobject)
## End(Not run)
```

plotICG 183

plotICG plotICG

## **Description**

Create barplot to visualize interaction changed genes

## Usage

```
plotICG(
   gobject,
   cpgObject,
   source_type,
   source_markers,
   ICG_genes,
   cell_color_code = NULL,
   show_plot = NA,
   return_plot = NA,
   save_plot = NA,
   save_param = list(),
   default_save_name = "plotICG"
)
```

# Arguments

```
gobject
                  giotto object
                  cell proximity gene score object
cpgObject
source_type
                  cell type of the source cell
source_markers markers for the source cell type
ICG_genes
                  named character vector of ICG genes
cell_color_code
                  cell color code for the interacting cell types
show_plot
                  show plots
return_plot
                  return plotting object
                  directly save the plot [boolean]
save_plot
save_param
                  list of saving parameters from all_plots_save_function
default_save_name
                  default save name for saving, don't change, change save_name in save_param
```

#### Value

plot

```
plotICG(CPGscores)
```

```
plotInteraction {\tt Changed Genes} \\ plotInteraction {\tt Changed Genes} \\
```

# Description

Create barplot to visualize interaction changed genes

## Usage

```
plotInteractionChangedGenes(
   gobject,
   cpgObject,
   source_type,
   source_markers,
   ICG_genes,
   cell_color_code = NULL,
   show_plot = NA,
   return_plot = NA,
   save_plot = NA,
   save_param = list(),
   default_save_name = "plotInteractionChangedGenes")
```

# Arguments

```
gobject
                  giotto object
cpgObject
                  cell proximity gene score object
                  cell type of the source cell
source_type
source_markers markers for the source cell type
ICG_genes
                  named character vector of ICG genes
cell_color_code
                  cell color code for the interacting cell types
show_plot
                  show plots
return_plot
                  return plotting object
save_plot
                  directly save the plot [boolean]
                  list of saving parameters from all_plots_save_function
save_param
default_save_name
                  default save name for saving, don't change, change save_name in save_param
```

## Value

plot

```
plotInteractionChangedGenes(CPGscores)
```

plotly\_axis\_scale\_2D 185

```
plotly_axis_scale_2D plotly_axis_scale_2D
```

# Description

adjust the axis scale in 3D plotly plot

# Usage

```
plotly_axis_scale_2D(
  cell_locations,
  sdimx = NULL,
  sdimy = NULL,
  mode = c("cube", "real", "custom"),
  custom_ratio = NULL
)
```

#### **Arguments**

```
cell_locations spatial_loc in giotto object sdimx x axis of cell spatial location sdimy y axis of cell spatial location mode axis adjustment mode custom_ratio set the ratio artificially
```

#### Value

edges in spatial grid as data.table()

## **Examples**

```
plotly_axis_scale_2D(gobject)
```

```
plotly_axis_scale_3D plotly_axis_scale_3D
```

# Description

adjust the axis scale in 3D plotly plot

```
plotly_axis_scale_3D(
  cell_locations,
  sdimx = NULL,
  sdimy = NULL,
  sdimz = NULL,
  mode = c("cube", "real", "custom"),
  custom_ratio = NULL
)
```

186 plotly\_grid

## **Arguments**

#### Value

edges in spatial grid as data.table()

# **Examples**

```
plotly_axis_scale_3D(gobject)
```

plotly\_grid

plotly\_grid

# Description

provide grid segment to draw in plot\_ly()

# Usage

```
plotly_grid(
   spatial_grid,
   x_start = "x_start",
   y_start = "y_start",
   x_end = "x_end",
   y_end = "y_end"
)
```

# Arguments

```
spatial_grid spatial_grid in giotto object
```

# Value

edges in spatial grid as data.table()

```
plotly_grid(gobject)
```

plotly\_network 187

plotly\_network

plotly\_network

## **Description**

provide network segment to draw in 3D plot\_ly()

## Usage

```
plotly_network(
  network,
  x = "sdimx_begin",
  y = "sdimy_begin",
  z = "sdimz_begin",
  x_end = "sdimx_end",
  y_end = "sdimy_end",
  z_end = "sdimz_end"
)
```

#### **Arguments**

gobject network in giotto object

#### Value

edges in network as data.table()

### **Examples**

```
plotly_network(gobject)
```

```
plotMetaDataCellsHeatmap
```

plotMetaDataCellsHeatmap

# Description

Creates heatmap for numeric cell metadata within aggregated clusters.

```
plotMetaDataCellsHeatmap(
  gobject,
  metadata_cols = NULL,
  spat_enr_names = NULL,
  value_cols = NULL,
  first_meta_col = NULL,
  second_meta_col = NULL,
  show_values = c("zscores", "original", "zscores_rescaled"),
  custom_cluster_order = NULL,
```

save\_plot

save\_param

default\_save\_name

custom\_gene\_order

```
clus_cor_method = "pearson",
      clus_cluster_method = "complete",
      custom_values_order = NULL,
      values_cor_method = "pearson",
      values_cluster_method = "complete",
      midpoint = 0,
      x_{text_size} = 8,
      x_{text_angle} = 45,
      y_text_size = 8,
      strip_text_size = 8,
      show_plot = NA,
      return_plot = NA,
      save_plot = NA,
      save_param = list(),
      default_save_name = "plotMetaDataCellsHeatmap"
    )
Arguments
    gobject
                     giotto object
                     annotation columns found in pDataDT(gobject)
    metadata_cols
    spat_enr_names spatial enrichment results to include
    value_cols
                     value columns to use
    first_meta_col if more than 1 metadata column, select the x-axis factor
    second_meta_col
                     if more than 1 metadata column, select the facetting factor
    show_values
                     which values to show on heatmap
    custom_cluster_order
                     custom cluster order (default = NULL)
    clus_cor_method
                     correlation method for clusters
    clus_cluster_method
                     hierarchical cluster method for the clusters
                     midpoint of show_values
    midpoint
                     size of x-axis text
    x_text_size
    x_text_angle
                     angle of x-axis text
    y_text_size
                     size of y-axis text
    strip_text_size
                     size of strip text
    show_plot
                     show plot
    return_plot
                     return ggplot object
```

directly save the plot [boolean]

custom gene order (default = NULL)

list of saving parameters, see showSaveParameters

default save name for saving, don't change, change save\_name in save\_param

plotMetaDataHeatmap 189

#### **Details**

Creates heatmap for the average values of selected value columns in the different annotation groups.

## Value

ggplot or data.table

#### See Also

plotMetaDataHeatmap for gene expression instead of numeric cell annotation data.

#### **Examples**

```
plotMetaDataCellsHeatmap(gobject)
```

```
plotMetaDataHeatmap
```

## **Description**

Creates heatmap for genes within aggregated clusters.

```
plotMetaDataHeatmap(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  metadata_cols = NULL,
  selected_genes = NULL,
  first_meta_col = NULL,
  second_meta_col = NULL,
  show_values = c("zscores", "original", "zscores_rescaled"),
  custom_cluster_order = NULL,
  clus_cor_method = "pearson",
  clus_cluster_method = "complete",
  custom_gene_order = NULL,
  gene_cor_method = "pearson",
  gene_cluster_method = "complete",
  gradient_color = c("blue", "white", "red"),
  gradient_midpoint = 0,
  gradient_limits = NULL,
  x_{text_size} = 10,
  x_{text_angle} = 45,
  y_{text_size} = 10,
  strip_text_size = 8,
  show_plot = NA,
```

```
return_plot = NA,
save_plot = NA,
save_param = list(),
default_save_name = "plotMetaDataHeatmap"
)
```

```
gobject
                 giotto object
expression_values
                 expression values to use
metadata_cols annotation columns found in pDataDT(gobject)
selected_genes subset of genes to use
first_meta_col if more than 1 metadata column, select the x-axis factor
second_meta_col
                 if more than 1 metadata column, select the facetting factor
show_values
                 which values to show on heatmap
custom_cluster_order
                 custom cluster order (default = NULL)
clus_cor_method
                 correlation method for clusters
clus_cluster_method
                 hierarchical cluster method for the clusters
custom_gene_order
                 custom gene order (default = NULL)
gene_cor_method
                 correlation method for genes
gene_cluster_method
                 hierarchical cluster method for the genes
gradient_color vector with 3 colors for numeric data
gradient_midpoint
                  midpoint for color gradient
gradient_limits
                 vector with lower and upper limits
                 size of x-axis text
x_text_size
x_text_angle
                 angle of x-axis text
y_text_size
                 size of y-axis text
strip_text_size
                 size of strip text
show_plot
                 show plot
return_plot
                 return ggplot object
save_plot
                 directly save the plot [boolean]
                 list of saving parameters, see showSaveParameters
save_param
default_save_name
                 default save name
```

plotPCA 191

#### **Details**

Creates heatmap for the average expression of selected genes in the different annotation/cluster groups. Calculation of cluster or gene order is done on the provided expression values, but visualization is by default on the z-scores. Other options are the original values or z-scores rescaled per gene (-1 to 1).

#### Value

ggplot or data.table

#### See Also

plotMetaDataCellsHeatmap for numeric cell annotation instead of gene expression.

### **Examples**

```
plotMetaDataHeatmap(gobject)
```

plotPCA

plotPCA

## **Description**

Short wrapper for PCA visualization

### Usage

```
plotPCA(gobject, dim_reduction_name = "pca", default_save_name = "PCA", ...)
```

```
gobject
                 giotto object
dim_reduction_name
                 name of PCA
default_save_name
                 default save name of PCA plot
                 Arguments passed on to dimPlot2D
                 group_by_subset subset the group_by factor column
                 dim1_to_use dimension to use on x-axis
                 dim2_to_use dimension to use on y-axis
                 spat_enr_names names of spatial enrichment results to include
                 show_NN_network show underlying NN network
                 nn_network_to_use type of NN network to use (kNN vs sNN)
                 network_name name of NN network to use, if show_NN_network = TRUE
                 cell_color color for cells (see details)
                 color_as_factor convert color column to factor
                 cell_color_code named vector with colors
                 cell_color_gradient vector with 3 colors for numeric data
                 gradient_midpoint midpoint for color gradient
```

192 plotPCA

```
gradient_limits vector with lower and upper limits
select_cell_groups select subset of cells/clusters based on cell color param-
select_cells select subset of cells based on cell IDs
show_other_cells display not selected cells
other_cell_color color of not selected cells
other_point_size size of not selected cells
show_cluster_center plot center of selected clusters
show_center_label plot label of selected clusters
center_point_size size of center points
label_size size of labels
label_fontface font of labels
edge_alpha column to use for alpha of the edges
point_shape point with border or not (border or no_border)
point_size size of point (cell)
point_alpha transparancy of point
point_border_col color of border around points
point_border_stroke stroke size of border around points
title title for plot, defaults to cell_color parameter
show_legend show legend
legend_text size of legend text
legend_symbol_size size of legend symbols
background_color color of plot background
axis_text size of axis text
axis_title size of axis title
cow_n_col cowplot param: how many columns
cow_rel_h cowplot param: relative height
cow_rel_w cowplot param: relative width
cow_align cowplot param: how to align
show_plot show plot
return_plot return ggplot object
save_plot directly save the plot [boolean]
save_param list of saving parameters, see showSaveParameters
```

#### Details

Description of parameters, see dimPlot2D. For 3D plots see plotPCA\_3D

#### Value

ggplot

#### See Also

```
Other reduced dimension visualizations: dimPlot2D(), dimPlot3D(), dimPlot(), plotPCA_2D(), plotPCA_3D(), plotTSNE_3D(), plotTSNE(), plotUMAP_2D(), plotUMAP_3D(), plotUMAP()
```

plotPCA\_2D 193

#### **Examples**

```
plotPCA(gobject)
```

plotPCA\_2D

plotPCA\_2D

# **Description**

Short wrapper for PCA visualization

## Usage

```
plotPCA_2D(
  gobject,
  dim_reduction_name = "pca",
  default_save_name = "PCA_2D",
   ...
)
```

```
gobject
                 giotto object
dim_reduction_name
                 name of PCA
default_save_name
                 default save name of PCA plot
                 Arguments passed on to dimPlot2D
. . .
                 group_by_subset subset the group_by factor column
                 dim1_to_use dimension to use on x-axis
                 dim2_to_use dimension to use on y-axis
                 spat_enr_names names of spatial enrichment results to include
                 show_NN_network show underlying NN network
                 nn_network_to_use type of NN network to use (kNN vs sNN)
                 network_name name of NN network to use, if show_NN_network = TRUE
                 cell_color color for cells (see details)
                 color_as_factor convert color column to factor
                 cell_color_code named vector with colors
                 cell_color_gradient vector with 3 colors for numeric data
                 gradient_midpoint midpoint for color gradient
                 gradient_limits vector with lower and upper limits
                 select_cell_groups select subset of cells/clusters based on cell_color param-
                 select_cells select subset of cells based on cell IDs
                 show_other_cells display not selected cells
                 other_cell_color color of not selected cells
                 other_point_size size of not selected cells
                 show_cluster_center plot center of selected clusters
```

194 plotPCA\_2D

```
show_center_label plot label of selected clusters
center_point_size size of center points
label_size size of labels
label_fontface font of labels
edge_alpha column to use for alpha of the edges
point_shape point with border or not (border or no_border)
point_size size of point (cell)
point_alpha transparancy of point
point_border_col color of border around points
point_border_stroke stroke size of border around points
title title for plot, defaults to cell_color parameter
show_legend show legend
legend_text size of legend text
legend_symbol_size size of legend symbols
background_color color of plot background
axis_text size of axis text
axis_title size of axis title
cow_n_col cowplot param: how many columns
cow_rel_h cowplot param: relative height
cow_rel_w cowplot param: relative width
cow_align cowplot param: how to align
show_plot show plot
return_plot return ggplot object
save_plot directly save the plot [boolean]
save_param list of saving parameters, see showSaveParameters
```

#### **Details**

Description of parameters, see dimPlot2D. For 3D plots see plotPCA\_3D

# Value

ggplot

#### See Also

```
Other reduced dimension visualizations: dimPlot2D(), dimPlot3D(), dimPlot(), plotPCA_3D(), plotPCA(), plotTSNE_2D(), plotTSNE_3D(), plotTSNE(), plotUMAP_2D(), plotUMAP_3D(), plotUMAP()
```

```
plotPCA_2D(gobject)
```

plotPCA\_3D 195

plotPCA\_3D

#### **Description**

Visualize cells according to 3D PCA dimension reduction

plotPCA\_3D

#### Usage

```
plotPCA_3D(
  gobject,
  dim_reduction_name = "pca",
  default_save_name = "PCA_3D",
   ...
)
```

```
gobject
                 giotto object
dim_reduction_name
                 name of PCA
default_save_name
                 default save name of PCA plot
                 Arguments passed on to dimPlot3D
                 dim1_to_use dimension to use on x-axis
                 dim2_to_use dimension to use on y-axis
                 dim3_to_use dimension to use on z-axis
                 show_NN_network show underlying NN network
                 nn_network_to_use type of NN network to use (kNN vs sNN)
                 network_name name of NN network to use, if show NN network = TRUE
                 cell_color color for cells (see details)
                 color_as_factor convert color column to factor
                 cell_color_code named vector with colors
                 select_cell_groups select subset of cells/clusters based on cell_color param-
                      eter
                 select_cells select subset of cells based on cell IDs
                 show_other_cells display not selected cells
                 other_cell_color color of not selected cells
                 other_point_size size of not selected cells
                 show_cluster_center plot center of selected clusters
                 show_center_label plot label of selected clusters
                 center_point_size size of center points
                 label_size size of labels
                 edge_alpha column to use for alpha of the edges
                 point_size size of point (cell)
                 show_plot show plot
                 return_plot return ggplot object
                  save_plot directly save the plot [boolean]
                 save_param list of saving parameters, see showSaveParameters
```

#### **Details**

Description of parameters.

#### Value

plotly

#### See Also

```
Other reduced dimension visualizations: dimPlot2D(), dimPlot3D(), dimPlot(), plotPCA_2D(), plotPCA_2D(), plotTSNE_2D(), plotTSNE_3D(), plotTSNE(), plotUMAP_2D(), plotUMAP_3D(), plotUMAP()
```

## **Examples**

```
plotPCA_3D(gobject)
```

plotRankSpatvsExpr

plotRankSpatvsExpr

#### **Description**

Plots dotplot to compare ligand-receptor rankings from spatial and expression information

## Usage

```
plotRankSpatvsExpr(
  gobject,
  combCC,
  expr_rnk_column = "LR_expr_rnk",
  spat_rnk_column = "LR_spat_rnk",
  midpoint = 10,
  size\_range = c(0.01, 1.5),
  xlims = NULL,
  ylims = NULL,
  selected_ranks = c(1, 10, 20),
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "plotRankSpatvsExpr"
)
```

plotRecovery 197

```
midpoint of colors
midpoint
size_range
                  size ranges of dotplot
                  x-limits, numerical vector of 2
xlims
ylims
                  y-limits, numerical vector of 2
selected_ranks numerical vector, will be used to print out the percentage of top spatial ranks are
                  recovered
show_plot
                  show plots
return_plot
                  return plotting object
save_plot
                  directly save the plot [boolean]
                  list of saving parameters from all_plots_save_function
save_param
default_save_name
                  default save name for saving, don't change, change save_name in save_param
```

#### Value

ggplot

# **Examples**

```
plotRankSpatvsExpr(CPGscores)
```

plotRecovery

plotRecovery

# Description

Plots recovery plot to compare ligand-receptor rankings from spatial and expression information

```
plotRecovery(
  gobject,
  combCC,
  expr_rnk_column = "exprPI_rnk",
  spat_rnk_column = "spatPI_rnk",
  ground_truth = c("spatial", "expression"),
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "plotRecovery"
)
```

198 plotRecovery\_sub

### **Arguments**

gobject giotto object

combCC combined communinication scores from combCCcom

expr\_rnk\_column

column with expression rank information to use

spat\_rnk\_column

column with spatial rank information to use

ground\_truth what to consider as ground truth (default: spatial)

show\_plot show plots

return\_plot return plotting object

save\_plot directly save the plot [boolean]

save\_param list of saving parameters from all\_plots\_save\_function

default\_save\_name

default save name for saving, don't change, change save\_name in save\_param

#### Value

ggplot

# **Examples**

plotRecovery(CPGscores)

plotRecovery\_sub plotRecovery\_sub

# Description

Plots recovery plot to compare ligand-receptor rankings from spatial and expression information

# Usage

```
plotRecovery_sub(combCC, first_col = "LR_expr_rnk", second_col = "LR_spat_rnk")
```

# Arguments

combCC combined communinication scores from combCCcom

first\_col first column to use second\_col second column to use

```
plotRecovery_sub(CPGscores)
```

```
plotStatDelaunayNetwork
```

plotStatDelaunayNetwork

# Description

Plots network statistics for a Delaunay network..

# Usage

```
plotStatDelaunayNetwork(
  gobject,
  method = c("deldir", "delaunayn_geometry", "RTriangle"),
  dimensions = "all",
  maximum_distance = "auto",
  minimum_k = 0,
  options = "Pp",
  Y = TRUE,
  j = TRUE,
  S = 0,
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "plotStatDelaunayNetwork",
)
```

gobject	giotto object
dimensions	which spatial dimensions to use (maximum 2 dimensions)
maximum_distanc	ne e
	distance cuttof for Delaunay neighbors to consider
minimum_k	minimum neigbhours if maximum_distance != NULL
options	(geometry) String containing extra control options for the underlying Qhull command; see the Qhull documentation (/doc/qhull/html/qdelaun.html) for the available options. (default = 'Pp', do not report precision problems)
Υ	(RTriangle) If TRUE prohibits the insertion of Steiner points on the mesh boundary.
j	(RTriangle) If TRUE jettisons vertices that are not part of the final triangulation from the output.
S	(RTriangle) Specifies the maximum number of added Steiner points.
show_plot	show plots
return_plot	return ggplot object
save_plot	directly save the plot [boolean]
save_param	list of saving parameters from all_plots_save_function

200 plotTSNE

```
default_save_name

default save name for saving, don't change, change save_name in save_param

Other parameters of the triangulate function

name

name for spatial network (default = 'delaunay_network')
```

#### **Details**

Plots statistics for a spatial Delaunay network as explained in triangulate. This can be used to further finetune the createSpatialDelaunayNetwork function.

#### Value

giotto object with updated spatial network slot

## **Examples**

```
plotStatDelaunayNetwork(gobject)
```

plotTSNE

*plotTSNE* 

#### **Description**

Short wrapper for tSNE visualization

# Usage

```
plotTSNE(gobject, dim_reduction_name = "tsne", default_save_name = "tSNE", ...)
```

```
gobject
                 giotto object
dim_reduction_name
                 name of TSNE
default_save_name
                 default save name of TSNE plot
                 Arguments passed on to dimPlot2D
                 group_by_subset subset the group_by factor column
                 dim1_to_use dimension to use on x-axis
                 dim2_to_use dimension to use on y-axis
                 spat_enr_names names of spatial enrichment results to include
                 show_NN_network show underlying NN network
                 nn_network_to_use type of NN network to use (kNN vs sNN)
                 network_name name of NN network to use, if show_NN_network = TRUE
                 cell_color color for cells (see details)
                 color_as_factor convert color column to factor
                 cell_color_code named vector with colors
                 cell_color_gradient vector with 3 colors for numeric data
                 gradient_midpoint midpoint for color gradient
```

plotTSNE 201

```
gradient_limits vector with lower and upper limits
select_cell_groups select subset of cells/clusters based on cell color param-
select_cells select subset of cells based on cell IDs
show_other_cells display not selected cells
other_cell_color color of not selected cells
other_point_size size of not selected cells
show_cluster_center plot center of selected clusters
show_center_label plot label of selected clusters
center_point_size size of center points
label_size size of labels
label_fontface font of labels
edge_alpha column to use for alpha of the edges
point_shape point with border or not (border or no_border)
point_size size of point (cell)
point_alpha transparancy of point
point_border_col color of border around points
point_border_stroke stroke size of border around points
title title for plot, defaults to cell_color parameter
show_legend show legend
legend_text size of legend text
legend_symbol_size size of legend symbols
background_color color of plot background
axis_text size of axis text
axis_title size of axis title
cow_n_col cowplot param: how many columns
cow_rel_h cowplot param: relative height
cow_rel_w cowplot param: relative width
cow_align cowplot param: how to align
show_plot show plot
return_plot return ggplot object
save_plot directly save the plot [boolean]
save_param list of saving parameters, see showSaveParameters
```

#### Details

Description of parameters, see dimPlot2D. For 3D plots see plotTSNE\_3D

#### Value

ggplot

#### See Also

```
Other reduced dimension visualizations: dimPlot2D(), dimPlot3D(), dimPlot(), plotPCA_2D(), plotPCA_3D(), plotPCA_3D(), plotTSNE_2D(), plotTSNE_3D(), plotUMAP_2D(), plotUMAP_3D(), plotUMAP()
```

202 plotTSNE\_2D

#### **Examples**

```
plotTSNE(gobject)
```

plotTSNE\_2D

plotTSNE\_2D

# **Description**

Short wrapper for tSNE visualization

# Usage

```
plotTSNE_2D(
  gobject,
  dim_reduction_name = "tsne",
  default_save_name = "tSNE_2D",
  ...
)
```

```
gobject
                 giotto object
dim_reduction_name
                 name of TSNE
default_save_name
                 default save name of TSNE plot
                 Arguments passed on to dimPlot2D
. . .
                 group_by_subset subset the group_by factor column
                 dim1_to_use dimension to use on x-axis
                 dim2_to_use dimension to use on y-axis
                 spat_enr_names names of spatial enrichment results to include
                 show_NN_network show underlying NN network
                 nn_network_to_use type of NN network to use (kNN vs sNN)
                 network_name name of NN network to use, if show_NN_network = TRUE
                 cell_color color for cells (see details)
                 color_as_factor convert color column to factor
                 cell_color_code named vector with colors
                 cell_color_gradient vector with 3 colors for numeric data
                 gradient_midpoint midpoint for color gradient
                 gradient_limits vector with lower and upper limits
                 select_cell_groups select subset of cells/clusters based on cell_color param-
                 select_cells select subset of cells based on cell IDs
                 show_other_cells display not selected cells
                 other_cell_color color of not selected cells
                 other_point_size size of not selected cells
                 show_cluster_center plot center of selected clusters
```

plotTSNE\_2D 203

```
show_center_label plot label of selected clusters
center_point_size size of center points
label_size size of labels
label_fontface font of labels
edge_alpha column to use for alpha of the edges
point_shape point with border or not (border or no_border)
point_size size of point (cell)
point_alpha transparancy of point
point_border_col color of border around points
point_border_stroke stroke size of border around points
title title for plot, defaults to cell_color parameter
show_legend show legend
legend_text size of legend text
legend_symbol_size size of legend symbols
background_color color of plot background
axis_text size of axis text
axis_title size of axis title
cow_n_col cowplot param: how many columns
cow_rel_h cowplot param: relative height
cow_rel_w cowplot param: relative width
cow_align cowplot param: how to align
show_plot show plot
return_plot return ggplot object
save_plot directly save the plot [boolean]
save_param list of saving parameters, see showSaveParameters
```

#### **Details**

Description of parameters, see dimPlot2D. For 3D plots see plotTSNE\_3D

# Value

ggplot

#### See Also

```
Other reduced dimension visualizations: dimPlot2D(), dimPlot3D(), dimPlot(), plotPCA_2D(), plotPCA_3D(), plotPCA(), plotTSNE_3D(), plotTSNE(), plotUMAP_2D(), plotUMAP_3D(), plotUMAP()
```

```
plotTSNE_2D(gobject)
```

204 plotTSNE\_3D

plotTSNE\_3D

plotTSNE 3D

#### **Description**

Visualize cells according to dimension reduction coordinates

#### Usage

```
plotTSNE_3D(
  gobject,
  dim_reduction_name = "tsne"
  default_save_name = "TSNE_3D",
)
```

```
gobject
                 giotto object
dim_reduction_name
                 name of TSNE
default_save_name
                 default save name of TSNE plot
                 Arguments passed on to dimPlot3D
                 dim1_to_use dimension to use on x-axis
                 dim2_to_use dimension to use on y-axis
                 dim3_to_use dimension to use on z-axis
                 show_NN_network show underlying NN network
                 nn_network_to_use type of NN network to use (kNN vs sNN)
                 network_name name of NN network to use, if show NN network = TRUE
                 cell_color color for cells (see details)
                 color_as_factor convert color column to factor
                 cell_color_code named vector with colors
                 select_cell_groups select subset of cells/clusters based on cell_color param-
                      eter
                 select_cells select subset of cells based on cell IDs
                 show_other_cells display not selected cells
                 other_cell_color color of not selected cells
                 other_point_size size of not selected cells
                 show_cluster_center plot center of selected clusters
                 show_center_label plot label of selected clusters
                 center_point_size size of center points
                 label_size size of labels
                 edge_alpha column to use for alpha of the edges
                 point_size size of point (cell)
                 show_plot show plot
                 return_plot return ggplot object
                  save_plot directly save the plot [boolean]
                  save_param list of saving parameters, see showSaveParameters
```

plotUMAP 205

#### **Details**

Description of parameters.

#### Value

plotly

#### See Also

```
Other reduced dimension visualizations: dimPlot2D(), dimPlot3D(), dimPlot(), plotPCA_2D(), plotPCA_3D(), plotPCA(), plotTSNE_2D(), plotTSNE(), plotUMAP_2D(), plotUMAP_3D(), plotUMAP()
```

## **Examples**

```
plotTSNE_3D(gobject)
```

plotUMAP

plotUMAP

## **Description**

Short wrapper for UMAP visualization

## Usage

```
plotUMAP(gobject, dim_reduction_name = "umap", default_save_name = "UMAP", ...)
```

```
gobject
                 giotto object
dim_reduction_name
                 name of UMAP
default_save_name
                 default save name of UMAP plot
                 Arguments passed on to dimPlot2D
. . .
                 group_by_subset subset the group_by factor column
                 dim1_to_use dimension to use on x-axis
                 dim2_to_use dimension to use on y-axis
                 spat_enr_names names of spatial enrichment results to include
                 show_NN_network show underlying NN network
                 nn_network_to_use type of NN network to use (kNN vs sNN)
                 network_name name of NN network to use, if show_NN_network = TRUE
                 cell_color color for cells (see details)
                 color_as_factor convert color column to factor
                 cell_color_code named vector with colors
                 cell_color_gradient vector with 3 colors for numeric data
                 gradient_midpoint midpoint for color gradient
                 gradient_limits vector with lower and upper limits
```

206 plotUMAP

```
select_cell_groups select subset of cells/clusters based on cell color param-
select_cells select subset of cells based on cell IDs
show_other_cells display not selected cells
other_cell_color color of not selected cells
other_point_size size of not selected cells
show_cluster_center plot center of selected clusters
show_center_label plot label of selected clusters
center_point_size size of center points
label_size size of labels
label_fontface font of labels
edge_alpha column to use for alpha of the edges
point_shape point with border or not (border or no_border)
point_size size of point (cell)
point_alpha transparancy of point
point_border_col color of border around points
point_border_stroke stroke size of border around points
title title for plot, defaults to cell_color parameter
show_legend show legend
legend_text size of legend text
legend_symbol_size size of legend symbols
background_color color of plot background
axis_text size of axis text
axis_title size of axis title
cow_n_col cowplot param: how many columns
cow_rel_h cowplot param: relative height
cow_rel_w cowplot param: relative width
cow_align cowplot param: how to align
show_plot show plot
return_plot return ggplot object
save_plot directly save the plot [boolean]
save_param list of saving parameters, see showSaveParameters
```

# **Details**

Description of parameters, see dimPlot2D. For 3D plots see plotUMAP\_3D

## Value

ggplot

#### See Also

```
Other reduced dimension visualizations: dimPlot2D(), dimPlot3D(), dimPlot(), plotPCA_2D(), plotPCA_3D(), plotPCA(), plotTSNE_2D(), plotTSNE_3D(), plotTSNE(), plotUMAP_2D(), plotUMAP_3D()
```

```
plotUMAP(gobject)
```

plotUMAP\_2D 207

plotUMAP\_2D

plotUMAP\_2D

#### **Description**

Short wrapper for UMAP visualization

# Usage

```
plotUMAP_2D(
  gobject,
  dim_reduction_name = "umap";
  default_save_name = "UMAP_2D",
)
```

```
giotto object
gobject
dim_reduction_name
                 name of UMAP
default_save_name
                 default save name of UMAP plot
                 Arguments passed on to dimPlot2D
                 group_by_subset subset the group by factor column
                 dim1_to_use dimension to use on x-axis
                 dim2_to_use dimension to use on y-axis
                 spat_enr_names names of spatial enrichment results to include
                 show_NN_network show underlying NN network
                 nn_network_to_use type of NN network to use (kNN vs sNN)
                 network_name name of NN network to use, if show_NN_network = TRUE
                 cell_color color for cells (see details)
                 color_as_factor convert color column to factor
                 cell_color_code named vector with colors
                 cell_color_gradient vector with 3 colors for numeric data
                 gradient_midpoint midpoint for color gradient
                 gradient_limits vector with lower and upper limits
                 select_cell_groups select subset of cells/clusters based on cell_color param-
                 select_cells select subset of cells based on cell IDs
                 show_other_cells display not selected cells
                 other_cell_color color of not selected cells
                 other_point_size size of not selected cells
                  show_cluster_center plot center of selected clusters
                 show_center_label plot label of selected clusters
                 center_point_size size of center points
                 label_size size of labels
```

208 plotUMAP\_2D

```
label_fontface font of labels
edge_alpha column to use for alpha of the edges
point_shape point with border or not (border or no border)
point_size size of point (cell)
point_alpha transparancy of point
point_border_col color of border around points
point_border_stroke stroke size of border around points
title title for plot, defaults to cell_color parameter
show_legend show legend
legend_text size of legend text
legend_symbol_size size of legend symbols
background_color color of plot background
axis_text size of axis text
axis_title size of axis title
cow_n_col cowplot param: how many columns
cow_rel_h cowplot param: relative height
cow_rel_w cowplot param: relative width
cow_align cowplot param: how to align
show_plot show plot
return_plot return ggplot object
save_plot directly save the plot [boolean]
save_param list of saving parameters, see showSaveParameters
```

#### **Details**

Description of parameters, see dimPlot2D. For 3D plots see plotUMAP\_3D

#### Value

ggplot

#### See Also

```
Other reduced dimension visualizations: dimPlot2D(), dimPlot3D(), dimPlot(), plotPCA_2D(), plotPCA_3D(), plotPCA(), plotTSNE_2D(), plotTSNE_3D(), plotTSNE(), plotUMAP_3D(), plotUMAP()
```

```
plotUMAP_2D(gobject)
```

plotUMAP\_3D 209

plotUMAP\_3D

plotUMAP\_3D

#### **Description**

Visualize cells according to dimension reduction coordinates

#### Usage

```
plotUMAP_3D(
  gobject,
  dim_reduction_name = "umap";
  default_save_name = "UMAP_3D",
)
```

```
gobject
                 giotto object
dim_reduction_name
                 name of UMAP
default_save_name
                 default save name of UMAP plot
                 Arguments passed on to dimPlot3D
                 dim1_to_use dimension to use on x-axis
                 dim2_to_use dimension to use on y-axis
                 dim3_to_use dimension to use on z-axis
                 show_NN_network show underlying NN network
                 nn_network_to_use type of NN network to use (kNN vs sNN)
                 network_name name of NN network to use, if show NN network = TRUE
                 cell_color color for cells (see details)
                 color_as_factor convert color column to factor
                 cell_color_code named vector with colors
                 select_cell_groups select subset of cells/clusters based on cell_color param-
                      eter
                 select_cells select subset of cells based on cell IDs
                 show_other_cells display not selected cells
                 other_cell_color color of not selected cells
                 other_point_size size of not selected cells
                 show_cluster_center plot center of selected clusters
                 show_center_label plot label of selected clusters
                 center_point_size size of center points
                 label_size size of labels
                 edge_alpha column to use for alpha of the edges
                 point_size size of point (cell)
                 show_plot show plot
                 return_plot return ggplot object
                  save_plot directly save the plot [boolean]
                  save_param list of saving parameters, see showSaveParameters
```

210 rankEnrich

#### **Details**

Description of parameters.

#### Value

plotly

#### See Also

```
Other reduced dimension visualizations: dimPlot2D(), dimPlot3D(), dimPlot(), plotPCA_2D(), plotPCA_3D(), plotPCA(), plotTSNE_2D(), plotTSNE_3D(), plotTSNE(), plotUMAP_2D(), plotUMAP()
```

#### **Examples**

```
plotUMAP_3D(gobject)
```

print.giotto

print method for giotto class

## **Description**

print method for giotto class. Prints the chosen number of genes (rows) and cells (columns) from the raw count matrix. Also print the spatial locations for the chosen number of cells.

## Usage

```
print.giotto(object, ...)
```

# Arguments

nr\_genes number of genes (rows) to print nr\_cells number of cells (columns) to print

rankEnrich

rankEnrich

## **Description**

Function to calculate gene signature enrichment scores per spatial position using a rank based approach.

```
rankEnrich(
  gobject,
  sign_matrix,
  expression_values = c("normalized", "scaled", "custom"),
  reverse_log_scale = TRUE,
  logbase = 2,
  output_enrichment = c("original", "zscore")
)
```

rankSpatialCorGroups 211

### **Arguments**

```
gobject Giotto object

sign_matrix Matrix of signature genes for each cell type / process

expression_values

expression values to use

reverse_log_scale

reverse expression values from log scale

logbase log base to use if reverse_log_scale = TRUE

output_enrichment

how to return enrichment output
```

# now to return entremnent outpo

#### **Details**

sign\_matrix: a rank-fold matrix with genes as row names and cell-types as column names. Alternatively a scRNA-seq matrix and vector with clusters can be provided to makeSignMatrixRank, which will create the matrix for you.

First a new rank is calculated as  $R = (R1*R2)^{\Lambda}(1/2)$ , where R1 is the rank of fold-change for each gene in each spot and R2 is the rank of each marker in each cell type. The Rank-Biased Precision is then calculated as:  $RBP = (1 - 0.99) * (0.99)^{\Lambda}(R - 1)$  and the final enrichment score is then calculated as the sum of top 100 RBPs.

#### Value

data.table with enrichment results

# See Also

```
make Sign Matrix Rank
```

## **Examples**

```
rankEnrich(gobject)
```

```
rankSpatialCorGroups rankSpatialCorGroups
```

# **Description**

Rank spatial correlated clusters according to correlation structure

```
rankSpatialCorGroups(
  gobject,
  spatCorObject,
  use_clus_name = NULL,
  show_plot = NA,
  return_plot = FALSE,
  save_plot = NA,
```

212 readExprMatrix

```
save_param = list(),
  default_save_name = "rankSpatialCorGroups"
)
```

# **Arguments**

gobject giotto object

spatCorObject spatial correlation object

 $use\_clus\_name \quad name \ of \ clusters \ to \ visualize \ (from \ clusterSpatialCorGenes())$ 

show\_plot show plot

return\_plot return ggplot object

save\_plot directly save the plot [boolean]

save\_param list of saving parameters from all\_plots\_save\_function

default\_save\_name

default save name for saving, don't change, change save\_name in save\_param

## Value

data.table with positive (within group) and negative (outside group) scores

## **Examples**

```
rankSpatialCorGroups(gobject)
```

rank\_binarize

rank\_binarize

# Description

create binarized scores from a vector using arbitrary rank

# Usage

```
rank_binarize(x, max_rank = 200)
```

 ${\tt readExprMatrix}$ 

readExprMatrix

# Description

Function to read an expression matrix into a sparse matrix.

```
readExprMatrix(path, cores = NA, transpose = FALSE)
```

readGiottoInstructions 213

## **Arguments**

path path to the expression matrix

cores number of cores to use

transpose transpose matrix

#### **Details**

The expression matrix needs to have both unique column names and row names

# Value

sparse matrix

# **Examples**

readExprMatrix()

readGiottoInstructions

read Giot to Instrunctions

# Description

Retrieves the instruction associated with the provided parameter

## Usage

```
readGiottoInstructions(giotto_instructions, param = NULL)
```

# **Arguments**

giotto\_instructions

 $giot to\ object\ or\ result\ from\ create Giot to Instructions()$ 

param parameter to retrieve

#### Value

specific parameter

# Examples

readGiottoInstrunctions()

214 removeGeneAnnotation

removeCellAnnotation removeCellAnnotation

# Description

removes cell annotation of giotto object

## Usage

```
removeCellAnnotation(gobject, columns = NULL, return_gobject = TRUE)
```

#### **Arguments**

gobject giotto object

columns names of columns to remove

return\_gobject boolean: return giotto object (default = TRUE)

#### **Details**

if return\_gobject = FALSE, it will return the cell metadata

## Value

giotto object

# **Examples**

removeCellAnnotation(gobject)

removeGeneAnnotation removeGeneAnnotation

# **Description**

removes gene annotation of giotto object

# Usage

```
removeGeneAnnotation(gobject, columns = NULL, return_gobject = TRUE)
```

## **Arguments**

gobject giotto object

columns names of columns to remove

return\_gobject boolean: return giotto object (default = TRUE)

## **Details**

if return\_gobject = FALSE, it will return the gene metadata

## Value

giotto object

# **Examples**

removeGeneAnnotation(gobject)

replaceGiottoInstructions

replace Giot to Instructions

# Description

Function to replace all instructions from giotto object

# Usage

```
replaceGiottoInstructions(gobject, instructions = NULL)
```

# Arguments

gobject giotto object

instructions new instructions (e.g. result from createGiottoInstructions)

# Value

giotto object with replaces instructions

# Examples

replaceGiottoInstructions()

runPCA runPCA

# Description

runs a Principal Component Analysis

216 runPCA

#### Usage

```
runPCA(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  reduction = c("cells", "genes"),
  name = "pca",
  genes_to_use = "hvg",
  return_gobject = TRUE,
  center = F,
  scale_unit = F,
  ncp = 100,
  method = c("irlba", "factominer"),
  rev = FALSE,
  verbose = TRUE,
  ...
)
```

# Arguments

gobject giotto object expression\_values expression values to use reduction cells or genes arbitrary name for PCA run name subset of genes to use for PCA genes\_to\_use return\_gobject boolean: return giotto object (default = TRUE) center data first (default = FALSE) center scale features before PCA (default = FALSE) scale\_unit ncp number of principal components to calculate which implementation to use method do a reverse PCA rev verbose verbosity of the function additional parameters for PCA (see details)

## Details

See prcomp\_irlba and PCA for more information about other parameters.

- genes\_to\_use = NULL: will use all genes from the selected matrix
- genes\_to\_use = <hvg name>: can be used to select a column name of highly variable genes, created by (see calculateHVG)
- genes\_to\_use = c('geneA', 'geneB', ...): will use all manually provided genes

# Value

giotto object with updated PCA dimension recuction

runPCA\_factominer 217

#### **Examples**

```
# 1. create giotto object
expr_path = system.file("extdata", "seqfish_field_expr.txt", package = 'Giotto')
loc_path = system.file("extdata", "seqfish_field_locs.txt", package = 'Giotto')
VC_small <- createGiottoObject(raw_exprs = expr_path, spatial_locs = loc_path)</pre>
# 2. normalize giotto
VC_small <- normalizeGiotto(gobject = VC_small, scalefactor = 6000)</pre>
VC_small <- addStatistics(gobject = VC_small)</pre>
# 3. dimension reduction
VC_small <- calculateHVG(gobject = VC_small)</pre>
VC_small <- runPCA(gobject = VC_small)</pre>
plotPCA(VC_small)
```

runPCA\_factominer

runPCA\_factominer

## **Description**

performs PCA based on the factominer package

## Usage

```
runPCA_factominer(x, ncp = 100, scale = TRUE, rev = FALSE, ...)
```

# **Arguments**

matrix or object that can be converted to matrix number of principal components to calculate ncp scale features scale

reverse PCA rev

## Value

list of eigenvalues, loadings and pca coordinates

```
runPCA_prcomp_irlba
                       runPCA_prcomp_irlba
```

# **Description**

performs PCA based on the irlba package

218 runtSNE

#### Usage

```
runPCA_prcomp_irlba(
    x,
    ncp = 100,
    center = TRUE,
    scale = TRUE,
    rev = FALSE,
    ...
)
```

# **Arguments**

x matrix or object that can be converted to matrix
ncp number of principal components to calculate
center center data
scale scale features
rev reverse PCA

#### Value

list of eigenvalues, loadings and pca coordinates

runtSNE runtSNE

# Description

run tSNE

# Usage

```
runtSNE(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  reduction = c("cells", "genes"),
  dim_reduction_to_use = "pca",
  dim_reduction_name = "pca",
  dimensions_to_use = 1:10,
  name = "tsne",
  genes_to_use = NULL,
  return_gobject = TRUE,
  dims = 2,
  perplexity = 30,
  theta = 0.5,
  do_PCA_first = F,
  set_seed = T,
  seed_number = 1234,
  verbose = TRUE,
)
```

runtSNE 219

#### **Arguments**

```
gobject
                 giotto object
expression_values
                 expression values to use
reduction
                 cells or genes
dim_reduction_to_use
                 use another dimension reduction set as input
dim_reduction_name
                 name of dimension reduction set to use
dimensions_to_use
                 number of dimensions to use as input
name
                 arbitrary name for tSNE run
                 if dim_reduction_to_use = NULL, which genes to use
genes_to_use
return_gobject boolean: return giotto object (default = TRUE)
                 tSNE param: number of dimensions to return
dims
perplexity
                 tSNE param: perplexity
                 tSNE param: theta
theta
                 tSNE param: do PCA before tSNE (default = FALSE)
do_PCA_first
set seed
                 use of seed
seed_number
                 seed number to use
                 verbosity of the function
verbose
                 additional tSNE parameters
. . .
```

#### **Details**

See Rtsne for more information about these and other parameters.

- Input for tSNE dimension reduction can be another dimension reduction (default = 'pca')
- To use gene expression as input set dim\_reduction\_to\_use = NULL
- If dim\_reduction\_to\_use = NULL, genes\_to\_use can be used to select a column name of highly variable genes (see calculateHVG) or simply provide a vector of genes
- multiple tSNE results can be stored by changing the *name* of the analysis

#### Value

giotto object with updated tSNE dimension recuction

#### **Examples**

```
# 1. create giotto object
expr_path = system.file("extdata", "seqfish_field_expr.txt", package = 'Giotto')
loc_path = system.file("extdata", "seqfish_field_locs.txt", package = 'Giotto')
VC_small <- createGiottoObject(raw_exprs = expr_path, spatial_locs = loc_path)
# 2. normalize giotto
VC_small <- normalizeGiotto(gobject = VC_small, scalefactor = 6000)
VC_small <- addStatistics(gobject = VC_small)</pre>
```

220 runUMAP

```
# 3. dimension reduction
VC_small <- calculateHVG(gobject = VC_small)
VC_small <- runPCA(gobject = VC_small)
VC_small <- runTSNE(VC_small, dimensions_to_use = 1:5, n_threads = 2)
plotTSNE(gobject = VC_small)</pre>
```

runUMAP

runUMAP

## **Description**

run UMAP

## Usage

```
runUMAP(
 gobject,
 expression_values = c("normalized", "scaled", "custom"),
 reduction = c("cells", "genes"),
 dim_reduction_to_use = "pca",
 dim_reduction_name = "pca",
 dimensions_to_use = 1:10,
 name = "umap",
 genes_to_use = NULL,
 return_gobject = TRUE,
 n_neighbors = 40,
 n_{components} = 2,
 n_{epochs} = 400,
 min_dist = 0.01,
 n_{threads} = 1,
 spread = 5,
 set\_seed = T,
 seed_number = 1234,
 verbose = T,
)
```

# Arguments

```
gobject giotto object
expression_values
expression values to use

reduction cells or genes
dim_reduction_to_use
use another dimension reduction set as input
dim_reduction_name
name of dimension reduction set to use
dimensions_to_use
number of dimensions to use as input
```

runUMAP 221

```
name
                 arbitrary name for UMAP run
                 if dim_reduction_to_use = NULL, which genes to use
genes_to_use
return_gobject boolean: return giotto object (default = TRUE)
n_neighbors
                 UMAP param: number of neighbors
n_components
                 UMAP param: number of components
                 UMAP param: number of epochs
n_epochs
                 UMAP param: minimum distance
min_dist
n_threads
                 UMAP param: threads to use
spread
                 UMAP param: spread
set_seed
                 use of seed
                 seed number to use
seed_number
                 verbosity of function
verbose
                 additional UMAP parameters
```

#### **Details**

See umap for more information about these and other parameters.

- Input for UMAP dimension reduction can be another dimension reduction (default = 'pca')
- To use gene expression as input set dim\_reduction\_to\_use = NULL
- If dim\_reduction\_to\_use = NULL, genes\_to\_use can be used to select a column name of highly variable genes (see calculateHVG) or simply provide a vector of genes
- multiple UMAP results can be stored by changing the *name* of the analysis

#### Value

giotto object with updated UMAP dimension recuction

## **Examples**

```
# 1. create giotto object
expr_path = system.file("extdata", "seqfish_field_expr.txt", package = 'Giotto')
loc_path = system.file("extdata", "seqfish_field_locs.txt", package = 'Giotto')
VC_small <- createGiottoObject(raw_exprs = expr_path, spatial_locs = loc_path)
# 2. normalize giotto
VC_small <- normalizeGiotto(gobject = VC_small, scalefactor = 6000)
VC_small <- addStatistics(gobject = VC_small)
# 3. dimension reduction
VC_small <- calculateHVG(gobject = VC_small)
VC_small <- runPCA(gobject = VC_small)
VC_small <- runUMAP(VC_small, dimensions_to_use = 1:5, n_threads = 2)
plotUMAP(gobject = VC_small)</pre>
```

222 screePlot

screePlot screePlot

## **Description**

identify significant prinicipal components (PCs) using an screeplot (a.k.a. elbowplot)

## Usage

```
screePlot(
 gobject,
 name = "pca",
 expression_values = c("normalized", "scaled", "custom"),
 reduction = c("cells", "genes"),
 method = c("irlba", "factominer"),
 rev = FALSE,
  genes_to_use = NULL,
  center = F,
  scale_unit = F,
 ncp = 100,
 ylim = c(0, 20),
  verbose = T,
  show_plot = NA,
 return_plot = NA,
  save_plot = NA,
  save_param = list(),
 default_save_name = "screePlot",
)
```

# **Arguments**

gobject giotto object name of PCA object if available name expression\_values expression values to use cells or genes reduction which implementation to use method do a reverse PCA rev genes\_to\_use subset of genes to use for PCA center data before PCA center scale features before PCA scale\_unit number of principal components to calculate ncp ylim y-axis limits on scree plot verbose verobsity show\_plot show plot return\_plot return ggplot object

selectPatternGenes 223

#### Details

Screeplot works by plotting the explained variance of each individual PC in a barplot allowing you to identify which PC provides a significant contribution (a.k.a 'elbow method'). Screeplot will use an available pca object, based on the parameter 'name', or it will create it if it's not available (see runPCA)

#### Value

ggplot object for scree method

## **Examples**

```
screePlot(gobject)
```

selectPatternGenes

selectPatternGenes

## **Description**

Select genes correlated with spatial patterns

## Usage

```
selectPatternGenes(
   spatPatObj,
   dimensions = 1:5,
   top_pos_genes = 10,
   top_neg_genes = 10,
   min_pos_cor = 0.5,
   min_neg_cor = -0.5,
   return_top_selection = FALSE
)
```

#### **Arguments**

spatPatObj	Output from detectSpatialPatterns
dimensions	dimensions to identify correlated genes for.
top_pos_genes	Top positively correlated genes.
top_neg_genes	Top negatively correlated genes.
min_pos_cor	Minimum positive correlation score to include a gene.
min_neg_cor	Minimum negative correlation score to include a gene.

224 select\_spatialNetwork

#### **Details**

Description.

## Value

Data.table with genes associated with selected dimension (PC).

# **Examples**

```
selectPatternGenes(gobject)
```

```
select\_expression\_values \\ select\_expression\_values
```

# Description

helper function to select expression values

# Usage

```
select_expression_values(gobject, values)
```

# Arguments

gobject giotto object

values expression values to extract

#### Value

expression matrix

```
select_spatialNetwork
```

# Description

function to select a spatial network

# Usage

```
select_spatialNetwork(gobject, name = NULL, return_network_Obj = FALSE)
```

set\_giotto\_python\_path 225

# Description

sets the python path and/or install miniconda and the python modules

# Usage

```
set_giotto_python_path(
   python_path = NULL,
   packages_to_install = c("pandas", "networkx", "python-igraph", "leidenalg",
        "python-louvain", "python.app", "scikit-learn")
)
```

show,giotto-method

show method for giotto class

# Description

show method for giotto class

# Usage

```
## S4 method for signature 'giotto'
show(object)
```

 $show {\tt ClusterDendrogram} \ \ \textit{show ClusterDendrogram}$ 

## **Description**

Creates dendrogram for selected clusters.

## Usage

```
showClusterDendrogram(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  cluster_column,
  cor = c("pearson", "spearman"),
  distance = "ward.D",
  h = NULL,
  h_color = "red",
  rotate = FALSE,
  show_plot = NA,
```

226 showClusterHeatmap

```
return_plot = NA,
save_plot = NA,
save_param = list(),
default_save_name = "showClusterDendrogram",
...
)
```

## **Arguments**

```
gobject
                  giotto object
expression_values
                  expression values to use
cluster_column name of column to use for clusters
cor
                  correlation score to calculate distance
                  distance method to use for hierarchical clustering
distance
h
                  height of horizontal lines to plot
h_color
                  color of horizontal lines
rotate
                  rotate dendrogram 90 degrees
show_plot
                  show plot
return_plot
                  return ggplot object
save_plot
                  directly save the plot [boolean]
save_param
                  list of saving parameters, see showSaveParameters
default_save_name
                  default save name for saving, don't change, change save_name in save_param
```

additional parameters for ggdendrogram()

## **Details**

Expression correlation dendrogram for selected clusters.

#### Value

ggplot

## **Examples**

showClusterDendrogram(gobject)

showClusterHeatmap showClusterHeatmap

# Description

Creates heatmap based on identified clusters

showClusterHeatmap 227

#### Usage

```
showClusterHeatmap(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  genes = "all",
  cluster_column,
  cor = c("pearson", "spearman"),
  distance = "ward.D",
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "showClusterHeatmap",
  ...
)
```

## **Arguments**

giotto object gobject expression\_values expression values to use vector of genes to use, default to 'all' genes cluster\_column name of column to use for clusters correlation score to calculate distance cor distance distance method to use for hierarchical clustering show\_plot show plot return\_plot return ggplot object save\_plot directly save the plot [boolean] save\_param list of saving parameters, see showSaveParameters default\_save\_name default save name for saving, don't change, change save\_name in save\_param

additional parameters for the Heatmap function from ComplexHeatmap

## **Details**

Correlation heatmap of selected clusters.

#### Value

ggplot

## **Examples**

```
showClusterHeatmap(gobject)
```

228 showGiottoInstructions

 $show {\tt GiottoImageNames} \quad \textit{show GiottoImageNames}$ 

# Description

Prints the available giotto images that are attached to the Giotto object

## Usage

```
showGiottoImageNames(gobject, verbose = TRUE)
```

## **Arguments**

gobject a giotto object

verbose verbosity of function

## Value

a vector of giotto image names attached to the giotto object

# **Examples**

```
showGiottoImageNames(gobject)
```

 ${\tt showGiottoInstructions}$ 

showGiottoInstructions

# Description

Function to display all instructions from giotto object

# Usage

```
showGiottoInstructions(gobject)
```

# Arguments

gobject giotto object

#### Value

named vector with giotto instructions

# **Examples**

showGiottoInstructions()

showGrids 229

showGrids

showGrids

# Description

Prints the available spatial grids that are attached to the Giotto object

# Usage

```
showGrids(gobject, verbose = TRUE)
```

# Arguments

gobject

a giotto object

verbose

verbosity of function#'

# Value

vector

# **Examples**

showGrids()

showNetworks

showNetworks

# Description

Prints the available spatial networks that are attached to the Giotto object

## Usage

```
showNetworks(gobject, verbose = TRUE)
```

# **Arguments**

gobject

a giotto object

verbose

verbosity of function#'

#### Value

vector

# **Examples**

showNetworks()

showPattern

showPattern

showPattern

# Description

```
show patterns for 2D spatial data
```

# Usage

```
showPattern(gobject, spatPatObj, ...)
```

# Arguments

gobject giotto object

spatPatObj Output from detectSpatialPatterns

dimension dimension to plot

trim Trim ends of the PC values.

background\_color

background color for plot

grid\_border\_color

color for grid

show\_legend show legend of ggplot

show\_plot show plot

return\_plot return ggplot object

save\_plot directly save the plot [boolean]

save\_param list of saving parameters from all\_plots\_save\_function

default\_save\_name

default save name for saving, don't change, change save\_name in save\_param

## Value

ggplot

## See Also

showPattern2D

## **Examples**

```
showPattern(gobject)
```

showPattern2D 231

showPattern2D

showPattern2D

## **Description**

show patterns for 2D spatial data

# Usage

```
showPattern2D(
  gobject,
  spatPatObj,
  dimension = 1,
  trim = c(0.02, 0.98),
  background_color = "white",
  grid_border_color = "grey",
  show_legend = T,
  point_size = 1,
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "showPattern2D")
```

## **Arguments**

```
gobject
                 giotto object
                 Output from detectSpatialPatterns
spatPatObj
dimension
                 dimension to plot
trim
                 Trim ends of the PC values.
background_color
                 background color for plot
grid_border_color
                 color for grid
                 show legend of ggplot
show_legend
show_plot
                 show plot
return_plot
                 return ggplot object
                 directly save the plot [boolean]
save_plot
                 list of saving parameters from all_plots_save_function
save_param
default_save_name
```

default save name for saving, don't change, change save\_name in save\_param

#### Value

ggplot

## **Examples**

```
{\tt showPattern2D(gobject)}
```

232 showPattern3D

showPattern3D

showPattern3D

## **Description**

show patterns for 3D spatial data

#### Usage

```
showPattern3D(
 gobject,
 spatPatObj,
 dimension = 1,
  trim = c(0.02, 0.98),
 background_color = "white",
 grid_border_color = "grey",
  show_legend = T,
 point_size = 1,
 axis_scale = c("cube", "real", "custom"),
 custom_ratio = NULL,
 x_ticks = NULL,
 y_ticks = NULL,
 z_ticks = NULL,
 show_plot = NA,
 return_plot = NA,
 save_plot = NA,
 save_param = list(),
 default_save_name = "showPattern3D"
```

## Arguments

```
gobject
                  giotto object
spatPatObj
                  Output from detectSpatialPatterns
dimension
                  dimension to plot
                  Trim ends of the PC values.
trim
background_color
                  background color for plot
grid_border_color
                  color for grid
                  show legend of plot
show_legend
point_size
                  adjust the point size
axis_scale
                  scale the axis
                  cutomize the scale of the axis
custom_ratio
x_ticks
                  the tick number of x_axis
                  the tick number of y_axis
y_ticks
z_ticks
                  the tick number of z_axis
```

showPatternGenes 233

```
show_plot show plot

return_plot return plot object

save_plot directly save the plot [boolean]

save_param list of saving parameters from all_plots_save_function

default_save_name

default save name for saving, don't change, change save_name in save_param
```

#### Value

plotly

## **Examples**

```
showPattern3D(gobject)
```

showPatternGenes

showPatternGenes

## **Description**

show genes correlated with spatial patterns

## Usage

```
showPatternGenes(
  gobject,
  spatPatObj,
  dimension = 1,
  top_pos_genes = 5,
  top_neg_genes = 5,
  point_size = 1,
  return_DT = FALSE,
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "showPatternGenes"
)
```

## **Arguments**

```
gobject
                  giotto object
                  Output from detectSpatialPatterns
spatPatObj
dimension
                  dimension to plot genes for.
                  Top positively correlated genes.
top_pos_genes
top_neg_genes
                  Top negatively correlated genes.
point_size
                  size of points
return_DT
                  if TRUE, it will return the data.table used to generate the plots
show_plot
                  show plot
```

234 showProcessingSteps

return\_plot return ggplot object

save\_plot directly save the plot [boolean]

save\_param list of saving parameters from all\_plots\_save\_function()

default\_save\_name

default save name for saving, don't change, change save\_name in save\_param

## Value

ggplot

# **Examples**

showPatternGenes(gobject)

 $show Processing Steps \qquad show Processing Steps$ 

## Description

shows the sequential processing steps that were performed in a summarized format

#### Usage

showProcessingSteps(gobject)

# **Arguments**

gobject giotto object

#### Value

list of processing steps and names

## **Examples**

showProcessingSteps(gobject)

showSaveParameters 235

showSaveParameters showSaveParameters

# Description

Description of Giotto saving options, links to all\_plots\_save\_function

#### Usage

```
showSaveParameters()
```

#### Value

Instruction on how to use the automatic plot saving options within Giotto

#### **Examples**

```
showSaveParameters()
```

showSpatialCorGenes

show Spatial Cor Genes

# Description

Shows and filters spatially correlated genes

## Usage

```
showSpatialCorGenes(
  spatCorObject,
  use_clus_name = NULL,
  selected_clusters = NULL,
  genes = NULL,
  min_spat_cor = 0.5,
  min_expr_cor = NULL,
  min_cor_diff = NULL,
  min_rank_diff = NULL,
  show_top_genes = NULL
)
```

#### **Arguments**

236 signPCA

```
min_cor_diff filter on minimum correlation difference (spatial vs expression)
min_rank_diff filter on minimum correlation rank difference (spatial vs expression)
show_top_genes show top genes per gene
```

#### Value

data.table with filtered information

# **Examples**

```
showSpatialCorGenes(gobject)
```

signPCA

signPCA

## **Description**

identify significant prinicipal components (PCs)

# Usage

```
signPCA(
 gobject,
 name = "pca",
 method = c("screeplot", "jackstraw"),
 expression_values = c("normalized", "scaled", "custom"),
 reduction = c("cells", "genes"),
 pca_method = c("irlba", "factominer"),
 rev = FALSE,
 genes_to_use = NULL,
 center = T,
  scale_unit = T,
 ncp = 50,
 scree_ylim = c(0, 10),
  jack_iter = 10,
  jack_threshold = 0.01,
  jack_ylim = c(0, 1),
 verbose = TRUE,
  show_plot = NA,
 return_plot = NA,
 save_plot = NA,
 save_param = list(),
 default_save_name = "signPCA"
)
```

# Arguments

```
gobject giotto object

name name of PCA object if available

method method to use to identify significant PCs
```

signPCA 237

expression\_values

expression values to use

reduction cells or genes

pca\_method which implementation to use

rev do a reverse PCA

genes\_to\_use subset of genes to use for PCA

center center data before PCA

scale\_unit scale features before PCA

ncp number of principal components to calculate

scree\_ylim y-axis limits on scree plot

jack\_iter number of interations for jackstraw

jack\_threshold p-value threshold to call a PC significant

jack\_ylim y-axis limits on jackstraw plot

verbose verbosity show\_plot show plot

return\_plot return ggplot object

save\_plot directly save the plot [boolean]

save\_param list of saving parameters from all\_plots\_save\_function()

default\_save\_name

default save name for saving, don't change, change save\_name in save\_param

#### **Details**

Two different methods can be used to assess the number of relevant or significant prinicipal components (PC's).

- 1. Screeplot works by plotting the explained variance of each individual PC in a barplot allowing you to identify which PC provides a significant contribution (a.k.a. 'elbow method').
- 2. The Jackstraw method uses the permutationPA function. By systematically permuting genes it identifies robust, and thus significant, PCs.

#### Value

ggplot object for scree method and maxtrix of p-values for jackstraw

## **Examples**

signPCA(gobject)

238 silhouetteRank

silhouetteRank si

silhouetteRank

## **Description**

Previously: calculate\_spatial\_genes\_python. This method computes a silhouette score per gene based on the spatial distribution of two partitions of cells (expressed L1, and non-expressed L0). Here, rather than L2 Euclidean norm, it uses a rank-transformed, exponentially weighted function to represent the local physical distance between two cells.

# Usage

```
silhouetteRank(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  metric = "euclidean",
  subset_genes = NULL,
  rbp_p = 0.95,
  examine_top = 0.3,
  python_path = NULL
)
```

specify specific path to python if required

# Arguments

```
gobject giotto object
expression_values
expression values to use

metric distance metric to use

subset_genes only run on this subset of genes

rbp_p fractional binarization threshold

examine_top top fraction to evaluate with silhouette
```

## Value

python\_path

data.table with spatial scores

# **Examples**

```
silhouetteRank(gobject)
```

spatCellCellcom 239

spatCellCellcom spatCellCellcom

#### **Description**

Spatial Cell-Cell communication scores based on spatial expression of interacting cells

#### Usage

```
spatCellCellcom(
  gobject,
 spatial_network_name = "Delaunay_network",
 cluster_column = "cell_types",
  random_iter = 1000,
 gene_set_1,
 gene_set_2,
  log2FC_addendum = 0.1,
 min_observations = 2,
 adjust_method = c("fdr", "bonferroni", "BH", "holm", "hochberg", "hommel", "BY",
    "none"),
  adjust_target = c("genes", "cells"),
  do_parallel = TRUE,
 cores = NA,
  verbose = c("a little", "a lot", "none")
)
```

#### **Arguments**

```
gobject
                  giotto object to use
spatial_network_name
                  spatial network to use for identifying interacting cells
cluster_column cluster column with cell type information
                  number of iterations
random_iter
                  first specific gene set from gene pairs
gene_set_1
gene_set_2
                  second specific gene set from gene pairs
log2FC_addendum
                  addendum to add when calculating log2FC
min_observations
                  minimum number of interactions needed to be considered
                  which method to adjust p-values
adjust_method
                  adjust multiple hypotheses at the cell or gene level
adjust_target
                  run calculations in parallel with mclapply
do_parallel
                  number of cores to use if do_parallel = TRUE
cores
                  verbose
verbose
```

# **Details**

Statistical framework to identify if pairs of genes (such as ligand-receptor combinations) are expressed at higher levels than expected based on a reshuffled null distribution of gene expression values in cells that are spatially in proximity to eachother.. More details will follow soon.

240 spatCellPlot

#### Value

Cell-Cell communication scores for gene pairs based on spatial interaction

#### **Examples**

```
spatCellCellcom(gobject)
```

spatCellPlot

spatCellPlot

## **Description**

Visualize cells according to spatial coordinates

## Usage

```
spatCellPlot(...)
```

# Arguments . . .

```
Arguments passed on to spatCellPlot2D
gobject giotto object
show_image show a tissue background image
gimage a giotto image
image_name name of a giotto image
sdimx x-axis dimension name (default = 'sdimx')
sdimy y-axis dimension name (default = 'sdimy')
spat_enr_names names of spatial enrichment results to include
cell_annotation_values numeric cell annotation columns
cell_color_gradient vector with 3 colors for numeric data
gradient_midpoint midpoint for color gradient
gradient_limits vector with lower and upper limits
select_cell_groups select subset of cells/clusters based on cell_color param-
    eter
select_cells select subset of cells based on cell IDs
point_shape shape of points (border, no_border or voronoi)
point_size size of point (cell)
point_alpha transparancy of spatial points
point_border_col color of border around points
point_border_stroke stroke size of border around points
show_cluster_center plot center of selected clusters
show_center_label plot label of selected clusters
center_point_size size of center points
label_size size of labels
label_fontface font of labels
show_network show underlying spatial network
spatial_network_name name of spatial network to use
```

spatCellPlot 241

```
network_color color of spatial network
network_alpha alpha of spatial network
show_grid show spatial grid
spatial_grid_name name of spatial grid to use
grid_color color of spatial grid
show_other_cells display not selected cells
other_cell_color color of not selected cells
other_point_size point size of not selected cells
other_cells_alpha alpha of not selected cells
coord_fix_ratio fix ratio between x and y-axis
show_legend show legend
legend_text size of legend text
legend_symbol_size size of legend symbols
background_color color of plot background
vor_border_color border colorr for voronoi plot
vor_max_radius maximum radius for voronoi 'cells'
vor_alpha transparancy of voronoi 'cells'
axis_text size of axis text
axis_title size of axis title
show_plot show plot
return_plot return ggplot object
save_plot directly save the plot [boolean]
save_param list of saving parameters, see showSaveParameters
default_save_name default save name for saving, don't change, change save_name
    in save_param
```

## **Details**

Description of parameters.

#### Value

ggplot

#### See Also

Other spatial cell annotation visualizations: spatCellPlot2D()

## **Examples**

spatCellPlot(gobject)

spatCellPlot2D

spatCellPlot2D

## **Description**

Visualize cells according to spatial coordinates

## Usage

```
spatCellPlot2D(
 gobject,
  show_image = F,
  gimage = NULL,
  image_name = "image",
  sdimx = "sdimx",
  sdimy = "sdimy"
  spat_enr_names = NULL,
  cell_annotation_values = NULL,
 cell_color_gradient = c("blue", "white", "red"),
 gradient_midpoint = NULL,
 gradient_limits = NULL,
 select_cell_groups = NULL,
  select_cells = NULL,
 point_shape = c("border", "no_border", "voronoi"),
 point_size = 3,
 point_alpha = 1,
 point_border_col = "black",
 point_border_stroke = 0.1,
  show_cluster_center = F,
  show\_center\_label = F,
  center_point_size = 4,
  center_point_border_col = "black",
  center_point_border_stroke = 0.1,
  label_size = 4,
  label_fontface = "bold",
  show_network = F,
  spatial_network_name = "Delaunay_network",
 network_color = NULL,
 network_alpha = 1,
  show_grid = F,
  spatial_grid_name = "spatial_grid",
 grid_color = NULL,
  show_other_cells = T,
 other_cell_color = "lightgrey",
 other_point_size = 1,
 other_cells_alpha = 0.1,
  coord_fix_ratio = NULL,
  show_legend = T,
  legend_text = 8,
  legend_symbol_size = 1,
 background_color = "white",
```

```
vor_border_color = "white",
      vor_max_radius = 200,
      vor_alpha = 1,
      axis_text = 8,
      axis_title = 8,
      cow_n_col = 2,
      cow_rel_h = 1,
      cow_rel_w = 1,
      cow_align = "h",
      show_plot = NA,
      return_plot = NA,
      save_plot = NA,
      save_param = list(),
      default_save_name = "spatCellPlot2D"
    )
Arguments
   gobject
                     giotto object
    show_image
                     show a tissue background image
    gimage
                     a giotto image
    image_name
                     name of a giotto image
    sdimx
                     x-axis dimension name (default = 'sdimx')
                     y-axis dimension name (default = 'sdimy')
    sdimy
    spat_enr_names names of spatial enrichment results to include
    cell_annotation_values
                     numeric cell annotation columns
    cell_color_gradient
                     vector with 3 colors for numeric data
    gradient_midpoint
                     midpoint for color gradient
    gradient_limits
                     vector with lower and upper limits
    select_cell_groups
                     select subset of cells/clusters based on cell_color parameter
                     select subset of cells based on cell IDs
    select_cells
                     shape of points (border, no_border or voronoi)
    point_shape
   point_size
                     size of point (cell)
    point_alpha
                     transparancy of spatial points
   point_border_col
                     color of border around points
   point_border_stroke
                     stroke size of border around points
    show_cluster_center
                     plot center of selected clusters
```

plot label of selected clusters

show\_center\_label

center\_point\_size size of center points label\_size size of labels label\_fontface font of labels show\_network show underlying spatial network spatial\_network\_name name of spatial network to use color of spatial network network\_color network\_alpha alpha of spatial network show\_grid show spatial grid spatial\_grid\_name name of spatial grid to use grid\_color color of spatial grid show\_other\_cells display not selected cells other\_cell\_color color of not selected cells other\_point\_size point size of not selected cells other\_cells\_alpha alpha of not selected cells coord\_fix\_ratio fix ratio between x and y-axis show\_legend show legend legend\_text size of legend text legend\_symbol\_size size of legend symbols background\_color color of plot background vor\_border\_color border colorr for voronoi plot vor\_max\_radius maximum radius for voronoi 'cells' vor\_alpha transparancy of voronoi 'cells' axis\_text size of axis text axis\_title size of axis title show\_plot show plot return ggplot object return\_plot directly save the plot [boolean] save\_plot list of saving parameters, see showSaveParameters save\_param

default save name for saving, don't change, change save\_name in save\_param

## **Details**

Description of parameters.

default\_save\_name

#### Value

ggplot

#### See Also

Other spatial cell annotation visualizations: spatCellPlot()

#### **Examples**

```
spatCellPlot2D(gobject)
```

spatDimCellPlot

spatDimCellPlot

# Description

Visualize numerical features of cells according to spatial AND dimension reduction coordinates in 2D

## Usage

```
spatDimCellPlot(...)
```

#### **Arguments**

Arguments passed on to spatDimCellPlot2D . . . gobject giotto object show\_image show a tissue background image gimage a giotto image image\_name name of a giotto image plot\_alignment direction to align plot spat\_enr\_names names of spatial enrichment results to include cell\_annotation\_values numeric cell annotation columns dim\_reduction\_to\_use dimension reduction to use dim\_reduction\_name dimension reduction name dim1\_to\_use dimension to use on x-axis dim2\_to\_use dimension to use on y-axis sdimx = spatial dimension to use on x-axis sdimy = spatial dimension to use on y-axis cell\_color\_gradient vector with 3 colors for numeric data gradient\_midpoint midpoint for color gradient gradient\_limits vector with lower and upper limits select\_cell\_groups select subset of cells/clusters based on cell\_color paramselect\_cells select subset of cells based on cell IDs dim\_point\_shape dim reduction points with border or not (border or no\_border) dim\_point\_size size of points in dim. reduction space

dim\_point\_alpha transparancy of dim. reduction points

```
dim_point_border_col border color of points in dim. reduction space
dim_point_border_stroke border stroke of points in dim. reduction space
spat_point_shape shape of points (border, no_border or voronoi)
spat_point_size size of spatial points
spat_point_alpha transparancy of spatial points
spat_point_border_col border color of spatial points
spat_point_border_stroke border stroke of spatial points
dim_show_cluster_center show the center of each cluster
dim_show_center_label provide a label for each cluster
dim_center_point_size size of the center point
dim_center_point_border_col border color of center point
dim_center_point_border_stroke stroke size of center point
dim_label_size size of the center label
dim_label_fontface font of the center label
spat show cluster center show the center of each cluster
spat_show_center_label provide a label for each cluster
spat_center_point_size size of the center point
spat_label_size size of the center label
spat_label_fontface font of the center label
show_NN_network show underlying NN network
nn_network_to_use type of NN network to use (kNN vs sNN)
nn_network_name name of NN network to use, if show_NN_network = TRUE
dim_edge_alpha column to use for alpha of the edges
spat_show_network show spatial network
spatial_network_name name of spatial network to use
spat_network_color color of spatial network
spat_show_grid show spatial grid
spatial_grid_name name of spatial grid to use
spat_grid_color color of spatial grid
show_other_cells display not selected cells
other_cell_color color of not selected cells
dim_other_point_size size of not selected dim cells
spat_other_point_size size of not selected spat cells
spat_other_cells_alpha alpha of not selected spat cells
coord_fix_ratio ratio for coordinates
cow_n_col cowplot param: how many columns
cow_rel_h cowplot param: relative height
cow_rel_w cowplot param: relative width
cow_align cowplot param: how to align
show_legend show legend
legend_text size of legend text
legend_symbol_size size of legend symbols
dim_background_color background color of points in dim. reduction space
spat_background_color background color of spatial points
vor_border_color border colorr for voronoi plot
```

```
vor_max_radius maximum radius for voronoi 'cells'
vor_alpha transparancy of voronoi 'cells'
axis_text size of axis text
axis_title size of axis title
show_plot show plot
return_plot return ggplot object
save_plot directly save the plot [boolean]
save_param list of saving parameters, see showSaveParameters
default_save_name default save name for saving, don't change, change save_name
in save_param
```

#### **Details**

Description of parameters.

#### Value

ggplot

#### See Also

Other spatial and dimension reduction cell annotation visualizations: spatDimCellPlot2D()

#### **Examples**

```
spatDimCellPlot(gobject)
```

spatDimCellPlot2D

spatDimCellPlot2D

# Description

Visualize numerical features of cells according to spatial AND dimension reduction coordinates in 2D

## Usage

```
spatDimCellPlot2D(
  gobject,
  show_image = F,
  gimage = NULL,
  image_name = "image",
  plot_alignment = c("vertical", "horizontal"),
  spat_enr_names = NULL,
  cell_annotation_values = NULL,
  dim_reduction_to_use = "umap",
  dim_reduction_name = "umap",
  dim1_to_use = 1,
  dim2_to_use = 2,
  sdimx = "sdimx",
  sdimy = "sdimy",
```

```
cell_color_gradient = c("blue", "white", "red"),
gradient_midpoint = NULL,
gradient_limits = NULL,
select_cell_groups = NULL,
select_cells = NULL,
dim_point_shape = c("border", "no_border"),
dim_point_size = 1,
dim_point_alpha = 1,
dim_point_border_col = "black",
dim_point_border_stroke = 0.1,
spat_point_shape = c("border", "no_border", "voronoi"),
spat_point_size = 1,
spat_point_alpha = 1,
spat_point_border_col = "black",
spat_point_border_stroke = 0.1,
dim_show_cluster_center = F,
dim_show_center_label = T,
dim_center_point_size = 4,
dim_center_point_border_col = "black",
dim_center_point_border_stroke = 0.1,
dim_label_size = 4,
dim_label_fontface = "bold",
spat_show_cluster_center = F,
spat_show_center_label = F,
spat_center_point_size = 4,
spat_center_point_border_col = "black",
spat_center_point_border_stroke = 0.1,
spat_label_size = 4,
spat_label_fontface = "bold",
show_NN_network = F,
nn_network_to_use = "sNN",
nn_network_name = "sNN.pca",
dim_edge_alpha = 0.5,
spat_show_network = F,
spatial_network_name = "Delaunay_network",
spat_network_color = "red",
spat_network_alpha = 0.5,
spat_show_grid = F,
spatial_grid_name = "spatial_grid",
spat_grid_color = "green",
show_other_cells = TRUE,
other_cell_color = "grey",
dim_other_point_size = 0.5,
spat_other_point_size = 0.5,
spat_other_cells_alpha = 0.5,
show_legend = T,
legend_text = 8,
legend_symbol_size = 1,
dim_background_color = "white",
spat_background_color = "white",
vor_border_color = "white",
vor_max_radius = 200,
```

```
vor_alpha = 1,
      axis_text = 8,
      axis_title = 8,
      coord_fix_ratio = NULL,
      cow_n_col = 2,
      cow_rel_h = 1,
      cow_rel_w = 1,
      cow_align = "h",
      show_plot = NA,
      return_plot = NA,
      save_plot = NA,
      save_param = list(),
      default_save_name = "spatDimCellPlot2D"
    )
Arguments
    gobject
                     giotto object
                     show a tissue background image
    show_image
    gimage
                     a giotto image
                     name of a giotto image
    image_name
    plot_alignment direction to align plot
    spat_enr_names names of spatial enrichment results to include
    cell_annotation_values
                     numeric cell annotation columns
    dim_reduction_to_use
                     dimension reduction to use
    dim_reduction_name
                     dimension reduction name
    dim1_to_use
                     dimension to use on x-axis
    dim2_to_use
                     dimension to use on y-axis
    sdimx
                     = spatial dimension to use on x-axis
    sdimv
                     = spatial dimension to use on y-axis
    cell_color_gradient
                      vector with 3 colors for numeric data
    gradient_midpoint
                     midpoint for color gradient
    gradient_limits
                     vector with lower and upper limits
    select_cell_groups
                     select subset of cells/clusters based on cell_color parameter
                     select subset of cells based on cell IDs
    select_cells
    dim_point_shape
                     dim reduction points with border or not (border or no_border)
    dim_point_size size of points in dim. reduction space
```

transparancy of dim. reduction points

dim\_point\_alpha

dim\_point\_border\_col border color of points in dim. reduction space dim\_point\_border\_stroke border stroke of points in dim. reduction space spat\_point\_shape shape of points (border, no\_border or voronoi) spat\_point\_size size of spatial points spat\_point\_alpha transparancy of spatial points spat\_point\_border\_col border color of spatial points spat\_point\_border\_stroke border stroke of spatial points dim\_show\_cluster\_center show the center of each cluster dim\_show\_center\_label provide a label for each cluster dim\_center\_point\_size size of the center point dim\_center\_point\_border\_col border color of center point dim\_center\_point\_border\_stroke stroke size of center point dim\_label\_size size of the center label dim\_label\_fontface font of the center label spat\_show\_cluster\_center show the center of each cluster spat\_show\_center\_label provide a label for each cluster  $spat\_center\_point\_size$ size of the center point spat\_label\_size size of the center label spat\_label\_fontface font of the center label show\_NN\_network show underlying NN network nn\_network\_to\_use type of NN network to use (kNN vs sNN) nn\_network\_name name of NN network to use, if show\_NN\_network = TRUE dim\_edge\_alpha column to use for alpha of the edges spat\_show\_network show spatial network spatial\_network\_name name of spatial network to use

```
spat_network_color
                  color of spatial network
spat_show_grid show spatial grid
spatial_grid_name
                  name of spatial grid to use
spat_grid_color
                  color of spatial grid
show_other_cells
                  display not selected cells
other_cell_color
                  color of not selected cells
dim_other_point_size
                  size of not selected dim cells
spat_other_point_size
                  size of not selected spat cells
spat_other_cells_alpha
                  alpha of not selected spat cells
                  show legend
show_legend
legend_text
                  size of legend text
legend_symbol_size
                  size of legend symbols
dim_background_color
                  background color of points in dim. reduction space
spat_background_color
                  background color of spatial points
vor_border_color
                  border colorr for voronoi plot
vor_max_radius maximum radius for voronoi 'cells'
vor_alpha
                  transparancy of voronoi 'cells'
axis_text
                  size of axis text
                  size of axis title
axis_title
coord_fix_ratio
                  ratio for coordinates
cow_n_col
                  cowplot param: how many columns
cow_rel_h
                  cowplot param: relative height
                  cowplot param: relative width
cow_rel_w
                  cowplot param: how to align
cow_align
show_plot
                  show plot
return_plot
                  return ggplot object
                  directly save the plot [boolean]
save_plot
                  list of saving parameters, see showSaveParameters
save_param
default_save_name
                  default save name for saving, don't change, change save_name in save_param
```

## **Details**

Description of parameters.

252 spatDimGenePlot

#### Value

ggplot

#### See Also

Other spatial and dimension reduction cell annotation visualizations: spatDimCellPlot()

#### **Examples**

```
spatDimCellPlot2D(gobject)
```

spatDimGenePlot

spatDimGenePlot

## **Description**

Visualize cells according to spatial AND dimension reduction coordinates in ggplot mode

#### Usage

```
spatDimGenePlot(...)
```

#### **Arguments**

Arguments passed on to spatDimGenePlot2D . . . gobject giotto object show\_image show a tissue background image gimage a giotto image image\_name name of a giotto image expression\_values gene expression values to use plot\_alignment direction to align plot genes genes to show dim\_reduction\_to\_use dimension reduction to use dim\_reduction\_name dimension reduction name dim1\_to\_use dimension to use on x-axis dim2\_to\_use dimension to use on y-axis dim\_point\_shape dim reduction points with border or not (border or no\_border) dim\_point\_size dim reduction plot: point size dim\_point\_alpha transparancy of dim. reduction points dim\_point\_border\_col color of border around points dim\_point\_border\_stroke stroke size of border around points show\_NN\_network show underlying NN network nn\_network\_to\_use type of NN network to use (kNN vs sNN) network\_name name of NN network to use, if show\_NN\_network = TRUE dim\_network\_color color of NN network edge\_alpha\_dim dim reduction plot: column to use for alpha of the edges scale\_alpha\_with\_expression scale expression with ggplot alpha parameter

spatDimGenePlot 253

```
sdimx spatial x-axis dimension name (default = 'sdimx')
sdimy spatial y-axis dimension name (default = 'sdimy')
spatial_network_name name of spatial network to use
spatial_network_color color of spatial network
spatial_grid_name name of spatial grid to use
spat_point_shape spatial points with border or not (border or no_border)
spat_point_size spatial plot: point size
spat_point_alpha transparancy of spatial points
spat_point_border_col color of border around points
spat_point_border_stroke stroke size of border around points
cell_color_gradient vector with 3 colors for numeric data
gradient_midpoint midpoint for color gradient
gradient_limits vector with lower and upper limits
show_legend show legend
legend_text_size of legend text
dim_background_color color of plot background for dimension plot
spat_background_color color of plot background for spatial plot
vor_border_color border colorr for voronoi plot
vor_max_radius maximum radius for voronoi 'cells'
vor_alpha transparancy of voronoi 'cells'
axis_text size of axis text
axis_title size of axis title
cow_n_col cowplot param: how many columns
cow_rel_h cowplot param: relative height
cow_rel_w cowplot param: relative width
cow_align cowplot param: how to align
show_plot show plots
return_plot return ggplot object
save_plot directly save the plot [boolean]
save_param list of saving parameters, see showSaveParameters
default_save_name default save name for saving, don't change, change save_name
    in save_param
```

#### **Details**

Description of parameters.

## Value

ggplot

#### See Also

```
spatDimGenePlot3D
```

Other spatial and dimension reduction gene expression visualizations: spatDimGenePlot2D(), spatDimGenePlot3D()

## **Examples**

```
{\tt spatDimGenePlot(gobject)}
```

254 spatDimGenePlot2D

spatDimGenePlot2D

spatDimGenePlot2D

## **Description**

Visualize cells according to spatial AND dimension reduction coordinates in ggplot mode

```
spatDimGenePlot2D(
 gobject,
  show_image = F,
  gimage = NULL,
  image_name = "image",
  expression_values = c("normalized", "scaled", "custom"),
 plot_alignment = c("vertical", "horizontal"),
  genes,
 dim_reduction_to_use = "umap",
 dim_reduction_name = "umap",
 dim1_to_use = 1,
 dim2_to_use = 2,
 dim_point_shape = c("border", "no_border"),
 dim_point_size = 1,
 dim_point_alpha = 1,
 dim_point_border_col = "black",
 dim_point_border_stroke = 0.1,
  show_NN_network = F,
  show_spatial_network = F,
 dim_network_color = "gray",
  show\_spatial\_grid = F,
 nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  edge_alpha_dim = NULL,
  scale_alpha_with_expression = FALSE,
  sdimx = "sdimx",
  sdimy = "sdimy",
  spatial_network_name = "Delaunay_network",
  spatial_network_color = NULL,
  spatial_grid_name = "spatial_grid",
  spat_point_shape = c("border", "no_border", "voronoi"),
  spat_point_size = 1,
  spat_point_alpha = 1,
  spat_point_border_col = "black",
  spat_point_border_stroke = 0.1,
 cell_color_gradient = c("blue", "white", "red"),
 gradient_midpoint = NULL,
 gradient_limits = NULL,
  cow_n_col = 2,
 cow_rel_h = 1,
  cow_rel_w = 1,
  cow_align = "h",
```

spatDimGenePlot2D 255

```
show_legend = T,
      legend_text = 8,
      dim_background_color = "white",
      spat_background_color = "white",
      vor_border_color = "white",
      vor_max_radius = 200,
      vor_alpha = 1,
      axis_text = 8,
      axis_title = 8,
      show_plot = NA,
      return_plot = NA,
      save_plot = NA,
      save_param = list(),
      default_save_name = "spatDimGenePlot2D"
    )
Arguments
    gobject
                     giotto object
                     show a tissue background image
    show_image
    gimage
                     a giotto image
    image_name
                     name of a giotto image
    expression_values
                     gene expression values to use
    plot_alignment direction to align plot
                     genes to show
    genes
    dim_reduction_to_use
                     dimension reduction to use
    dim_reduction_name
                     dimension reduction name
                     dimension to use on x-axis
    dim1_to_use
    dim2_to_use
                     dimension to use on y-axis
    dim_point_shape
                     dim reduction points with border or not (border or no_border)
    dim_point_size dim reduction plot: point size
    dim_point_alpha
                     transparancy of dim. reduction points
    dim_point_border_col
                     color of border around points
    dim_point_border_stroke
                     stroke size of border around points
    show_NN_network
                     show underlying NN network
    dim_network_color
                     color of NN network
    nn_network_to_use
                     type of NN network to use (kNN vs sNN)
```

name of NN network to use, if show\_NN\_network = TRUE

network\_name

edge\_alpha\_dim dim reduction plot: column to use for alpha of the edges scale\_alpha\_with\_expression scale expression with ggplot alpha parameter spatial x-axis dimension name (default = 'sdimx') sdimx spatial y-axis dimension name (default = 'sdimy') sdimy spatial\_network\_name name of spatial network to use spatial\_network\_color color of spatial network spatial\_grid\_name name of spatial grid to use spat\_point\_shape spatial points with border or not (border or no\_border) spat\_point\_size spatial plot: point size spat\_point\_alpha transparancy of spatial points spat\_point\_border\_col color of border around points spat\_point\_border\_stroke stroke size of border around points cell\_color\_gradient vector with 3 colors for numeric data gradient\_midpoint midpoint for color gradient gradient\_limits vector with lower and upper limits cowplot param: how many columns cow\_n\_col cowplot param: relative height cow\_rel\_h cow\_rel\_w cowplot param: relative width cow\_align cowplot param: how to align show\_legend show legend legend\_text size of legend text dim\_background\_color color of plot background for dimension plot spat\_background\_color color of plot background for spatial plot vor\_border\_color border colorr for voronoi plot vor\_max\_radius maximum radius for voronoi 'cells' transparancy of voronoi 'cells' vor\_alpha size of axis text axis\_text size of axis title axis\_title show\_plot show plots return\_plot return ggplot object directly save the plot [boolean] save\_plot list of saving parameters, see showSaveParameters save\_param default\_save\_name default save name for saving, don't change, change save\_name in save\_param spatDimGenePlot3D 257

#### **Details**

Description of parameters.

#### Value

ggplot

#### See Also

```
spatDimGenePlot3D
```

Other spatial and dimension reduction gene expression visualizations: spatDimGenePlot3D(), spatDimGenePlot()

## **Examples**

```
spatDimGenePlot2D(gobject)
```

spatDimGenePlot3D

spatDimGenePlot3D

# Description

Visualize cells according to spatial AND dimension reduction coordinates in ggplot mode

```
spatDimGenePlot3D(
 gobject,
 expression_values = c("normalized", "scaled", "custom"),
 plot_alignment = c("horizontal", "vertical"),
 dim_reduction_to_use = "umap",
 dim_reduction_name = "umap",
 dim1_to_use = 1,
 dim2_to_use = 2,
 dim3_to_use = NULL,
  sdimx = "sdimx",
  sdimy = "sdimy",
  sdimz = "sdimz",
  genes,
 cluster_column = NULL,
  select_cell_groups = NULL,
 select_cells = NULL,
  show_other_cells = T,
 other_cell_color = "lightgrey",
 other_point_size = 1.5,
  show_NN_network = F,
 nn_network_to_use = "sNN",
 network_name = "sNN.pca",
 label_size = 16,
 genes_low_color = "blue",
 genes_mid_color = "white",
```

258 spatDimGenePlot3D

genes\_high\_color = "red",

```
dim_point_size = 3,
      nn_network_alpha = 0.5,
      show_spatial_network = F,
      spatial_network_name = "Delaunay_network",
      network_color = "lightgray",
      spatial_network_alpha = 0.5,
      show_spatial_grid = F,
      spatial_grid_name = "spatial_grid",
      spatial_grid_color = NULL,
      spatial_grid_alpha = 0.5,
      spatial_point_size = 3,
      legend_text_size = 12,
      axis_scale = c("cube", "real", "custom"),
      custom_ratio = NULL,
      x_ticks = NULL,
      y_ticks = NULL,
      z_{ticks} = NULL,
      show_plot = NA,
      return_plot = NA,
      save_plot = NA,
      save_param = list(),
      default_save_name = "spatDimGenePlot3D"
Arguments
    gobject
                    giotto object
    expression_values
                    gene expression values to use
   plot_alignment direction to align plot
    dim_reduction_to_use
                    dimension reduction to use
   dim_reduction_name
                    dimension reduction name
   dim1_to_use
                    dimension to use on x-axis
   dim2_to_use
                    dimension to use on y-axis
                    dimension to use on z-axis
   dim3_to_use
                    genes to show
    genes
    show_NN_network
                    show underlying NN network
    nn\_network\_to\_use
                    type of NN network to use (kNN vs sNN)
                    name of NN network to use, if show_NN_network = TRUE
    network_name
    dim_point_size dim reduction plot: point size
    spatial_network_name
                    name of spatial network to use
    spatial_grid_name
                    name of spatial grid to use
```

spatDimPlot 259

```
spatial_point_size
```

spatial plot: point size

show\_plot show plots

return\_plot return plotly object

save\_plot directly save the plot [boolean]

show legend

save\_param list of saving parameters, see showSaveParameters

default\_save\_name

show\_legend

default save name for saving, don't change, change save\_name in save\_param

edge\_alpha\_dim dim reduction plot: column to use for alpha of the edges

scale\_alpha\_with\_expression

scale expression with ggplot alpha parameter

point\_size size of point (cell)

# Details

Description of parameters.

#### Value

plotly

## See Also

Other spatial and dimension reduction gene expression visualizations: spatDimGenePlot2D(), spatDimGenePlot()

## **Examples**

```
spatDimGenePlot3D(gobject)
```

spatDimPlot

spatDimPlot

# Description

Visualize cells according to spatial AND dimension reduction coordinates 2D

```
spatDimPlot(...)
```

260 spatDimPlot

## **Arguments**

Arguments passed on to spatDimPlot2D gobject giotto object show\_image show a tissue background image gimage a giotto image image\_name name of a giotto image plot\_alignment direction to align plot dim\_reduction\_to\_use dimension reduction to use dim\_reduction\_name dimension reduction name dim1\_to\_use dimension to use on x-axis dim2\_to\_use dimension to use on y-axis sdimx = spatial dimension to use on x-axis sdimy = spatial dimension to use on y-axis spat\_enr\_names names of spatial enrichment results to include cell\_color color for cells (see details) color\_as\_factor convert color column to factor cell\_color\_code named vector with colors cell\_color\_gradient vector with 3 colors for numeric data gradient\_midpoint midpoint for color gradient gradient\_limits vector with lower and upper limits select\_cell\_groups select subset of cells/clusters based on cell color paramselect\_cells select subset of cells based on cell IDs dim\_point\_shape point with border or not (border or no\_border) dim\_point\_size size of points in dim. reduction space dim\_point\_alpha transparancy of point in dim. reduction space dim\_point\_border\_col border color of points in dim. reduction space dim\_point\_border\_stroke border stroke of points in dim. reduction space spat\_point\_shape shape of points (border, no\_border or voronoi) spat\_point\_size size of spatial points spat\_point\_alpha transparancy of spatial points spat\_point\_border\_col border color of spatial points spat\_point\_border\_stroke border stroke of spatial points dim\_show\_cluster\_center show the center of each cluster dim\_show\_center\_label provide a label for each cluster dim\_center\_point\_size size of the center point dim\_center\_point\_border\_col border color of center point dim\_center\_point\_border\_stroke stroke size of center point dim\_label\_size size of the center label dim\_label\_fontface font of the center label spat\_show\_cluster\_center show the center of each cluster spat\_show\_center\_label provide a label for each cluster spat\_center\_point\_size size of the center point spat\_label\_size size of the center label spat\_label\_fontface font of the center label

spatDimPlot 261

```
show_NN_network show underlying NN network
nn_network_to_use type of NN network to use (kNN vs sNN)
network_name name of NN network to use, if show_NN_network = TRUE
nn_network_alpha column to use for alpha of the edges
show_spatial_network show spatial network
spat_network_name name of spatial network to use
spat_network_color color of spatial network
show_spatial_grid show spatial grid
spat_grid_name name of spatial grid to use
spat_grid_color color of spatial grid
show_other_cells display not selected cells
other_cell_color color of not selected cells
dim_other_point_size size of not selected dim cells
spat_other_point_size size of not selected spat cells
spat_other_cells_alpha alpha of not selected spat cells
dim_show_legend show legend of dimension reduction plot
spat_show_legend show legend of spatial plot
legend_text size of legend text
legend_symbol_size size of legend symbols
dim_background_color background color of points in dim. reduction space
spat_background_color background color of spatial points
vor_border_color border colorr for voronoi plot
vor_max_radius maximum radius for voronoi 'cells'
vor_alpha transparancy of voronoi 'cells'
axis_text size of axis text
axis_title size of axis title
show_plot show plot
return_plot return ggplot object
save_plot directly save the plot [boolean]
save_param list of saving parameters, see showSaveParameters
default_save_name default save name for saving, don't change, change save name
    in save_param
```

## **Details**

Description of parameters.

## Value

ggplot

#### See Also

```
spatDimPlot2D and spatDimPlot3D for 3D visualization.

Other spatial and dimension reduction visualizations: spatDimPlot2D(), spatDimPlot3D()
```

## Examples

```
spatDimPlot(gobject)
```

262 spatDimPlot2D

spatDimPlot2D

spatDimPlot2D

#### **Description**

Visualize cells according to spatial AND dimension reduction coordinates 2D

```
spatDimPlot2D(
 gobject,
  show_image = F,
  gimage = NULL,
  image_name = "image",
 plot_alignment = c("vertical", "horizontal"),
 dim_reduction_to_use = "umap",
 dim_reduction_name = "umap",
 dim1_to_use = 1,
 dim2_to_use = 2,
  sdimx = "sdimx",
  sdimy = "sdimy",
  spat_enr_names = NULL,
 cell_color = NULL,
  color_as_factor = T,
  cell_color_code = NULL,
 cell_color_gradient = c("blue", "white", "red"),
 gradient_midpoint = NULL,
  gradient_limits = NULL,
  select_cell_groups = NULL,
  select_cells = NULL,
  dim_point_shape = c("border", "no_border"),
  dim_point_size = 1,
 dim_point_alpha = 1,
 dim_point_border_col = "black",
 dim_point_border_stroke = 0.1,
  spat_point_shape = c("border", "no_border", "voronoi"),
  spat_point_size = 1,
  spat_point_alpha = 1,
  spat_point_border_col = "black",
  spat_point_border_stroke = 0.1,
 dim_show_cluster_center = F,
 dim_show_center_label = T,
 dim_center_point_size = 4,
 dim_center_point_border_col = "black",
 dim_center_point_border_stroke = 0.1,
 dim_label_size = 4,
 dim_label_fontface = "bold",
  spat_show_cluster_center = F,
  spat_show_center_label = F,
  spat_center_point_size = 4,
  spat_label_size = 4,
```

spatDimPlot2D 263

```
spat_label_fontface = "bold",
 show_NN_network = F,
 nn_network_to_use = "sNN",
 network_name = "sNN.pca",
 nn_network_alpha = 0.05,
 show_spatial_network = F,
 spat_network_name = "Delaunay_network",
 spat_network_color = "blue",
  spat_network_alpha = 0.5,
 show_spatial_grid = F,
 spat_grid_name = "spatial_grid",
 spat_grid_color = "blue",
 show_other_cells = T,
 other_cell_color = "lightgrey",
 dim\_other\_point\_size = 1,
 spat_other_point_size = 1,
 spat_other_cells_alpha = 0.5,
 dim_show_legend = F,
 spat_show_legend = F,
 legend_text = 8,
 legend_symbol_size = 1,
 dim_background_color = "white",
 spat_background_color = "white",
 vor_border_color = "white",
 vor_max_radius = 200,
 vor_alpha = 1,
 axis_text = 8,
 axis_title = 8,
 show_plot = NA,
 return_plot = NA,
 save_plot = NA,
 save_param = list(),
 default_save_name = "spatDimPlot2D"
)
```

# Arguments

```
gobject
                  giotto object
                  show a tissue background image
show_image
                  a giotto image
gimage
image_name
                  name of a giotto image
plot_alignment direction to align plot
dim_reduction_to_use
                  dimension reduction to use
dim_reduction_name
                  dimension reduction name
dim1_to_use
                  dimension to use on x-axis
dim2_to_use
                  dimension to use on y-axis
sdimx
                  = spatial dimension to use on x-axis
sdimy
                  = spatial dimension to use on y-axis
```

264 spatDimPlot2D

spat\_enr\_names names of spatial enrichment results to include color for cells (see details) cell\_color color\_as\_factor convert color column to factor cell\_color\_code named vector with colors cell\_color\_gradient vector with 3 colors for numeric data gradient\_midpoint midpoint for color gradient gradient\_limits vector with lower and upper limits select\_cell\_groups select subset of cells/clusters based on cell\_color parameter select subset of cells based on cell IDs select\_cells dim\_point\_shape point with border or not (border or no\_border) dim\_point\_size size of points in dim. reduction space dim\_point\_alpha transparancy of point in dim. reduction space dim\_point\_border\_col border color of points in dim. reduction space dim\_point\_border\_stroke border stroke of points in dim. reduction space spat\_point\_shape shape of points (border, no border or voronoi) spat\_point\_size size of spatial points spat\_point\_alpha transparancy of spatial points spat\_point\_border\_col border color of spatial points spat\_point\_border\_stroke border stroke of spatial points dim\_show\_cluster\_center show the center of each cluster dim\_show\_center\_label provide a label for each cluster dim\_center\_point\_size size of the center point dim\_center\_point\_border\_col border color of center point dim\_center\_point\_border\_stroke stroke size of center point dim\_label\_size size of the center label dim\_label\_fontface font of the center label

spat\_background\_color

spat\_show\_cluster\_center show the center of each cluster spat\_show\_center\_label provide a label for each cluster spat\_center\_point\_size size of the center point spat\_label\_size size of the center label spat\_label\_fontface font of the center label show\_NN\_network show underlying NN network  $nn\_network\_to\_use$ type of NN network to use (kNN vs sNN) name of NN network to use, if show\_NN\_network = TRUE network\_name nn\_network\_alpha column to use for alpha of the edges show\_spatial\_network show spatial network spat\_network\_name name of spatial network to use spat\_network\_color color of spatial network show\_spatial\_grid show spatial grid spat\_grid\_name name of spatial grid to use spat\_grid\_color color of spatial grid show\_other\_cells display not selected cells other\_cell\_color color of not selected cells  ${\tt dim\_other\_point\_size}$ size of not selected dim cells spat\_other\_point\_size size of not selected spat cells spat\_other\_cells\_alpha alpha of not selected spat cells dim\_show\_legend show legend of dimension reduction plot spat\_show\_legend show legend of spatial plot legend\_text size of legend text legend\_symbol\_size size of legend symbols dim\_background\_color background color of points in dim. reduction space

background color of spatial points

266 spatDimPlot3D

```
vor_border_color
                  border colorr for voronoi plot
vor_max_radius maximum radius for voronoi 'cells'
                  transparancy of voronoi 'cells'
vor_alpha
axis_text
                  size of axis text
axis_title
                  size of axis title
show_plot
                  show plot
return_plot
                  return ggplot object
save_plot
                  directly save the plot [boolean]
save_param
                  list of saving parameters, see showSaveParameters
default_save_name
                  default save name for saving, don't change, change save_name in save_param
```

#### **Details**

Description of parameters.

#### Value

ggplot

## See Also

```
spatDimPlot3D
```

Other spatial and dimension reduction visualizations: spatDimPlot3D(), spatDimPlot()

## **Examples**

```
spatDimPlot2D(gobject)
```

spatDimPlot3D

# Description

Visualize cells according to spatial AND dimension reduction coordinates in plotly mode

```
spatDimPlot3D(
  gobject,
  plot_alignment = c("horizontal", "vertical"),
  dim_reduction_to_use = "umap",
  dim_reduction_name = "umap",
  dim1_to_use = 1,
  dim2_to_use = 2,
  dim3_to_use = 3,
  sdimx = "sdimx",
  sdimy = "sdimy",
```

spatDimPlot3D 267

```
sdimz = "sdimz",
show_NN_network = F,
nn_network_to_use = "sNN",
network_name = "sNN.pca",
show_cluster_center = F,
show_center_label = T,
center_point_size = 4,
label_size = 16,
select_cell_groups = NULL,
select_cells = NULL,
show_other_cells = T,
other_cell_color = "lightgrey",
other_point_size = 1.5,
cell_color = NULL,
color_as_factor = T,
cell_color_code = NULL,
dim_point_size = 3,
nn_network_alpha = 0.5,
show_spatial_network = F,
spatial_network_name = "Delaunay_network",
network_color = "lightgray",
spatial_network_alpha = 0.5,
show_spatial_grid = F,
spatial_grid_name = "spatial_grid",
spatial_grid_color = NULL,
spatial_grid_alpha = 0.5,
spatial_point_size = 3,
axis_scale = c("cube", "real", "custom"),
custom_ratio = NULL,
x_ticks = NULL,
y_ticks = NULL,
z_ticks = NULL,
legend_text_size = 12,
show_plot = NA,
return_plot = NA,
save_plot = NA,
save_param = list(),
default_save_name = "spatDimPlot3D"
```

#### **Arguments**

268 spatDimPlot3D

sdimx = spatial dimension to use on x-axis sdimy = spatial dimension to use on y-axis sdimz = spatial dimension to use on z-axis show\_NN\_network show underlying NN network nn\_network\_to\_use type of NN network to use (kNN vs sNN) name of NN network to use, if show\_NN\_network = TRUE network\_name show\_cluster\_center show the center of each cluster show\_center\_label provide a label for each cluster center\_point\_size size of the center point label\_size size of the center label select\_cell\_groups select subset of cells/clusters based on cell\_color parameter select\_cells select subset of cells based on cell IDs show\_other\_cells display not selected cells other\_cell\_color color of not selected cells other\_point\_size size of not selected cells color for cells (see details) cell\_color color\_as\_factor convert color column to factor cell\_color\_code named vector with colors dim\_point\_size size of points in dim. reduction space nn\_network\_alpha column to use for alpha of the edges show\_spatial\_network show spatial network spatial\_network\_name name of spatial network to use spatial\_network\_alpha alpha of spatial network show\_spatial\_grid show spatial grid spatial\_grid\_name name of spatial grid to use spatial\_grid\_color color of spatial grid spatial\_point\_size size of spatial points show plot show\_plot

spatGenePlot 269

```
return ggplot object
return_plot
save_plot
                  directly save the plot [boolean]
save_param
                  list of saving parameters, see showSaveParameters
default_save_name
                  default save name for saving, don't change, change save_name in save_param
dim_point_border_col
                  border color of points in dim. reduction space
dim_point_border_stroke
                  border stroke of points in dim. reduction space
spatial_network_color
                  color of spatial network
spatial_other_point_size
                  size of not selected spatial points
spatial_other_cells_alpha
                  alpha of not selected spatial points
dim_other_point_size
                  size of not selected dim. reduction points
```

## **Details**

Description of parameters.

show legend

show\_legend

## Value

plotly

# See Also

Other spatial and dimension reduction visualizations: spatDimPlot2D(), spatDimPlot()

## **Examples**

```
spatDimPlot3D(gobject)
```

spatGenePlot

spatGenePlot

# Description

Visualize cells and gene expression according to spatial coordinates

```
spatGenePlot(...)
```

270 spatGenePlot

# Arguments . . .

Arguments passed on to spatGenePlot2D gobject giotto object show\_image show a tissue background image gimage a giotto image image\_name name of a giotto image sdimx x-axis dimension name (default = 'sdimx') sdimy y-axis dimension name (default = 'sdimy') expression\_values gene expression values to use genes genes to show cell\_color\_gradient vector with 3 colors for numeric data gradient\_midpoint midpoint for color gradient gradient\_limits vector with lower and upper limits show\_network show underlying spatial network network\_color color of spatial network spatial\_network\_name name of spatial network to use show\_grid show spatial grid grid\_color color of spatial grid spatial\_grid\_name name of spatial grid to use midpoint expression midpoint scale\_alpha\_with\_expression scale expression with ggplot alpha parameter point\_shape shape of points (border, no\_border or voronoi) point\_size size of point (cell) point\_alpha transparancy of points point\_border\_col color of border around points point\_border\_stroke stroke size of border around points cow\_n\_col cowplot param: how many columns cow\_rel\_h cowplot param: relative height cow\_rel\_w cowplot param: relative width cow\_align cowplot param: how to align show\_legend show legend legend\_text size of legend text background\_color color of plot background vor\_border\_color border colorr for voronoi plot vor\_max\_radius maximum radius for voronoi 'cells' vor\_alpha transparancy of voronoi 'cells' axis\_text size of axis text axis\_title size of axis title show\_plot show plots return\_plot return ggplot object save\_plot directly save the plot [boolean] save\_param list of saving parameters, see showSaveParameters default\_save\_name default save name for saving, don't change, change save\_name in save\_param

spatGenePlot2D 271

#### **Details**

Description of parameters.

#### Value

ggplot

#### See Also

```
spatGenePlot3D and spatGenePlot2D
```

Other spatial gene expression visualizations: spatGenePlot2D(), spatGenePlot3D()

## **Examples**

```
spatGenePlot(gobject)
```

spatGenePlot2D

spatGenePlot2D

# Description

Visualize cells and gene expression according to spatial coordinates

```
spatGenePlot2D(
  gobject,
  show_image = F,
  gimage = NULL,
  image_name = "image",
  sdimx = "sdimx",
  sdimy = "sdimy",
  expression_values = c("normalized", "scaled", "custom"),
  genes,
  cell_color_gradient = c("blue", "white", "red"),
  gradient_midpoint = NULL,
  gradient_limits = NULL,
  show_network = F,
  network_color = NULL,
  spatial_network_name = "Delaunay_network",
  edge_alpha = NULL,
  show\_grid = F,
  grid_color = NULL,
  spatial_grid_name = "spatial_grid",
  midpoint = 0,
  scale_alpha_with_expression = FALSE,
  point_shape = c("border", "no_border", "voronoi"),
  point_size = 1,
  point_alpha = 1,
  point_border_col = "black",
  point_border_stroke = 0.1,
```

272 spatGenePlot2D

```
show_legend = T,
      legend_text = 8,
      background_color = "white",
      vor_border_color = "white",
      vor_alpha = 1,
      vor_max_radius = 200,
      axis_text = 8,
      axis_title = 8,
      cow_n_col = 2,
      cow_rel_h = 1,
      cow_rel_w = 1,
      cow_align = "h",
      show_plot = NA,
      return_plot = NA,
      save_plot = NA,
      save_param = list(),
      default_save_name = "spatGenePlot2D"
    )
Arguments
   gobject
                     giotto object
                     show a tissue background image
    show_image
                     a giotto image
    gimage
                     name of a giotto image
    image_name
                     x-axis dimension name (default = 'sdimx')
    sdimx
                     y-axis dimension name (default = 'sdimy')
    sdimy
    expression_values
                     gene expression values to use
                     genes to show
    genes
   cell_color_gradient
                     vector with 3 colors for numeric data
    gradient_midpoint
                     midpoint for color gradient
    gradient_limits
                     vector with lower and upper limits
                     show underlying spatial network
    show_network
    network_color
                     color of spatial network
    spatial_network_name
                     name of spatial network to use
    show_grid
                     show spatial grid
                     color of spatial grid
    grid_color
    spatial_grid_name
                     name of spatial grid to use
   midpoint
                     expression midpoint
    scale_alpha_with_expression
```

scale expression with ggplot alpha parameter shape of points (border, no\_border or voronoi)

point\_shape

spatGenePlot2D 273

```
size of point (cell)
point_size
                  transparancy of points
point_alpha
point_border_col
                  color of border around points
point_border_stroke
                  stroke size of border around points
show_legend
                  show legend
legend_text
                  size of legend text
background_color
                  color of plot background
vor_border_color
                  border colorr for voronoi plot
                  transparancy of voronoi 'cells'
vor_alpha
vor_max_radius maximum radius for voronoi 'cells'
                  size of axis text
axis_text
axis_title
                  size of axis title
                  cowplot param: how many columns
cow_n_col
cow_rel_h
                  cowplot param: relative height
cow_rel_w
                  cowplot param: relative width
                  cowplot param: how to align
cow_align
show_plot
                  show plots
return_plot
                  return ggplot object
                  directly save the plot [boolean]
save_plot
save_param
                  list of saving parameters, see showSaveParameters
default_save_name
                  default save name for saving, don't change, change save_name in save_param
                  additional parameters for cowplot::save_plot()
. . .
```

## **Details**

Description of parameters.

## Value

ggplot

#### See Also

```
spatGenePlot3D
```

Other spatial gene expression visualizations: spatGenePlot3D(), spatGenePlot()

## **Examples**

```
spatGenePlot2D(gobject)
```

274 spatGenePlot3D

spatGenePlot3D spatGenePlot3D

# Description

Visualize cells and gene expression according to spatial coordinates

#### Usage

```
spatGenePlot3D(
 gobject,
  expression_values = c("normalized", "scaled", "custom"),
 genes,
 show_network = F,
 network_color = NULL,
  spatial_network_name = "Delaunay_network",
 edge_alpha = NULL,
 show\_grid = F,
 cluster_column = NULL,
  select_cell_groups = NULL,
 select_cells = NULL,
 show_other_cells = T,
 other_cell_color = "lightgrey",
 other_point_size = 1,
 genes_high_color = NULL,
 genes_mid_color = "white",
 genes_low_color = "blue",
  spatial_grid_name = "spatial_grid",
 point_size = 2,
  show_legend = T,
  axis_scale = c("cube", "real", "custom"),
  custom_ratio = NULL,
 x_ticks = NULL,
 y_ticks = NULL,
 z_ticks = NULL,
 show_plot = NA,
 return_plot = NA,
 save_plot = NA,
 save_param = list(),
 default_save_name = "spatGenePlot3D"
)
```

# **Arguments**

```
gobject giotto object
expression_values
gene expression values to use
genes genes to show
show_network show underlying spatial network
network_color color of spatial network
```

spatGenePlot3D 275

```
spatial_network_name
                  name of spatial network to use
show_grid
                  show spatial grid
genes_high_color
                  color represents high gene expression
genes_mid_color
                  color represents middle gene expression
genes_low_color
                  color represents low gene expression
spatial_grid_name
                  name of spatial grid to use
                  size of point (cell)
point_size
show_legend
                  show legend
show_plot
                  show plots
                  return ggplot object
return_plot
                  directly save the plot [boolean]
save_plot
save_param
                  list of saving parameters, see showSaveParameters
default_save_name
                  default save name for saving, don't change, change save_name in save_param
grid_color
                  color of spatial grid
midpoint
                  expression midpoint
scale_alpha_with_expression
                  scale expression with ggplot alpha parameter
                  additional parameters for cowplot::save_plot()
```

# Details

Description of parameters.

## Value

ggplot

## See Also

Other spatial gene expression visualizations: spatGenePlot2D(), spatGenePlot()

# **Examples**

```
spatGenePlot3D(gobject)
```

276 spatialAEH

spatialAEH

spatialAEH

# Description

Compute spatial variable genes with spatialDE method

# Usage

```
spatialAEH(
  gobject = NULL,
  SpatialDE_results = NULL,
  name_pattern = "AEH_patterns",
  expression_values = c("raw", "normalized", "scaled", "custom"),
  pattern_num = 6,
  l = 1.05,
  python_path = NULL,
  return_gobject = TRUE
)
```

## **Arguments**

```
gobject Giotto object

SpatialDE_results
results of SpatialDE function

name_pattern name for the computed spatial patterns
expression_values
gene expression values to use

pattern_num number of spatial patterns to look for

lengthscale

python_path specify specific path to python if required

return_gobject show plot
```

# **Details**

This function is a wrapper for the SpatialAEH method implemented in the ...

#### Value

An updated giotto object

## **Examples**

```
spatialAEH(gobject)
```

spatialDE 277

spatialDE spatialDE

## **Description**

Compute spatial variable genes with spatialDE method

## Usage

```
spatialDE(
  gobject = NULL,
  expression_values = c("raw", "normalized", "scaled", "custom"),
  size = c(4, 2, 1),
  color = c("blue", "green", "red"),
  sig_alpha = 0.5,
  unsig_alpha = 0.5,
  python_path = NULL,
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "SpatialDE"
)
```

## **Arguments**

```
gobject
                  Giotto object
expression_values
                  gene expression values to use
size
                  size of plot
color
                  low/medium/high color scheme for plot
                  alpha value for significance
sig_alpha
                  alpha value for unsignificance
unsig_alpha
                  specify specific path to python if required
python_path
show_plot
                  show plot
return_plot
                  return ggplot object
save_plot
                  directly save the plot [boolean]
                  list of saving parameters from all_plots_save_function()
save_param
default_save_name
                  default save name for saving, don't change, change save_name in save_param
```

#### **Details**

This function is a wrapper for the SpatialDE method implemented in the ...

#### Value

a list of data.frames with results and plot (optional)

278 spatNetwDistributions

#### **Examples**

```
spatialDE(gobject)
```

 $spatNetwDistributions\ spatNetwDistributionsDistance$ 

## **Description**

This function return histograms displaying the distance distribution for each spatial k-neighbor

# Usage

```
spatNetwDistributions(
  gobject,
  spatial_network_name = "spatial_network",
  distribution = c("distance", "k_neighbors"),
  hist_bins = 30,
  test_distance_limit = NULL,
  ncol = 1,
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "spatNetwDistributions"
)
```

#### **Arguments**

```
Giotto object
gobject
spatial_network_name
                  name of spatial network
                  show the distribution of cell-to-cell distance or number of k neighbors
distribution
hist_bins
                  number of binds to use for the histogram
test_distance_limit
                  effect of different distance threshold on k-neighbors
ncol
                  number of columns to visualize the histograms in
show_plot
                  show plot
return_plot
                  return ggplot object
save_plot
                  directly save the plot [boolean]
                  list of saving parameters from all_plots_save_function
save_param
default_save_name
                  default save name for saving, alternatively change save_name in save_param
```

#### **Details**

The **distance** option shows the spatial distance distribution for each nearest neighbor rank (1st, 2nd, 3th, ... neigbor). With this option the user can also test the effect of a distance limit on the spatial network. This distance limit can be used to remove neigbor cells that are considered to far away. The **k\_neighbors** option shows the number of k neighbors distribution over all cells.

#### Value

```
ggplot plot
```

#### **Examples**

```
spatNetwDistributionsDistance(gobject)
```

```
spat {\tt NetwDistributionsDistance} \\ spat {\tt NetwDistributionsDistance}
```

## **Description**

This function return histograms displaying the distance distribution for each spatial k-neighbor

## Usage

```
spatNetwDistributionsDistance(
  gobject,
  spatial_network_name = "spatial_network",
  hist_bins = 30,
  test_distance_limit = NULL,
  ncol = 1,
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "spatNetwDistributionsDistance")
```

## **Arguments**

```
gobject
                  Giotto object
spatial_network_name
                  name of spatial network
                  number of binds to use for the histogram
hist_bins
test_distance_limit
                  effect of different distance threshold on k-neighbors
ncol
                  number of columns to visualize the histograms in
show_plot
                  show plot
return_plot
                  return ggplot object
                  directly save the plot [boolean]
save_plot
                  list of saving parameters from all_plots_save_function
save_param
default_save_name
                  default save name for saving, alternatively change save_name in save_param
```

# Value

```
ggplot plot
```

## **Examples**

```
spatNetwDistributionsDistance(gobject)
```

```
spat Netw Distributions Kneighbors \\ spat Netw Distributions Kneighbors
```

## **Description**

This function returns a histogram displaying the number of k-neighbors distribution for each cell

## Usage

```
spatNetwDistributionsKneighbors(
  gobject,
  spatial_network_name = "spatial_network",
  hist_bins = 30,
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "spatNetwDistributionsKneighbors")
```

## **Arguments**

```
Giotto object
gobject
spatial_network_name
                  name of spatial network
hist_bins
                  number of binds to use for the histogram
show_plot
                  show plot
                  return ggplot object
return_plot
                  directly save the plot [boolean]
save_plot
                  list of saving parameters from all_plots_save_function
save_param
default_save_name
                  default save name for saving, alternatively change save_name in save_param
```

# Value

ggplot plot

## **Examples**

```
spatNetwDistributionsKneighbors(gobject)
```

spatPlot 281

spatPlot spatPlot

#### **Description**

Visualize cells according to spatial coordinates

#### Usage

```
spatPlot(...)
```

#### **Arguments**

```
Arguments passed on to spatPlot2D
. . .
                 gobject giotto object
                 show_image show a tissue background image
                 gimage a giotto image
                 image_name name of a giotto image
                 group_by_subset subset the group_by factor column
                 sdimx x-axis dimension name (default = 'sdimx')
                 sdimy y-axis dimension name (default = 'sdimy')
                 spat_enr_names names of spatial enrichment results to include
                 cell_color color for cells (see details)
                 color_as_factor convert color column to factor
                 cell_color_code named vector with colors
                 cell_color_gradient vector with 3 colors for numeric data
                 gradient_midpoint midpoint for color gradient
                 gradient_limits vector with lower and upper limits
                 select_cell_groups select subset of cells/clusters based on cell_color param-
                 select_cells select subset of cells based on cell IDs
                 point_shape shape of points (border, no_border or voronoi)
                 point_size size of point (cell)
                 point_alpha transparancy of point
                 point_border_col color of border around points
                 point_border_stroke stroke size of border around points
                 show_cluster_center plot center of selected clusters
                 show_center_label plot label of selected clusters
                 center_point_size size of center points
                 label_size size of labels
                 label_fontface font of labels
                 show_network show underlying spatial network
                 spatial_network_name name of spatial network to use
                 network_color color of spatial network
                 network_alpha alpha of spatial network
```

show\_grid show spatial grid

282 spatPlot

```
spatial_grid_name name of spatial grid to use
grid_color color of spatial grid
show_other_cells display not selected cells
other_cell_color color of not selected cells
other_point_size point size of not selected cells
other_cells_alpha alpha of not selected cells
coord_fix_ratio fix ratio between x and y-axis
title title of plot
show_legend show legend
legend_text size of legend text
legend_symbol_size size of legend symbols
background_color color of plot background
vor_border_color border colorr for voronoi plot
vor_max_radius maximum radius for voronoi 'cells'
vor_alpha transparancy of voronoi 'cells'
axis_text size of axis text
axis_title size of axis title
cow_n_col cowplot param: how many columns
cow_rel_h cowplot param: relative height
cow_rel_w cowplot param: relative width
cow_align cowplot param: how to align
show_plot show plot
return_plot return ggplot object
save_plot directly save the plot [boolean]
save_param list of saving parameters, see showSaveParameters
default_save_name default save name for saving, don't change, change save_name
    in save_param
```

#### **Details**

Description of parameters.

## Value

ggplot

### See Also

```
spatPlot3D
```

Other spatial visualizations: spatPlot2D(), spatPlot3D()

## **Examples**

```
spatPlot(gobject)
```

spatPlot2D 283

spatPlot2D

spatPlot2D

#### **Description**

Visualize cells according to spatial coordinates

```
spatPlot2D(
 gobject,
  show_image = F,
  gimage = NULL,
  image_name = "image",
 group_by = NULL,
 group_by_subset = NULL,
  sdimx = "sdimx",
  sdimy = "sdimy",
  spat_enr_names = NULL,
 cell_color = NULL,
 color_as_factor = T,
 cell_color_code = NULL,
 cell_color_gradient = c("blue", "white", "red"),
  gradient_midpoint = NULL,
  gradient_limits = NULL,
 select_cell_groups = NULL,
  select_cells = NULL,
 point_shape = c("border", "no_border", "voronoi"),
 point_size = 3,
 point_alpha = 1,
 point_border_col = "black",
  point_border_stroke = 0.1,
  show_cluster_center = F,
  show_center_label = F,
  center_point_size = 4,
  center_point_border_col = "black",
  center_point_border_stroke = 0.1,
  label_size = 4,
 label_fontface = "bold",
  show_network = F,
  spatial_network_name = "Delaunay_network",
 network_color = NULL,
 network_alpha = 1,
  show_grid = F,
  spatial_grid_name = "spatial_grid",
 grid_color = NULL,
  show_other_cells = T,
 other_cell_color = "lightgrey",
 other_point_size = 1,
 other_cells_alpha = 0.1,
  coord_fix_ratio = NULL,
```

284 spatPlot2D

```
title = NULL,
  show_legend = T,
  legend_text = 8,
  legend_symbol_size = 1,
  background_color = "white",
  vor_border_color = "white",
  vor_max_radius = 200,
  vor_alpha = 1,
  axis_text = 8,
  axis_title = 8,
  cow_n_col = 2,
  cow_rel_h = 1,
  cow_rel_w = 1,
  cow_align = "h",
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
 save_param = list(),
 default_save_name = "spatPlot2D"
)
```

## **Arguments**

```
gobject
                  giotto object
                  show a tissue background image
show_image
                  a giotto image
gimage
                  name of a giotto image
image_name
group_by_subset
                  subset the group_by factor column
                  x-axis dimension name (default = 'sdimx')
sdimx
                  y-axis dimension name (default = 'sdimy')
sdimy
spat_enr_names names of spatial enrichment results to include
cell_color
                  color for cells (see details)
color_as_factor
                  convert color column to factor
cell_color_code
                  named vector with colors
cell_color_gradient
                  vector with 3 colors for numeric data
gradient_midpoint
                  midpoint for color gradient
gradient_limits
                  vector with lower and upper limits
select_cell_groups
                  select subset of cells/clusters based on cell_color parameter
select_cells
                  select subset of cells based on cell IDs
                  shape of points (border, no_border or voronoi)
point_shape
                  size of point (cell)
point_size
point_alpha
                  transparancy of point
```

spatPlot2D 285

point\_border\_col color of border around points point\_border\_stroke stroke size of border around points show\_cluster\_center plot center of selected clusters show\_center\_label plot label of selected clusters center\_point\_size size of center points size of labels label\_size label\_fontface font of labels show\_network show underlying spatial network spatial\_network\_name name of spatial network to use color of spatial network network\_color network\_alpha alpha of spatial network show\_grid show spatial grid spatial\_grid\_name name of spatial grid to use grid\_color color of spatial grid show\_other\_cells display not selected cells other\_cell\_color color of not selected cells other\_point\_size point size of not selected cells other\_cells\_alpha alpha of not selected cells coord\_fix\_ratio fix ratio between x and y-axis title title of plot show\_legend show legend legend\_text size of legend text legend\_symbol\_size size of legend symbols background\_color color of plot background vor\_border\_color border colorr for voronoi plot vor\_max\_radius maximum radius for voronoi 'cells' vor\_alpha transparancy of voronoi 'cells' axis\_text size of axis text

size of axis title

cowplot param: how many columns

axis\_title

cow\_n\_col

286 spatPlot3D

```
cowplot param: relative height
cow_rel_h
                  cowplot param: relative width
cow_rel_w
                  cowplot param: how to align
cow_align
                  show plot
show_plot
                  return ggplot object
return_plot
save_plot
                  directly save the plot [boolean]
                  list of saving parameters, see showSaveParameters
save_param
default_save_name
                  default save name for saving, don't change, change save_name in save_param
                  create multiple plots based on cell annotation column
groub_by
```

#### **Details**

Description of parameters.

#### Value

ggplot

#### See Also

```
spatPlot3D
```

Other spatial visualizations: spatPlot3D(), spatPlot()

## **Examples**

```
spatPlot2D(gobject)
```

 ${\tt spatPlot3D}$ 

spatPlot3D

## **Description**

Visualize cells according to spatial coordinates

```
spatPlot3D(
  gobject,
  sdimx = "sdimx",
  sdimy = "sdimy",
  sdimz = "sdimz",
  point_size = 3,
  cell_color = NULL,
  cell_color_code = NULL,
  select_cell_groups = NULL,
  select_cells = NULL,
  show_other_cells = T,
  other_cell_color = "lightgrey",
```

spatPlot3D 287

```
other_point_size = 0.5,
  show_network = F,
 network_color = NULL,
 network_alpha = 1,
 other_cell_alpha = 0.5,
  spatial_network_name = "Delaunay_network",
  show\_grid = F,
  grid_color = NULL,
 grid_alpha = 1,
  spatial_grid_name = "spatial_grid",
  title = "",
  show_legend = T,
 axis_scale = c("cube", "real", "custom"),
 custom_ratio = NULL,
 x_ticks = NULL,
 y_ticks = NULL,
 z_ticks = NULL,
 show_plot = NA,
 return_plot = NA,
 save_plot = NA,
 save_param = list(),
 default_save_name = "spat3D"
)
```

## **Arguments**

```
gobject
                  giotto object
                  x-axis dimension name (default = 'sdimx')
sdimx
sdimy
                  y-axis dimension name (default = 'sdimy')
sdimz
                  z-axis dimension name (default = 'sdimy')
point_size
                  size of point (cell)
cell_color
                  color for cells (see details)
cell_color_code
                  named vector with colors
select_cell_groups
                  select subset of cells/clusters based on cell_color parameter
select_cells
                  select subset of cells based on cell IDs
show_other_cells
                  display not selected cells
other_cell_color
                  color of not selected cells
                  show underlying spatial network
show_network
network_color
                  color of spatial network
network_alpha
                  opacity of spatial network
spatial_network_name
                  name of spatial network to use
show_grid
                  show spatial grid
grid_color
                  color of spatial grid
```

288 spat\_fish\_func

grid\_alpha opacity of spatial grid

spatial\_grid\_name

name of spatial grid to use

title title of plot show\_legend show legend

axis\_scale the way to scale the axis

custom\_ratio customize the scale of the plot

 $x_{ticks}$  set the number of ticks on the x-axis  $y_{ticks}$  set the number of ticks on the y-axis  $z_{ticks}$  set the number of ticks on the z-axis

show\_plot show plot

return\_plot return ggplot object

save\_plot directly save the plot [boolean]

save\_param list of saving parameters, see showSaveParameters

default\_save\_name

default save name for saving, don't change, change save\_name in save\_param

#### **Details**

Description of parameters.

#### Value

ggplot

# See Also

Other spatial visualizations: spatPlot2D(), spatPlot()

#### **Examples**

spatPlot3D(gobject)

spat\_fish\_func
spat\_fish\_func

## Description

performs fisher exact test

```
spat_fish_func(gene, bin_matrix, spat_mat, calc_hub = F, hub_min_int = 3)
```

spat\_OR\_func 289

```
spat_OR_func
```

spat\_OR\_func

### **Description**

calculate odds-ratio

#### Usage

```
spat_OR_func(gene, bin_matrix, spat_mat, calc_hub = F, hub_min_int = 3)
```

```
specific Cell Cell communication Scores\\ specific Cell Cell communication Scores
```

# Description

Specific Cell-Cell communication scores based on spatial expression of interacting cells

### Usage

gene\_set\_1 first specific gene set from gene pairs
gene\_set\_2 second specific gene set from gene pairs

log2FC\_addendum

addendum to add when calculating log2FC

min\_observations

minimum number of interactions needed to be considered

adjust\_method which method to adjust p-values

adjust\_target adjust multiple hypotheses at the cell or gene level

verbose verbose

#### **Details**

Statistical framework to identify if pairs of genes (such as ligand-receptor combinations) are expressed at higher levels than expected based on a reshuffled null distribution of gene expression values in cells that are spatially in proximity to eachother.. More details will follow soon.

#### Value

Cell-Cell communication scores for gene pairs based on spatial interaction

### **Examples**

```
specificCellCellcommunicationScores(gobject)
```

### **Description**

Merge selected clusters based on pairwise correlation scores and size of cluster.

#### Usage

```
split_dendrogram_in_two(dend)
```

### Arguments

dend dendrogram object

# Value

list of two dendrograms and height of node

```
split_dendrogram_in_two(dend)
```

standardise\_giotto 291

```
standardise_giotto standardise_giotto
```

# Description

standardises a matrix

# Usage

```
standardise_giotto(x, center = TRUE, scale = TRUE)
```

# Arguments

```
x matrixcenter center datascale scale data
```

### Value

standardized matrix

```
stitchFieldCoordinates
```

stitchFieldCoordinates

# Description

Helper function to stitch field coordinates together to form one complete picture

```
stitchFieldCoordinates(
  location_file,
  offset_file,
  cumulate_offset_x = F,
  cumulate_offset_y = F,
  field_col = "Field of View",
  X_coord_col = "X",
  Y_coord_col = "Y",
  reverse_final_x = F,
  reverse_final_y = T
)
```

292 stitchTileCoordinates

#### **Arguments**

```
location_file location dataframe with X and Y coordinates
offset file
                  dataframe that describes the offset for each field (see details)
cumulate_offset_x
                  (boolean) Do the x-axis offset values need to be cumulated?
cumulate_offset_y
                  (boolean) Do the y-axis offset values need to be cumulated?
                  column that indicates the field within the location_file
field_col
X_coord_col
                  column that indicates the x coordinates
Y_coord_col
                  column that indicates the x coordinates
reverse_final_x
                  (boolean) Do the final x coordinates need to be reversed?
reverse_final_y
                  (boolean) Do the final y coordinates need to be reversed?
```

#### **Details**

Stitching of fields:

- 1. have cell locations: at least 3 columns: field, X, Y
- 2. create offset file: offset file has 3 columns: field, x\_offset, y\_offset
- 3. create new cell location file by stitching original cell locations with stitchFieldCoordinates
- 4. provide new cell location file to createGiottoObject

# Value

Updated location dataframe with new X ['X\_final'] and Y ['Y\_final'] coordinates

### **Examples**

```
stitchFieldCoordinates(gobject)
```

```
\verb|stitchTileCoordinates|| stitchTileCoordinates||
```

# Description

Helper function to stitch tile coordinates together to form one complete picture

# Usage

```
stitchTileCoordinates(location_file, Xtilespan, Ytilespan)
```

```
 \begin{array}{ll} \mbox{location\_file} & \mbox{location dataframe with $X$ and $Y$ coordinates} \\ \mbox{Xtilespan} & \mbox{numerical value specifying the width of each tile} \\ \mbox{Ytilespan} & \mbox{numerical value specifying the height of each tile} \\ \end{array}
```

subClusterCells 293

#### **Details**

•••

#### **Examples**

stitchTileCoordinates(gobject)

subClusterCells

subClusterCells

# Description

subcluster cells

# Usage

```
subClusterCells(
  gobject,
  name = "sub_clus",
  cluster_method = c("leiden", "louvain_community", "louvain_multinet"),
  cluster_column = NULL,
  selected_clusters = NULL,
 hvg_param = list(reverse_log_scale = T, difference_in_variance = 1, expression_values
    = "normalized"),
  hvg_min_perc_cells = 5,
  hvg_mean_expr_det = 1,
  use_all_genes_as_hvg = FALSE,
  min_nr_of_hvg = 5,
  pca_param = list(expression_values = "normalized", scale_unit = T),
  nn_param = list(dimensions_to_use = 1:20),
  k_neighbors = 10,
  resolution = 1,
  n_{iterations} = 1000,
  gamma = 1,
  omega = 1,
  python_path = NULL,
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  return_gobject = TRUE,
  verbose = T
)
```

```
gobject giotto object

name name for new clustering result

cluster_method clustering method to use

cluster_column cluster column to subcluster

selected_clusters

only do subclustering on these clusters
```

294 subClusterCells

hvg\_param parameters for calculateHVG

hvg\_min\_perc\_cells

threshold for detection in min percentage of cells

hvg\_mean\_expr\_det

threshold for mean expression level in cells with detection

use\_all\_genes\_as\_hvg

forces all genes to be HVG and to be used as input for PCA

min\_nr\_of\_hvg minimum number of HVG, or all genes will be used as input for PCA

pca\_param parameters for runPCA

nn\_param parameters for parameters for createNearestNetwork

 $k\_neighbors$  number of k for createNearestNetwork

resolution resolution

n\_iterations number of interations to run the Leiden algorithm.

gamma gamma omega omega

python\_path specify specific path to python if required

nn\_network\_to\_use

type of NN network to use (kNN vs sNN)

network\_name name of NN network to use

return\_gobject boolean: return giotto object (default = TRUE)

verbose verbose

#### **Details**

This function performs subclustering on selected clusters. The systematic steps are:

- 1. subset Giotto object
- 2. identify highly variable genes
- 3. run PCA
- 4. create nearest neighbouring network
- 5. do clustering

#### Value

giotto object with new subclusters appended to cell metadata

#### See Also

 ${\tt doLouvainCluster\_multinet}, {\tt doLouvainCluster\_community} \ and \ @see also \ {\tt doLeidenCluster\_community} \\$ 

#### **Examples**

subClusterCells(gobject)

subsetGiotto 295

subsetGiotto

subsetGiot to

# Description

subsets Giotto object including previous analyses.

# Usage

```
subsetGiotto(gobject, cell_ids = NULL, gene_ids = NULL, verbose = FALSE)
```

# Arguments

```
gobject giotto object
cell_ids cell IDs to keep
gene_ids gene IDs to keep
verbose be verbose
```

#### Value

giotto object

### **Examples**

```
subsetGiotto(gobject)
```

 ${\tt subsetGiottoLocs}$ 

subsetGiottoLocs

### **Description**

subsets Giotto object based on spatial locations

```
subsetGiottoLocs(
  gobject,
  x_max = NULL,
  x_min = NULL,
  y_max = NULL,
  y_min = NULL,
  z_max = NULL,
  z_min = NULL,
  return_gobject = T,
  verbose = FALSE
)
```

296 trendSceek

### **Arguments**

gobject	giotto object
x_max	maximum x-coordinate
x_min	minimum x-coordinate
y_max	maximum y-coordinate
y_min	minimum y-coordinate
z_max	maximum z-coordinate
z_min	minimum z-coordinate
return_gobject	return Giotto object

### **Details**

if return\_gobject = FALSE, then a filtered combined metadata data.table will be returned

### Value

giotto object

### **Examples**

```
subsetGiottoLocs(gobject)
```

trendSceek trendSceek

### **Description**

Compute spatial variable genes with trendsceek method

### Usage

```
trendSceek(
  gobject,
  expression_values = c("normalized", "raw"),
  subset_genes = NULL,
  nrand = 100,
  ncores = 8,
  ...
)
```

```
gobject Giotto object
expression_values
gene expression values to use
subset_genes subset of genes to run trendsceek on
nrand An integer specifying the number of random resamplings of the mark distribution as to create the null-distribution.

ncores An integer specifying the number of cores to be used by BiocParallel
... Additional parameters to the trendsceek_test function
```

updateGiottoImage 297

#### **Details**

This function is a wrapper for the trendsceek\_test method implemented in the trendsceek package

### Value

data.frame with trendsceek spatial genes results

#### **Examples**

```
trendSceek(gobject)
```

updateGiottoImage

updateGiottoImage

### **Description**

Updates the boundaries of a giotto image attached to a giotto object

### Usage

```
updateGiottoImage(
  gobject,
  image_name,
  xmax_adj = 0,
  xmin_adj = 0,
  ymax_adj = 0,
  ymin_adj = 0,
  return_gobject = TRUE
)
```

### **Arguments**

```
gobject giotto object
image_name spatial locations

xmax_adj adjustment of the maximum x-value to align the image

xmin_adj adjustment of the minimum x-value to align the image

ymax_adj adjustment of the maximum y-value to align the image

ymin_adj adjustment of the minimum y-value to align the image

return_gobject return a giotto object
```

#### Value

```
a giotto object or an updated giotto image if return_gobject = F
```

```
updateGiottoImage(gobject)
```

298 viewHMRFresults

viewHMRFresults

viewHMRFresults

# Description

View results from doHMRF.

# Usage

```
viewHMRFresults(
  gobject,
  HMRFoutput,
  k = NULL,
  betas_to_view = NULL,
  third_dim = NULL,
  ...
)
```

# **Arguments**

```
gobject giotto object
```

HMRF output from doHMRF k number of HMRF domains

betas\_to\_view results from different betas that you want to view

... paramters to visPlot()

# **Details**

Description ...

# Value

spatial plots with HMRF domains

### See Also

```
visPlot
```

```
viewHMRFresults(gobject)
```

viewHMRFresults2D 299

viewHMRFresults2D

viewHMRFresults2D

# Description

View results from doHMRF.

# Usage

```
viewHMRFresults2D(
  gobject,
  HMRFoutput,
  k = NULL,
  betas_to_view = NULL,
  third_dim = NULL,
  ...
)
```

# **Arguments**

```
gobject giotto object
```

HMRF output from doHMRF

k number of HMRF domains

betas\_to\_view results from different betas that you want to view

... paramters to visPlot()

# **Details**

Description ...

# Value

spatial plots with HMRF domains

### See Also

```
spatPlot2D
```

```
viewHMRFresults2D(gobject)
```

300 viewHMRFresults3D

viewHMRFresults3D

viewHMRFresults3D

# Description

View results from doHMRF.

# Usage

```
viewHMRFresults3D(
  gobject,
  HMRFoutput,
  k = NULL,
  betas_to_view = NULL,
  third_dim = NULL,
  ...
)
```

# **Arguments**

```
gobject giotto object
```

HMRF output from doHMRF k number of HMRF domains

betas\_to\_view results from different betas that you want to view

... paramters to visPlot()

# **Details**

Description ...

# Value

spatial plots with HMRF domains

### See Also

```
spatPlot3D
```

```
viewHMRFresults3D(gobject)
```

violinPlot 301

violinPlot

violinPlot

### **Description**

Creates violinplot for selected clusters

### Usage

```
violinPlot(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  genes,
  cluster_column,
  cluster_custom_order = NULL,
  color_violin = c("genes", "cluster"),
  cluster_color_code = NULL,
  strip_position = c("top", "right", "left", "bottom"),
  strip\_text = 7,
  axis_text_x_size = 10,
  axis_text_y_size = 6,
  show_plot = NA,
  return_plot = NA,
  save_plot = NA,
  save_param = list(),
  default_save_name = "violinPlot"
)
```

```
gobject
                  giotto object
expression_values
                  expression values to use
                  genes to plot
genes
cluster_column name of column to use for clusters
cluster_custom_order
                  custom order of clusters
color_violin
                  color violin according to genes or clusters
cluster_color_code
                  color code for clusters
strip_position position of gene labels
strip_text
                  size of strip text
\verb"axis_text_x_size"
                  size of x-axis text
axis_text_y_size
                  size of y-axis text
show_plot
                  show plot
return_plot
                  return ggplot object
```

302 visDimGenePlot

#### Value

ggplot

#### **Examples**

```
violinPlot(gobject)
```

visDimGenePlot

visDimGenePlot

### **Description**

Visualize cells and gene expression according to dimension reduction coordinates

```
visDimGenePlot(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  genes = NULL,
  dim_reduction_to_use = "umap",
  dim_reduction_name = "umap",
  dim1_to_use = 1,
  dim2_to_use = 2,
  dim3_to_use = NULL,
  show_NN_network = F,
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  network_color = "lightgray",
  edge_alpha = NULL,
  scale_alpha_with_expression = FALSE,
  point_size = 1,
  genes_high_color = NULL,
  genes_mid_color = "white",
  genes_low_color = "blue",
  point_border_col = "black",
  point_border_stroke = 0.1,
  midpoint = 0,
  cow_n_col = 2,
  cow_rel_h = 1,
  cow_rel_w = 1,
  cow_align = "h"
  show_legend = T,
  plot_method = c("ggplot", "plotly"),
  show_plots = F
)
```

visDimGenePlot 303

#### **Arguments**

gobject giotto object

expression\_values

gene expression values to use

genes genes to show

dim\_reduction\_to\_use

dimension reduction to use

dim\_reduction\_name

dimension reduction name

dim1\_to\_use dimension to use on x-axis dim2\_to\_use dimension to use on y-axis dim3\_to\_use dimension to use on z-axis

show\_NN\_network

show underlying NN network

nn\_network\_to\_use

type of NN network to use (kNN vs sNN)

 $network\_name$  name of NN network to use, if  $show\_NN\_network = TRUE$ 

edge\_alpha column to use for alpha of the edges

scale\_alpha\_with\_expression

scale expression with ggplot alpha parameter

point\_size size of point (cell)

point\_border\_col

color of border around points

point\_border\_stroke

stroke size of border around points

midpoint size of point (cell)

cow\_n\_col cowplot param: how many columns
cow\_rel\_h cowplot param: relative height
cow\_rel\_w cowplot param: relative width
cow\_align cowplot param: how to align

show\_legend show legend show\_plots show plots

### **Details**

Description of parameters.

#### Value

ggplot

# Examples

visDimGenePlot(gobject)

### **Description**

Visualize cells and gene expression according to dimension reduction coordinates

### Usage

```
visDimGenePlot_2D_ggplot(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  genes = NULL,
  dim_reduction_to_use = "umap",
  dim_reduction_name = "umap",
  dim1_to_use = 1,
  dim2\_to\_use = 2,
  show_NN_network = F,
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  network_color = "lightgray",
  edge_alpha = NULL,
  scale_alpha_with_expression = FALSE,
  point_size = 1,
  genes_high_color = "red",
  genes_mid_color = "white",
  genes_low_color = "blue",
  point_border_col = "black",
  point_border_stroke = 0.1,
  midpoint = 0,
  cow_n_col = 2,
  cow_rel_h = 1,
  cow_rel_w = 1,
  cow_align = "h",
  show_legend = T,
  show_plots = F
```

```
gobject giotto object
expression_values
gene expression values to use
genes genes to show
dim_reduction_to_use
dimension reduction to use
dim_reduction_name
dimension reduction name
dim1_to_use
dimension to use on x-axis
```

```
dim2_to_use
                 dimension to use on y-axis
show_NN_network
                 show underlying NN network
nn_network_to_use
                 type of NN network to use (kNN vs sNN)
                 name of NN network to use, if show_NN_network = TRUE
network_name
edge_alpha
                 column to use for alpha of the edges
scale_alpha_with_expression
                 scale expression with ggplot alpha parameter
                 size of point (cell)
point_size
point_border_col
                 color of border around points
point_border_stroke
                 stroke size of border around points
                 size of point (cell)
midpoint
cow_n_col
                 cowplot param: how many columns
                 cowplot param: relative height
cow_rel_h
                 cowplot param: relative width
cow_rel_w
cow_align
                 cowplot param: how to align
show_legend
                 show legend
show_plots
                 show plots
```

#### **Details**

Description of parameters.

### Value

ggplot

### **Examples**

visDimGenePlot\_2D\_ggplot(gobject)

### **Description**

Visualize cells and gene expression according to dimension reduction coordinates

#### Usage

```
visDimGenePlot_3D_plotly(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  genes = NULL,
  dim_reduction_to_use = "umap",
  dim_reduction_name = "umap",
  dim1_to_use = 1,
  dim2\_to\_use = 2,
  dim3_to_use = 3,
  show_NN_network = F,
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  network_color = "lightgray",
  edge_alpha = NULL,
  point_size = 1,
  genes_high_color = NULL,
  genes_mid_color = "white",
  genes_low_color = "blue",
  show_legend = T,
  show_plots = F
)
```

### **Arguments**

```
gobject
                 giotto object
expression_values
                 gene expression values to use
genes
                 genes to show
dim_reduction_to_use
                 dimension reduction to use
dim_reduction_name
                 dimension reduction name
dim1_to_use
                 dimension to use on x-axis
dim2_to_use
                 dimension to use on y-axis
dim3_to_use
                 dimension to use on z-axis
show_NN_network
                 show underlying NN network
nn_network_to_use
                 type of NN network to use (kNN vs sNN)
network_name
                 name of NN network to use, if show_NN_network = TRUE
edge_alpha
                 column to use for alpha of the edges
point_size
                 size of point (cell)
show_legend
                 show legend
                 show plots
show_plots
```

#### **Details**

Description of parameters.

visDimPlot 307

#### Value

ggplot

#### **Examples**

```
visDimGenePlot_3D_plotly(gobject)
```

visDimPlot

visDimPlot

#### **Description**

Visualize cells according to dimension reduction coordinates

```
visDimPlot(
  gobject,
  dim_reduction_to_use = "umap",
  dim_reduction_name = "umap",
  dim1_to_use = 1,
  dim2_to_use = 2,
  dim3_to_use = NULL,
  show_NN_network = F,
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  cell_color = NULL,
  color_as_factor = T,
  cell_color_code = NULL,
  select_cell_groups = NULL,
  select_cells = NULL,
  show_other_cells = T,
  other_cell_color = "lightgrey",
  other_point_size = 0.5,
  show_cluster_center = F,
  show_center_label = T,
  center_point_size = 4,
  center_point_border_col = "black",
  center_point_border_stroke = 0.1,
  label_size = 4,
  label_fontface = "bold",
  edge_alpha = NULL,
  point_size = 3,
  point_border_col = "black",
  point_border_stroke = 0.1,
  plot_method = c("ggplot", "plotly"),
  show_legend = T,
  show_plot = F,
  return_plot = TRUE,
  save_plot = F,
  save_dir = NULL,
```

308 visDimPlot

```
save_folder = NULL,
      save_name = NULL,
      save_format = NULL,
      show_saved_plot = F,
    )
Arguments
    gobject
                     giotto object
    dim_reduction_to_use
                     dimension reduction to use
    dim_reduction_name
                     dimension reduction name
    dim1_to_use
                     dimension to use on x-axis
    dim2_to_use
                     dimension to use on y-axis
                     dimension to use on z-axis
    dim3_to_use
    show_NN_network
                     show underlying NN network
    nn_network_to_use
                     type of NN network to use (kNN vs sNN)
                     name of NN network to use, if show_NN_network = TRUE
    network_name
    cell_color
                     color for cells (see details)
    color_as_factor
                     convert color column to factor
    cell_color_code
                     named vector with colors
    show_cluster_center
                     plot center of selected clusters
    show_center_label
                     plot label of selected clusters
    center_point_size
                     size of center points
    label_size
                     size of labels
    label_fontface font of labels
    edge_alpha
                     column to use for alpha of the edges
    point_size
                     size of point (cell)
    point_border_col
                     color of border around points
    point_border_stroke
                     stroke size of border around points
    show_legend
                     show legend
```

show\_plot

save\_plot
save\_dir

return\_plot

show plot

return ggplot object

directly save the plot [boolean]

directory to save the plot

visDimPlot\_2D\_ggplot

309

#### **Details**

Description of parameters.

#### Value

ggplot or plotly

### **Examples**

```
visDimPlot(gobject)
```

```
visDimPlot_2D_ggplot visDimPlot_2D_ggplot
```

### **Description**

Visualize cells according to dimension reduction coordinates

```
visDimPlot_2D_ggplot(
  gobject,
  dim_reduction_to_use = "umap",
  dim_reduction_name = "umap",
  dim1_to_use = 1,
  dim2_to_use = 2,
  show_NN_network = F,
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  cell_color = NULL,
  color_as_factor = T,
  cell_color_code = NULL,
  select_cell_groups = NULL,
  select_cells = NULL,
  show_other_cells = T,
  other_cell_color = "lightgrey",
  other_point_size = 0.5,
  show_cluster_center = F,
  show_center_label = T,
  center_point_size = 4,
  center_point_border_col = "black",
  center_point_border_stroke = 0.1,
  label_size = 4,
  label_fontface = "bold",
```

```
edge_alpha = NULL,
      point_size = 1,
      point_border_col = "black",
      point_border_stroke = 0.1,
      show_legend = T,
      show_plot = F,
      return_plot = TRUE,
      save_plot = F,
      save_dir = NULL,
      save_folder = NULL,
      save_name = NULL,
      save_format = NULL,
      show_saved_plot = F,
    )
Arguments
    gobject
                     giotto object
    dim_reduction_to_use
                     dimension reduction to use
    dim_reduction_name
                     dimension reduction name
    dim1_to_use
                     dimension to use on x-axis
    dim2_to_use
                     dimension to use on y-axis
    show_NN_network
                     show underlying NN network
    nn_network_to_use
                     type of NN network to use (kNN vs sNN)
    network_name
                     name of NN network to use, if show_NN_network = TRUE
                     color for cells (see details)
    cell_color
    color_as_factor
                     convert color column to factor
    cell_color_code
                     named vector with colors
    select_cell_groups
                     select subset of cells/clusters based on cell_color parameter
    select_cells
                     select subset of cells based on cell IDs
    show_other_cells
                     display not selected cells
    other_cell_color
                     color of not selected cells
    other_point_size
                     size of not selected cells
    show_cluster_center
                     plot center of selected clusters
    show_center_label
                     plot label of selected clusters
    center_point_size
```

size of center points

visDimPlot\_2D\_plotly 311

#### **Details**

Description of parameters.

#### Value

ggplot

### **Examples**

```
visDimPlot_2D_ggplot(gobject)
```

```
visDimPlot_2D_plotly visDimPlot_2D_plotly
```

### **Description**

Visualize cells according to dimension reduction coordinates

```
visDimPlot_2D_plotly(
  gobject,
  dim_reduction_to_use = "umap",
  dim_reduction_name = "umap",
  dim1_to_use = 1,
  dim2\_to\_use = 2,
  select_cell_groups = NULL,
  select_cells = NULL,
  show_other_cells = T,
  other_cell_color = "lightgrey",
  other_point_size = 0.5,
  show_NN_network = F,
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  color_as_factor = T,
  cell_color = NULL,
  cell_color_code = NULL,
  show_cluster_center = F,
  show_center_label = T,
```

```
center_point_size = 4,
label_size = 4,
edge_alpha = NULL,
point_size = 5
)
```

### **Arguments**

```
gobject
                 giotto object
dim_reduction_to_use
                 dimension reduction to use
dim_reduction_name
                 dimension reduction name
                 dimension to use on x-axis
dim1_to_use
dim2_to_use
                 dimension to use on y-axis
show_NN_network
                 show underlying NN network
nn_network_to_use
                 type of NN network to use (kNN vs sNN)
network_name
                 name of NN network to use, if show_NN_network = TRUE
color_as_factor
                 convert color column to factor
cell_color
                 color for cells (see details)
cell_color_code
                 named vector with colors
show_cluster_center
                 plot center of selected clusters
show_center_label
                 plot label of selected clusters
center_point_size
                 size of center points
label_size
                 size of labels
edge_alpha
                 column to use for alpha of the edges
point_size
                 size of point (cell)
```

#### **Details**

Description of parameters.

#### Value

plotly

```
visDimPlot_2D_plotly(gobject)
```

```
visDimPlot_3D_plotly
```

### **Description**

Visualize cells according to dimension reduction coordinates

### Usage

```
visDimPlot_3D_plotly(
  gobject,
  dim_reduction_to_use = "umap",
  dim_reduction_name = "umap",
  dim1_to_use = 1,
  dim2\_to\_use = 2,
  dim3_to_use = 3,
  select_cell_groups = NULL,
  select_cells = NULL,
  show_other_cells = T,
  other_cell_color = "lightgrey",
  other_point_size = 0.5,
  show_NN_network = F,
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  color_as_factor = T,
  cell_color = NULL,
  cell_color_code = NULL,
  show_cluster_center = F,
  show_center_label = T,
  center_point_size = 4,
  label_size = 4,
  edge_alpha = NULL,
  point_size = 1
```

```
giotto object
gobject
dim_reduction_to_use
                 dimension reduction to use
dim_reduction_name
                 dimension reduction name
dim1_to_use
                 dimension to use on x-axis
                 dimension to use on y-axis
dim2_to_use
dim3_to_use
                 dimension to use on z-axis
show_NN_network
                 show underlying NN network
nn_network_to_use
                 type of NN network to use (kNN vs sNN)
```

314 visForceLayoutPlot

```
name of NN network to use, if show_NN_network = TRUE
network_name
color_as_factor
                  convert color column to factor
cell_color
                  color for cells (see details)
cell_color_code
                  named vector with colors
show_cluster_center
                  plot center of selected clusters
show_center_label
                  plot label of selected clusters
center_point_size
                  size of center points
label_size
                  size of labels
                  column to use for alpha of the edges
edge_alpha
point_size
                  size of point (cell)
```

#### **Details**

Description of parameters.

#### Value

plotly

### **Examples**

```
visDimPlot_3D_plotly(gobject)
```

visForceLayoutPlot visForceLayoutPlot

# Description

Visualize cells according to forced layout algorithm coordinates

```
visForceLayoutPlot(
  gobject,
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  layout_name = "layout",
  dim1_to_use = 1,
  dim2_to_use = 2,
  show_NN_network = T,
  cell_color = NULL,
  color_as_factor = TRUE,
  cell_color_code = NULL,
  edge_alpha = NULL,
  point_size = 1,
```

visForceLayoutPlot 315

```
point_border_col = "black",
point_border_stroke = 0.1,
show_legend = T,
show_plot = F,
return_plot = TRUE,
save_plot = F,
save_dir = NULL,
save_folder = NULL,
save_format = NULL,
show_saved_plot = F,
...
)
```

```
gobject
                  giotto object
nn_network_to_use
                  type of NN network to use (kNN vs sNN)
network_name
                  NN network to use
                  name of layout to use
layout_name
dim1_to_use
                  dimension to use on x-axis
                  dimension to use on y-axis
dim2_to_use
show_NN_network
                  show underlying NN network
cell_color
                  color for cells (see details)
color_as_factor
                  convert color column to factor
cell_color_code
                  named vector with colors
                  column to use for alpha of the edges
edge_alpha
                  size of point (cell)
point_size
point_border_col
                  color of border around points
point_border_stroke
                  stroke size of border around points
                  show legend
show_legend
show_plot
                  show plot
return_plot
                  return ggplot object
                  directly save the plot [boolean]
save_plot
                  directory to save the plot
save_dir
save_folder
                  (optional) folder in directory to save the plot
                  name of plot
save_name
save_format
                  format of plot (e.g. tiff, png, pdf, ...)
show_saved_plot
                  load & display the saved plot
```

316 visGenePlot

#### **Details**

Description of parameters.

#### Value

ggplot

#### **Examples**

visForceLayoutPlot(gobject)

visGenePlot

visGenePlot

### **Description**

Visualize cells and gene expression according to spatial coordinates

```
visGenePlot(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  genes_high_color = NULL,
  genes_mid_color = "white",
  genes_low_color = "blue",
  show_network = F,
  network_color = NULL,
  spatial_network_name = "spatial_network",
  edge_alpha = NULL,
  show\_grid = F,
  grid_color = NULL,
  spatial_grid_name = "spatial_grid",
  midpoint = 0,
  scale_alpha_with_expression = FALSE,
  point_size = 1,
  point_border_col = "black",
  point_border_stroke = 0.1,
  show_legend = T,
  cow_n_col = 2,
  cow_rel_h = 1,
  cow_rel_w = 1,
  cow_align = "h",
  axis_scale = c("cube", "real", "custom"),
  custom_ratio = NULL,
  x_ticks = NULL,
  y_ticks = NULL,
  z_ticks = NULL,
  plot_method = c("ggplot", "plotly"),
  show_plots = F
```

visGenePlot 317

#### **Arguments**

gobject giotto object
expression\_values

gene expression values to use

genes genes to show

genes\_high\_color

color represents high gene expression

genes\_mid\_color

color represents middle gene expression

genes\_low\_color

color represents low gene expression

show\_network show underlying spatial network

network\_color color of spatial network

spatial\_network\_name

name of spatial network to use

show\_grid show spatial grid grid\_color color of spatial grid spatial\_grid\_name

name of spatial grid to use

midpoint expression midpoint
scale\_alpha\_with\_expression

scale expression with ggplot alpha parameter

point\_size size of point (cell)

point\_border\_col

color of border around points

point\_border\_stroke

stroke size of border around points

show\_legend show legend

cow\_n\_col cowplot param: how many columns cow\_rel\_h cowplot param: relative height cow\_rel\_w cowplot param: relative width cow\_align cowplot param: how to align three mode to adjust axis scale axis\_scale x\_ticks number of ticks on x axis number of ticks on y axis y\_ticks number of ticks on z axis z\_ticks plot\_method two methods of plot

show\_plots show plots

### **Details**

Description of parameters.

#### Value

ggplot or plotly

#### **Examples**

```
visGenePlot(gobject)
```

```
visGenePlot_2D_ggplot visGenePlot_2D_ggplot
```

### **Description**

Visualize cells and gene expression according to spatial coordinates

#### Usage

```
visGenePlot_2D_ggplot(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  genes,
  genes_high_color = "darkred",
  genes_mid_color = "white",
  genes_low_color = "darkblue",
  show_network = F,
  network_color = NULL,
  spatial_network_name = "spatial_network",
  edge_alpha = NULL,
  show\_grid = F,
  grid_color = NULL,
  spatial_grid_name = "spatial_grid",
  midpoint = 0,
  scale_alpha_with_expression = FALSE,
  point_size = 1,
  point_border_col = "black",
  point_border_stroke = 0.1,
  show_legend = T,
  cow_n_col = 2,
  cow_rel_h = 1,
  cow_rel_w = 1,
  cow_align = "h",
  show_plots = F
)
```

```
gobject giotto object
expression_values
gene expression values to use
genes genes to show
genes_high_color
color represents high gene expression
genes_mid_color
color represents middle gene expression
```

visGenePlot\_3D\_plotly

```
genes_low_color
```

color represents low gene expression

show\_network show underlying spatial network

network\_color color of spatial network

spatial\_network\_name

name of spatial network to use

show\_grid show spatial grid grid\_color color of spatial grid

spatial\_grid\_name

name of spatial grid to use

midpoint expression midpoint

 ${\tt scale\_alpha\_with\_expression}$ 

scale expression with ggplot alpha parameter

point\_size size of point (cell)

point\_border\_col

color of border around points

point\_border\_stroke

stroke size of border around points

show\_legend show legend

cow\_n\_colcowplot param: how many columnscow\_rel\_hcowplot param: relative heightcow\_rel\_wcowplot param: relative widthcow\_aligncowplot param: how to align

show\_plots show plots

### **Details**

Description of parameters.

# Value

ggplot

### **Examples**

visGenePlot\_2D\_ggplot(gobject)

 ${\tt visGenePlot\_3D\_plotly} \ \ {\it visGenePlot\_3D\_plotly}$ 

# Description

Visualize cells and gene expression according to spatial coordinates

#### Usage

network\_color

show\_grid

spatial\_network\_name

genes\_high\_color

genes\_mid\_color

genes\_low\_color

spatial\_grid\_name

point\_size show\_legend

axis\_scale

x\_ticks

y\_ticks

```
visGenePlot_3D_plotly(
     gobject,
      expression_values = c("normalized", "scaled", "custom"),
     genes,
      show_network = F,
     network_color = NULL,
      spatial_network_name = "spatial_network",
     edge_alpha = NULL,
      show\_grid = F,
     genes_high_color = NULL,
     genes_mid_color = "white",
     genes_low_color = "blue",
      spatial_grid_name = "spatial_grid",
     point_size = 1,
      show_legend = T,
     axis_scale = c("cube", "real", "custom"),
     custom_ratio = NULL,
     x_ticks = NULL,
     y_ticks = NULL,
     z_ticks = NULL,
      show_plots = F
Arguments
   gobject
                    giotto object
   expression_values
                    gene expression values to use
   genes
                    genes to show
                    show underlying spatial network
   show_network
```

color of spatial network

show spatial grid

name of spatial network to use

color represents high gene expression

color represents middle gene expression

color represents low gene expression

name of spatial grid to use

three mode to adjust axis scale

number of ticks on x axis number of ticks on y axis

size of point (cell)

show legend

visPlot 321

#### **Details**

Description of parameters.

#### Value

plotly

### **Examples**

```
visGenePlot_3D_plotly(gobject)
```

visPlot visPlot

# Description

Visualize cells according to spatial coordinates

```
visPlot(
  gobject,
  sdimx = NULL,
  sdimy = NULL,
  sdimz = NULL,
  point_size = 3,
  point_border_col = "black",
  point_border_stroke = 0.1,
  cell_color = NULL,
  cell_color_code = NULL,
  color_as_factor = T,
  select_cell_groups = NULL,
  select_cells = NULL,
  show_other_cells = T,
  other_cell_color = "lightgrey",
  show_network = F,
  network_color = NULL,
  network_alpha = 1,
  other_cell_alpha = 0.1,
  spatial_network_name = "spatial_network",
  show\_grid = F,
```

322 visPlot

```
grid_color = NULL,
      grid_alpha = 1,
      spatial_grid_name = "spatial_grid",
      coord_fix_ratio = 0.6,
      title = "",
      show_legend = T,
      axis_scale = c("cube", "real", "custom"),
      custom_ratio = NULL,
      x_{ticks} = NULL,
      y_ticks = NULL,
      z_ticks = NULL,
      plot_method = c("ggplot", "plotly"),
      show_plot = F,
      return_plot = TRUE,
      save_plot = F,
      save_dir = NULL,
      save_folder = NULL,
      save_name = NULL,
      save_format = NULL,
      show_saved_plot = F,
    )
Arguments
   gobject
                     giotto object
    sdimx
                     x-axis dimension name (default = 'sdimx')
    sdimy
                     y-axis dimension name (default = 'sdimy')
    sdimz
                     z-axis dimension name (default = 'sdimz')
   point_size
                     size of point (cell)
   point_border_col
                     color of border around points
   point_border_stroke
                     stroke size of border around points
    cell_color
                     color for cells (see details)
    cell_color_code
                     named vector with colors
    color_as_factor
                     convert color column to factor
    select_cell_groups
                     select subset of cells/clusters based on cell_color parameter
                     select subset of cells based on cell IDs
    select_cells
    show_other_cells
                     display not selected cells
   other_cell_color
                     color of not selected cells
    show_network
                     show underlying spatial network
```

color of spatial network

name of spatial network to use

network\_color

spatial\_network\_name

visPlot\_2D\_ggplot 323

```
show_grid
                  show spatial grid
grid_color
                  color of spatial grid
spatial_grid_name
                  name of spatial grid to use
coord_fix_ratio
                  fix ratio between x and y-axis
title
                  title of plot
show_legend
                  show legend
show_plot
                  show plot
return_plot
                  return ggplot object
save_plot
                  directly save the plot [boolean]
save_dir
                  directory to save the plot
                  (optional) folder in directory to save the plot
save_folder
                  name of plot
save_name
save_format
                  format of plot (e.g. tiff, png, pdf, ...)
show_saved_plot
                  load & display the saved plot
```

#### **Details**

Description of parameters.

#### Value

ggplot

# **Examples**

```
visPlot(gobject)
```

```
visPlot_2D_ggplot
visPlot_2D_ggplot
```

### **Description**

Visualize cells according to spatial coordinates

```
visPlot_2D_ggplot(
  gobject,
  sdimx = NULL,
  sdimy = NULL,
  point_size = 3,
  point_border_col = "black",
  point_border_stroke = 0.1,
  cell_color = NULL,
  cell_color_code = NULL,
```

324 visPlot\_2D\_ggplot

```
color_as_factor = T,
  select_cell_groups = NULL,
  select_cells = NULL,
  show_other_cells = T,
  other_cell_color = "lightgrey",
  show_network = F,
  network_color = NULL,
  network_alpha = 1,
  other_cells_alpha = 0.1,
  spatial_network_name = "spatial_network",
  show\_grid = F,
  grid_color = NULL,
  spatial_grid_name = "spatial_grid",
  coord_fix_ratio = 0.6,
  title = "",
  show_legend = T,
  axis_scale = c("cube", "real", "custom"),
  custom_ratio = NULL,
  x_{ticks} = NULL,
  y_ticks = NULL,
  z_ticks = NULL,
  show_plot = F,
  return_plot = TRUE,
  save_plot = F,
  save_dir = NULL,
  save_folder = NULL,
  save_name = NULL,
  save_format = NULL,
  show_saved_plot = F,
)
```

```
gobject
                  giotto object
sdimx
                  x-axis dimension name (default = 'sdimx')
sdimy
                  y-axis dimension name (default = 'sdimy')
point_size
                  size of point (cell)
point_border_col
                  color of border around points
point_border_stroke
                  stroke size of border around points
cell_color
                  color for cells (see details)
cell_color_code
                  named vector with colors
color_as_factor
                  convert color column to factor
select_cell_groups
                  select subset of cells/clusters based on cell_color parameter
                  select subset of cells based on cell IDs
select_cells
```

visPlot\_2D\_ggplot 325

show\_other\_cells

display not selected cells

other\_cell\_color

color of not selected cells

show\_network show underlying spatial network

network\_color color of spatial network

spatial\_network\_name

name of spatial network to use

show\_grid show spatial grid

grid\_color color of spatial grid

spatial\_grid\_name

name of spatial grid to use

coord\_fix\_ratio

fix ratio between x and y-axis

title title of plot

show\_legend show legend

show\_plot show plot

return\_plot return ggplot object

save\_plot directly save the plot [boolean]

save\_dir directory to save the plot

save\_folder (optional) folder in directory to save the plot

save\_name name of plot

save\_format format of plot (e.g. tiff, png, pdf, ...)

show\_saved\_plot

load & display the saved plot

# **Details**

Description of parameters.

#### Value

ggplot

#### **Examples**

visPlot\_2D\_ggplot(gobject)

326 visPlot\_2D\_plotly

```
visPlot_2D_plotly
```

## **Description**

Visualize cells according to spatial coordinates

#### Usage

```
visPlot_2D_plotly(
  gobject,
  sdimx = NULL,
  sdimy = NULL,
  point_size = 3,
  cell_color = NULL,
  cell_color_code = NULL,
  color_as_factor = T,
  select_cell_groups = NULL,
  select_cells = NULL,
  show_other_cells = T,
  other_cell_color = "lightgrey",
  other_point_size = 0.5,
  show_network = F,
  network_color = "lightgray",
  network_alpha = 1,
  other_cell_alpha = 0.5,
  spatial_network_name = "spatial_network",
  show\_grid = F,
  grid_color = NULL,
  grid_alpha = 1,
  spatial_grid_name = "spatial_grid",
  show_legend = T,
  axis_scale = c("cube", "real", "custom"),
  custom_ratio = NULL,
  x_ticks = NULL,
  y_ticks = NULL,
  show_plot = F
```

# **Arguments**

```
gobject giotto object

sdimx x-axis dimension name (default = 'sdimx')

sdimy y-axis dimension name (default = 'sdimy')

point_size size of point (cell)

cell_color color for cells (see details)

cell_color_code

named vector with colors

color_as_factor

convert color column to factor
```

visPlot\_3D\_plotly 327

```
select_cell_groups
                  select a subset of the groups from cell_color
                  show underlying spatial network
show_network
                  color of spatial network
network_color
spatial_network_name
                  name of spatial network to use
                  show spatial grid
show_grid
grid_color
                  color of spatial grid
                  alpha of spatial grid
grid_alpha
spatial_grid_name
                  name of spatial grid to use
                  show legend
show_legend
show_plot
                  show plot
```

#### **Details**

Description of parameters.

#### Value

plotly

# **Examples**

```
visPlot_2D_plotly(gobject)
```

```
visPlot_3D_plotly
```

# Description

Visualize cells according to spatial coordinates

```
visPlot_3D_plotly(
  gobject,
  sdimx = NULL,
  sdimy = NULL,
  sdimz = NULL,
  point_size = 3,
  cell_color = NULL,
  cell_color_code = NULL,
  select_cell_groups = NULL,
  select_cells = NULL,
  show_other_cells = T,
  other_cell_color = "lightgrey",
  other_point_size = 0.5,
  show_network = F,
```

328 visPlot\_3D\_plotly

```
network_color = NULL,
network_alpha = 1,
other_cell_alpha = 0.5,
spatial_network_name = "spatial_network",
spatial_grid_name = "spatial_grid",
title = "",
show_legend = T,
axis_scale = c("cube", "real", "custom"),
custom_ratio = NULL,
x_ticks = NULL,
y_ticks = NULL,
stow_plot = F
```

## **Arguments**

```
gobject
                  giotto object
sdimx
                  x-axis dimension name (default = 'sdimx')
sdimy
                  y-axis dimension name (default = 'sdimy')
sdimz
                  z-axis dimension name (default = 'sdimz')
point_size
                  size of point (cell)
cell_color
                  color for cells (see details)
cell_color_code
                  named vector with colors
select_cell_groups
                  select a subset of the groups from cell_color
                  show underlying spatial network
show_network
network_color
                  color of spatial network
spatial_network_name
                  name of spatial network to use
spatial_grid_name
                  name of spatial grid to use
                  title of plot
title
show_legend
                  show legend
show_plot
                  show plot
point_border_col
                  color of border around points
point_border_stroke
                  stroke size of border around points
color_as_factor
                  convert color column to factor
show_grid
                  show spatial grid
grid_color
                  color of spatial grid
coord_fix_ratio
                  fix ratio between x and y-axis
```

visSpatDimGenePlot 329

#### **Details**

Description of parameters.

#### Value

ggplot

#### **Examples**

```
visPlot_3D_plotly(gobject)
```

visSpatDimGenePlot

visSpatDimGenePlot

#### **Description**

integration of visSpatDimGenePlot\_2D(ggplot) and visSpatDimGenePlot\_3D(plotly)

```
visSpatDimGenePlot(
 gobject,
 plot_method = c("ggplot", "plotly"),
 expression_values = c("normalized", "scaled", "custom"),
 plot_alignment = c("horizontal", "vertical"),
 dim_reduction_to_use = "umap",
 dim_reduction_name = "umap",
 dim1_to_use = 1,
 dim2_to_use = 2,
 dim3_to_use = NULL,
  sdimx = NULL,
  sdimy = NULL,
  sdimz = NULL,
 genes,
 dim_point_border_col = "black",
 dim_point_border_stroke = 0.1,
  show_NN_network = F,
 nn_network_to_use = "sNN",
 network_name = "sNN.pca",
 edge_alpha_dim = NULL,
  scale_alpha_with_expression = FALSE,
 label_size = 16,
 genes_low_color = "blue",
 genes_mid_color = "white",
 genes_high_color = "red",
 dim_point_size = 3,
 nn_network_alpha = 0.5,
  show_spatial_network = F,
  spatial_network_name = "spatial_network",
 network_color = "lightgray",
  spatial_network_alpha = 0.5,
```

330 visSpatDimGenePlot

```
show_spatial_grid = F,
      spatial_grid_name = "spatial_grid",
      spatial_grid_color = NULL,
      spatial_grid_alpha = 0.5,
      spatial_point_size = 3,
      spatial_point_border_col = "black",
      spatial_point_border_stroke = 0.1,
      legend_text_size = 12,
      axis_scale = c("cube", "real", "custom"),
      custom_ratio = NULL,
     x_ticks = NULL,
     y_ticks = NULL,
     z_ticks = NULL,
     midpoint = 0,
     point_size = 1,
      cow_n_col = 2,
      cow_rel_h = 1,
     cow_rel_w = 1,
     cow_align = "h",
     show_legend = T,
      show_plots = F
   )
Arguments
   gobject
                    giotto object
   expression_values
                    gene expression values to use
   plot_alignment direction to align plot
   dim_reduction_to_use
                    dimension reduction to use
   dim_reduction_name
                    dimension reduction name
   dim1_to_use
                    dimension to use on x-axis
   dim2_to_use
                    dimension to use on y-axis
   dim3_to_use
                    dimension to use on z-axis
   sdimx
                    x-axis dimension name (default = 'sdimx')
   sdimy
                    y-axis dimension name (default = 'sdimy')
    sdimz
                    z-axis dimension name (default = 'sdimz')
   genes
                    genes to show
   dim_point_border_col
                    color of border around points
   dim_point_border_stroke
                    stroke size of border around points
   show_NN_network
                    show underlying NN network
   nn_network_to_use
                    type of NN network to use (kNN vs sNN)
                    name of NN network to use, if show_NN_network = TRUE
   network_name
```

edge\_alpha\_dim dim reduction plot: column to use for alpha of the edges scale\_alpha\_with\_expression scale expression with ggplot alpha parameter label\_size size for the label

genes\_low\_color

color to represent low expression of gene

genes\_high\_color

color to represent high expression of gene

dim\_point\_size dim reduction plot: point size

spatial\_network\_name

name of spatial network to use

spatial\_grid\_name

name of spatial grid to use

spatial\_point\_size

spatial plot: point size

spatial\_point\_border\_col

color of border around points

spatial\_point\_border\_stroke

stroke size of border around points

legend\_text\_size

the size of the text in legend

axis\_scale three modes to adjust axis scale ratio custom\_ratio set the axis scale ratio on custom

x\_ticks number of ticks on x axis y\_ticks number of ticks on y axis z\_ticks number of ticks on z axis

midpoint size of point (cell)
point\_size size of point (cell)

cow\_n\_col cowplot param: how many columns
cow\_rel\_h cowplot param: relative height
cow\_rel\_w cowplot param: relative width
cow\_align cowplot param: how to align

show\_legend show\_plot show plot

#### **Details**

Description of parameters.

#### Value

ggplot or plotly

## **Examples**

 $\verb|visSpatDimGenePlot(gobject)| \\$ 

visSpatDimGenePlot\_2D visSpatDimGenePlot\_2D

## **Description**

Visualize cells according to spatial AND dimension reduction coordinates in ggplot mode

## Usage

```
visSpatDimGenePlot_2D(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  plot_alignment = c("horizontal", "vertical"),
  genes,
  dim_reduction_to_use = "umap",
  dim_reduction_name = "umap",
  dim1_to_use = 1,
  dim2_to_use = 2,
  point_size = 1,
  dim_point_border_col = "black",
  dim_point_border_stroke = 0.1,
  show_NN_network = F,
  show_spatial_network = F,
  show_spatial_grid = F,
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  edge_alpha_dim = NULL,
  scale_alpha_with_expression = FALSE,
  spatial_network_name = "spatial_network",
  spatial_grid_name = "spatial_grid",
  spatial_point_size = 1,
  spatial_point_border_col = "black",
  spatial_point_border_stroke = 0.1,
  midpoint = 0,
  genes_high_color = "red",
  genes_mid_color = "white";
  genes_low_color = "blue",
  cow_n_col = 2,
  cow_rel_h = 1,
  cow_rel_w = 1,
  cow_align = "h",
  axis_scale = c("cube", "real", "custom"),
  custom_ratio = NULL,
  x_ticks = NULL,
  y_ticks = NULL,
  show_legend = T,
  show_plots = F
```

#### **Arguments**

gobject giotto object

expression\_values

gene expression values to use

plot\_alignment direction to align plot

genes genes to show

dim\_reduction\_to\_use

dimension reduction to use

dim\_reduction\_name

dimension reduction name

dim1\_to\_use dimension to use on x-axis dim2\_to\_use dimension to use on y-axis

point\_size size of point (cell)

dim\_point\_border\_col

color of border around points

dim\_point\_border\_stroke

stroke size of border around points

show\_NN\_network

show underlying NN network

nn\_network\_to\_use

type of NN network to use (kNN vs sNN)

 $network\_name \qquad name \ of \ NN \ network \ to \ use, \ if \ show\_NN\_network = TRUE$ 

 ${\tt edge\_alpha\_dim} \ \ dim \ reduction \ plot: \ column \ to \ use \ for \ alpha \ of \ the \ edges$ 

scale\_alpha\_with\_expression

scale expression with ggplot alpha parameter

spatial\_network\_name

name of spatial network to use

spatial\_grid\_name

name of spatial grid to use

spatial\_point\_size

spatial plot: point size

spatial\_point\_border\_col

color of border around points

 ${\tt spatial\_point\_border\_stroke}$ 

stroke size of border around points

midpoint size of point (cell)

cow\_n\_col cowplot param: how many columns

cow\_rel\_h cowplot param: relative height cow\_rel\_w cowplot param: relative width cow\_align cowplot param: how to align

show\_legend show legend

dim\_point\_size dim reduction plot: point size

show\_plot show plot

#### **Details**

Description of parameters.

#### Value

ggplot

#### **Examples**

```
visSpatDimGenePlot_2D(gobject)
```

```
visSpatDimGenePlot_3D visSpatDimGenePlot_3D
```

## **Description**

Visualize cells according to spatial AND dimension reduction coordinates in plotly mode

```
visSpatDimGenePlot_3D(
  gobject,
  expression_values = c("normalized", "scaled", "custom"),
  plot_alignment = c("horizontal", "vertical"),
  dim_reduction_to_use = "umap",
  dim_reduction_name = "umap",
  dim1_to_use = 1,
  dim2\_to\_use = 2,
  dim3_to_use = NULL,
  sdimx = NULL,
  sdimy = NULL,
  sdimz = NULL,
  genes,
  show_NN_network = F,
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  label_size = 16,
  genes_low_color = "blue",
  genes_mid_color = "white",
  genes_high_color = "red",
  dim_point_size = 3,
  nn_network_alpha = 0.5,
  show\_spatial\_network = F,
  spatial_network_name = "spatial_network",
  network_color = "lightgray",
  spatial_network_alpha = 0.5,
  show_spatial_grid = F,
  spatial_grid_name = "spatial_grid",
  spatial_grid_color = NULL,
  spatial_grid_alpha = 0.5,
  spatial_point_size = 3,
  legend_text_size = 12,
  axis_scale = c("cube", "real", "custom"),
  custom_ratio = NULL,
  x_{ticks} = NULL,
```

y\_ticks = NULL, z\_ticks = NULL

```
Arguments
   gobject
                     giotto object
    plot_alignment direction to align plot
   dim_reduction_to_use
                     dimension reduction to use
    dim_reduction_name
                     dimension reduction name
                     dimension to use on x-axis
   dim1_to_use
   dim2_to_use
                     dimension to use on y-axis
    dim3_to_use
                     dimension to use on z-axis
    show_NN_network
                     show underlying NN network
    nn_network_to_use
                     type of NN network to use (kNN vs sNN)
                     name of NN network to use, if show_NN_network = TRUE
   network_name
    genes_low_color
                     color represent high gene expression (see details)
    genes_high_color
                     color represent high gene expression (see details)
    nn_network_alpha
                     column to use for alpha of the edges
    show_spatial_network
                     show spatial network
    spatial_network_name
                     name of spatial network to use
    network_color color of spatial/nn network
    spatial_network_alpha
                     alpha of spatial network
    show_spatial_grid
                     show spatial grid
    spatial_grid_name
                     name of spatial grid to use
    spatial_grid_color
                     color of spatial grid
    spatial_grid_alpha
                     alpha of spatial grid
    legend_text_size
                     text size of legend
    show_legend
                     show legend
```

## **Details**

show\_plot

Description of parameters.

show plot

visSpatDimPlot

#### Value

plotly

#### **Examples**

```
visSpatDimPlot_3D(gobject)
```

visSpatDimPlot

visSpatDimPlot

## **Description**

integration of visSpatDimPlot\_2D and visSpatDimPlot\_3D

```
visSpatDimPlot(
  gobject,
  plot_method = c("ggplot", "plotly"),
  plot_alignment = NULL,
  dim_reduction_to_use = "umap",
  dim_reduction_name = "umap",
  dim1_to_use = 1,
  dim2\_to\_use = 2,
  dim3_to_use = NULL,
  sdimx = NULL,
  sdimy = NULL,
  sdimz = NULL,
  show_NN_network = F,
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  show_cluster_center = F,
  show_center_label = T,
  center_point_size = 4,
  label_size = NULL,
  label_fontface = "bold",
  cell_color = NULL,
  color_as_factor = T,
  cell_color_code = NULL,
  select_cell_groups = NULL,
  select_cells = NULL,
  show_other_cells = T,
  other_cell_color = "lightgrey",
  dim_point_size = 3,
  dim_point_border_col = "black",
  dim_point_border_stroke = 0.1,
  nn_network_alpha = NULL,
  show_spatial_network = F,
  spatial_network_name = "spatial_network",
  network_color = "lightgray",
  spatial_network_alpha = 0.5,
```

visSpatDimPlot 337

```
show_spatial_grid = F,
      spatial_grid_name = "spatial_grid",
      spatial_grid_color = NULL,
      spatial_grid_alpha = 0.5,
      spatial_point_size = 3,
      legend_text_size = 12,
      spatial_point_border_col = "black",
      spatial_point_border_stroke = 0.1,
      show_legend = T,
      axis_scale = c("cube", "real", "custom"),
      custom_ratio = NULL,
      x_ticks = NULL,
      y_ticks = NULL,
      z_ticks = NULL,
      show_plot = F
Arguments
    gobject
                     giotto object
    plot_alignment direction to align plot
    dim_reduction_to_use
                     dimension reduction to use
    dim_reduction_name
                     dimension reduction name
    dim1_to_use
                     dimension to use on x-axis
    dim2_to_use
                     dimension to use on y-axis
    dim3_to_use
                     dimension to use on z-axis
    show_NN_network
                     show underlying NN network
    nn_network_to_use
                     type of NN network to use (kNN vs sNN)
                     name of NN network to use, if show_NN_network = TRUE
    network_name
    cell_color
                     color for cells (see details)
    color_as_factor
                     convert color column to factor
    cell_color_code
                     named vector with colors
    select_cell_groups
                     select subset of cells/clusters based on cell_color parameter
                     select subset of cells based on cell IDs
    select_cells
    show_other_cells
                     display not selected cells
    other_cell_color
                     color of not selected cells
    nn_network_alpha
                     column to use for alpha of the edges
    show\_spatial\_network
                     show spatial network
```

338 visSpatDimPlot\_2D

```
spatial_network_name
                  name of spatial network to use
spatial_network_alpha
                  alpha of spatial network
show_spatial_grid
                  show spatial grid
spatial_grid_name
                  name of spatial grid to use
spatial\_grid\_color
                  color of spatial grid
spatial_grid_alpha
                  alpha of spatial grid
legend_text_size
                  text size of legend
show_legend
                  show legend
show_plot
                  show plot
plot_mode
                  choose the mode to draw plot: ggplot or plotly
spatial_network_color
                  color of spatial network
```

#### **Details**

Description of parameters.

#### Value

ggplot or plotly

#### **Examples**

```
visSpatDimPlot(gobject)
```

visSpatDimPlot\_2D
visSpatDimPlot\_2D

## **Description**

Visualize cells according to spatial AND dimension reduction coordinates in ggplot2 mode

```
visSpatDimPlot_2D(
  gobject,
  plot_alignment = c("vertical", "horizontal"),
  dim_reduction_to_use = "umap",
  dim_reduction_name = "umap",
  dim1_to_use = 1,
  dim2_to_use = 2,
  sdimx = NULL,
  sdimy = NULL,
```

visSpatDimPlot\_2D

 $show_NN_network = F,$ 

339

```
nn_network_to_use = "sNN",
     network_name = "sNN.pca",
      show\_cluster\_center = F,
      show_center_label = T,
      center_point_size = 4,
      label_size = 4,
      label_fontface = "bold",
      cell_color = NULL,
      color_as_factor = T,
      cell_color_code = NULL,
      select_cell_groups = NULL,
      select_cells = NULL,
      show_other_cells = T,
     other_cell_color = "lightgrey",
      dim_plot_mode = NULL,
     dim_point_size = 1,
     dim_point_border_col = "black",
     dim_point_border_stroke = 0.1,
     nn_network_alpha = 0.05,
      show_spatial_network = F,
      spatial_network_name = "spatial_network",
      spatial_network_color = NULL,
      show_spatial_grid = F,
      spatial_grid_name = "spatial_grid",
      spatial_grid_color = NULL,
      spatial_point_size = 1,
      spatial_point_border_col = "black",
      spatial_point_border_stroke = 0.1,
      show_legend = T,
      show_plot = F,
     plot_method = "ggplot"
Arguments
                    giotto object
   gobject
   plot_alignment direction to align plot
   dim_reduction_to_use
                    dimension reduction to use
   dim_reduction_name
                    dimension reduction name
   dim1_to_use
                    dimension to use on x-axis
                    dimension to use on y-axis
   dim2_to_use
   show_NN_network
                    show underlying NN network
   nn_network_to_use
```

type of NN network to use (kNN vs sNN)

color for cells (see details)

network\_name
cell\_color

name of NN network to use, if show\_NN\_network = TRUE

340 visSpatDimPlot\_2D

```
color_as_factor
                  convert color column to factor
cell_color_code
                  named vector with colors
select_cell_groups
                  select subset of cells/clusters based on cell_color parameter
                  select subset of cells based on cell IDs
select_cells
show_other_cells
                  display not selected cells
other_cell_color
                  color of not selected cells
nn_network_alpha
                  column to use for alpha of the edges
show_spatial_network
                  show spatial network
spatial_network_name
                  name of spatial network to use
spatial_network_color
                  color of spatial network
show_spatial_grid
                  show spatial grid
spatial_grid_name
                  name of spatial grid to use
spatial_grid_color
                  color of spatial grid
show_legend
                  show legend
show_plot
                  show plot
return_plot
                  return ggplot object
                  directly save the plot [boolean]
save_plot
save_dir
                  directory to save the plot
                  (optional) folder in directory to save the plot
save_folder
                  name of plot
save_name
save_format
                  format of plot (e.g. tiff, png, pdf, ...)
show_saved_plot
                  load & display the saved plot
```

#### **Details**

Description of parameters.

#### Value

ggplot

## **Examples**

```
visSpatDimPlot_2D(gobject)
```

visSpatDimPlot\_3D 341

visSpatDimPlot\_3D

visSpatDimPlot\_3D

# Description

Visualize cells according to spatial AND dimension reduction coordinates in plotly mode

## Usage

```
visSpatDimPlot_3D(
  gobject,
  plot_alignment = c("horizontal", "vertical"),
  dim_reduction_to_use = "umap",
  dim_reduction_name = "umap",
  dim1_to_use = 1,
  dim2\_to\_use = 2,
  dim3_to_use = NULL,
  sdimx = NULL,
  sdimy = NULL,
  sdimz = NULL,
  show_NN_network = F,
  nn_network_to_use = "sNN",
  network_name = "sNN.pca",
  show_cluster_center = F,
  show_center_label = T,
  center_point_size = 4,
  label_size = 16,
  cell_color = NULL,
  color_as_factor = T,
  cell_color_code = NULL,
  dim_point_size = 3,
  nn_network_alpha = 0.5,
  show_spatial_network = F,
  spatial_network_name = "spatial_network",
  network_color = "lightgray",
  spatial_network_alpha = 0.5,
  show_spatial_grid = F,
  spatial_grid_name = "spatial_grid",
  spatial_grid_color = NULL,
  spatial_grid_alpha = 0.5,
  spatial_point_size = 3,
  axis_scale = c("cube", "real", "custom"),
  custom_ratio = NULL,
  x_ticks = NULL,
  y_ticks = NULL,
  z_ticks = NULL,
  legend_text_size = 12
```

#### **Arguments**

gobject giotto object

342 visSpatDimPlot\_3D

```
plot_alignment direction to align plot
dim_reduction_to_use
                 dimension reduction to use
dim_reduction_name
                 dimension reduction name
                 dimension to use on x-axis
dim1_to_use
                 dimension to use on y-axis
dim2_to_use
dim3_to_use
                 dimension to use on z-axis
show_NN_network
                 show underlying NN network
nn_network_to_use
                 type of NN network to use (kNN vs sNN)
                 name of NN network to use, if show_NN_network = TRUE
network_name
cell_color
                 color for cells (see details)
color_as_factor
                 convert color column to factor
cell_color_code
                 named vector with colors
nn_network_alpha
                 column to use for alpha of the edges
show_spatial_network
                 show spatial network
spatial_network_name
                 name of spatial network to use
spatial_network_alpha
                 alpha of spatial network
show_spatial_grid
                 show spatial grid
spatial_grid_name
                 name of spatial grid to use
spatial_grid_color
                 color of spatial grid
spatial_grid_alpha
                 alpha of spatial grid
legend_text_size
                 text size of legend
spatial_network_color
                 color of spatial network
                 show legend
show_legend
                 show plot
show_plot
```

## **Details**

Description of parameters.

#### Value

plotly

writeHMRFresults 343

## **Examples**

```
visSpatDimPlot_3D(gobject)
```

writeHMRFresults

writeHMRFresults

#### **Description**

write results from doHMRF to a data.table.

#### Usage

```
writeHMRFresults(
  gobject,
  HMRFoutput,
  k = NULL,
  betas_to_view = NULL,
  print_command = F
)
```

# Arguments

gobject giotto object

HMRF output HMRF output from doHMRF

k k to write results for

betas\_to\_view results from different betas that you want to view

print\_command see the python command

# Value

data.table with HMRF results for each b and the selected k

# **Examples**

```
writeHMRFresults(gobject)
```

## **Description**

write out factor-like annotation data from a giotto object for the Viewer

```
write_giotto_viewer_annotation(
  annotation,
  annot_name = "test",
  output_directory = getwd()
)
```

#### **Arguments**

```
annotation annotation from the data.table from giotto object
annot_name name of the annotation
output_directory
directory where to save the files
```

#### Value

write a .txt and .annot file for the selection annotation

#### **Description**

write out dimensional reduction data from a giotto object for the Viewer

## Usage

```
write_giotto_viewer_dim_reduction(
  dim_reduction_cell,
  dim_red = NULL,
  dim_red_name = NULL,
  dim_red_rounding = NULL,
  dim_red_rescale = c(-20, 20),
  output_directory = getwd()
)
```

## **Arguments**

```
dim_reduction_cell

dimension reduction slot from giotto object

dim_red high level name of dimension reduction

dim_red_name specific name of dimension reduction to use

dim_red_rounding

numerical indicating how to round the coordinates

dim_red_rescale

numericals to rescale the coordinates

output_directory

directory where to save the files
```

#### Value

write a .txt and .annot file for the selection annotation

```
write\_giotto\_viewer\_numeric\_annotation \\ write\_giotto\_viewer\_numeric\_annotation
```

# Description

write out numeric annotation data from a giotto object for the Viewer

# Usage

```
write_giotto_viewer_numeric_annotation(
  annotation,
  annot_name = "test",
  output_directory = getwd()
)
```

# **Arguments**

```
annotation annotation from the data.table from giotto object
annot_name name of the annotation
output_directory
directory where to save the files
```

# Value

write a .txt and .annot file for the selection annotation

# Index

*Topic <b>giotto</b> ,	cellProximityHeatmap, 30
giotto-class, 154	cellProximityNetwork, 31
print.giotto, 210	cellProximityNetwork, 31
show, giotto-method, 225	cellProximitySpatrlot2D, 34, 34
*Topic <b>giotto</b>	
createGiottoObject, 59	cellProximitySpatPlot3D, 34, 36
*Topic <b>object</b>	cellProximityVisPlot, 39
giotto-class, 154	cellProximityVisPlot_2D_ggplot, 41
print.giotto, 210	cellProximityVisPlot_2D_plotly, 43
show, giotto-method, 225	cellProximityVisPlot_3D_plotly, 44
Show, grotto-method, 225	changeGiottoInstructions, 46
addCellIntMetadata,8	changeImageBg, 47
addCellMetadata, 9, 60	cluster_walktrap, 121
addCellStatistics, 10, 16	clusterCells, 47
addGeneMetadata, 11, 60	clusterSpatialCorGenes, 50
addGenesPerc, 11	combCCcom, 51, 196, 198
addGeneStatistics, 12, 16	combineCellProximityGenes, 51
addGiottoImage, 13	<pre>combineCellProximityGenes_per_interaction,</pre>
addGiottoImage, 13 addGiottoImageToSpatPlot, 13	53
addHMRF, 14	combineCPG, 53
addNetworkLayout, 15	combineMetadata, 54
addStatistics, 16	<pre>convert_mgImage_to_array_DT, 55</pre>
addistatistics, 10 adjustGiottoMatrix, 16	<pre>convertEnsemblToGeneSymbol, 55</pre>
all_plots_save_function, 17, 26, 29, 31,	<pre>create_average_detection_DT, 73</pre>
32, 34, 36, 38, 129, 132, 155, 158,	create_average_DT, 74
171–173, 175–178, 180, 183, 184,	<pre>create_cell_type_random_cell_IDs, 74</pre>
197–199, 212, 230, 231, 233, 235,	<pre>create_crossSection_object, 75</pre>
278–280	create_delaunayNetwork2D,76
annotate_spatlocs_with_spatgrid_2D, 21	<pre>create_delaunayNetwork3D,77</pre>
annotate_spatiocs_with_spatgrid_3D, 22	<pre>create_delaunayNetwork_deldir,77</pre>
annotateGiotto, 19	<pre>create_delaunayNetwork_geometry, 78</pre>
annotateSpatialGrid, 20	<pre>create_delaunayNetwork_geometry_3D, 78</pre>
annotateSpatialNetwork, 20	<pre>create_delaunayNetwork_RTriangle, 79</pre>
average_gene_gene_expression_in_groups,	<pre>create_genes_to_use_matrix, 79</pre>
22	create_KNNnetwork_dbscan, 80
22	create_screeplot, 80
binSpect, 23	createCrossSection, 56
5115peet, 25	createGiottoImage, 13, 57
calculate_distance_and_weight, 28	createGiottoInstructions, 58, 60, 61
calculateHVG, 25, 216, 219, 221	createGiottoObject, 59, 292
calculateMetaTable, 26	createGiottoVisiumObject, 61
calculateMetaTableCells, 27	createHeatmap_DT, 62
cellProximityBarplot, 28	createMetagenes, 63
cellProximityEnrichment, 29	createNearestNetwork, 64

INDEX 347

createSpatialDelaunayNetwork, 66, 200	fDataDT, 127
createSpatialEnrich, 67, 125	filterCellProximityGenes, 128
createSpatialGrid, 68	filterCombinations, 129, 132
createSpatialGrid_2D, 69	filterCPG, 130
createSpatialGrid_3D,70	filterDistributions, 131
createSpatialKNNnetwork, 71	filterGiotto, 132
createSpatialNetwork, 72, 86	find_grid_2D,147
crossSectionGenePlot, 81	find_grid_3D,147
crossSectionGenePlot3D, 82	find_grid_x, 148
crossSectionPlot, 83, 83	find_grid_y,148
crossSectionPlot3D, 84	find_grid_z,148
	findCellProximityGenes, 133
decide_cluster_order, 85	<pre>findCellProximityGenes_per_interaction</pre>
delaunayn, 67	135
deldir,67	findCPG, 135
detectSpatialCorGenes, 86	findGiniMarkers, 137, 139, 141
detectSpatialPatterns, 87	<pre>findGiniMarkers_one_vs_all, 138, 142</pre>
dimCellPlot, 88, 92	findMarkers, 140, <i>146</i>
dimCellPlot2D, 88, 89, 90	findMarkers_one_vs_all, 141
dimGenePlot, 93, 96, 98	findMastMarkers, <i>141</i> , 143, <i>144</i>
dimGenePlot2D, 93, 94, 94, 98	findMastMarkers_one_vs_all, 142, 144
dimGenePlot3D, 94, 96, 96	findNetworkNeighbors, 145
dimPlot, 98, 103, 105, 192, 194, 196, 201,	findScranMarkers, 141, 145, 147
203, 205, 206, 208, 210	findScranMarkers_one_vs_all, 142, 146
dimPlot2D, 98–100, 100, 105, 191–194, 196,	FSV_show, 149
200–203, 205–208, 210	
dimPlot3D, 92, 99, 100, 103, 103, 192,	general_save_function, $18$
194–196, 201, 203–206, 208–210	get10Xmatrix, 150
do_permuttest (do_permuttest_random),	get_os, 153
123	getClusterSimilarity, 150
do_permuttest_random, 123	getDendrogramSplits, 151
doHclust, 50, 105	<pre>getDistinctColors, 152</pre>
doHMRF, 106	<pre>getGiottoImage, 153</pre>
doKmeans, 50, 108	giotto (giotto-class), 154
doLeidenCluster, 50, 109, 112, 294	giotto-class, 154
doLeidenSubCluster, 110	glouvain_ml, <i>115</i>
doLouvainCluster, 50, 112	
doLouvainCluster_community, 50, 113, 113,	hclust, <i>106</i>
117, 119, 294	Heatmap, <i>155</i>
doLouvainCluster_multinet, 50, 113, 114,	heatmSpatialCorGenes, 154
117, 120, 294	hyperGeometricEnrich, 68, 156
doLouvainSubCluster, 115	<pre>insertCrossSectionGenePlot3D, 157</pre>
<pre>doLouvainSubCluster_community, 117</pre>	insertCrossSectionSpatPlot3D, 158
<pre>doLouvainSubCluster_multinet, 119</pre>	inscrete observations patricists, 150
doRandomWalkCluster, 50, 121	jackstrawPlot, 159
doSNNCluster, 50, 122	3
DT_removeNA, 123	kmeans, <i>109</i>
	kmeans_binarize, 161
estimateImageBg, 124	knn, 65
exportGiottoViewer, 124	
exprCellCellcom, 125, 171, 172	layout_with_drl, 15
extractNearestNetwork, 127	loadHMRF, 161

348 INDEX

make_simulated_network, 163	plotUMAP_2D, 100, 103, 105, 192, 194, 196,
makeSignMatrixPAGE, 162, 169	201, 203, 205, 206, 207, 210
makeSignMatrixRank, 163, 211	plotUMAP_3D, 100, 103, 105, 192, 194, 196,
mergeClusters, 164	201, 203, 205, 206, 208, 209
my_arowMeans, 165	prcomp_irlba, 216
my_growMeans, 165	print.giotto, 210
my_rowMeans, 165	
	rank_binarize, 212
nnDT_to_kNN, 166	rankEnrich, 68, 163, 210
node_clusters, 166	rankSpatialCorGroups, 211
normalizeGiotto, 167	readExprMatrix,212
	readGiottoInstructions, 213
PAGEEnrich, 68, 162, 168	removeBatchEffect, 17
PCA, 216	removeCellAnnotation, 214
pca_giotto, 169	removeGeneAnnotation, 214
pDataDT, 20, 170	replaceGiottoInstructions, 215
permutationPA, <i>160</i> , <i>237</i>	Rtsne, <i>219</i>
plotCCcomDotplot, 170	runPCA, 215, 223
plotCCcomHeatmap, 171	runPCA_factominer, 217
plotCellProximityGenes, 172	<pre>runPCA_prcomp_irlba, 217</pre>
plotCombineCCcom, 174	runtSNE, 218
plotCombineCellCellCommunication, 175	runUMAP, 220
plotCombineCellProximityGenes, 176	
plotCombineCPG, 178	screePlot, 222
plotCPG, 179	select_expression_values, 224
plotGiottoImage, 180	select_spatialNetwork, 224
plotHeatmap, 181	selectPatternGenes, 223
plotICG, 183	<pre>set_giotto_python_path, 225</pre>
plotInteractionChangedGenes, 184	show, giotto-method, 225
plotly_axis_scale_2D, 185	showClusterDendrogram, 225
plotly_axis_scale_3D, 185	showClusterHeatmap, 226
plotly_grid, 186	showGiottoImageNames, 153, 180, 228
plotly_network, 187	showGiottoInstructions, 228
plotMetaDataCellsHeatmap, 187, 191	showGrids, <i>20</i> , 229
plotMetaDataHeatmap, 189, 189	showNetworks, 229
plotPCA, 100, 103, 105, 191, 194, 196, 201,	showPattern, 230
203, 205, 206, 208, 210	showPattern2D, <i>230</i> , 231
plotPCA_2D, 100, 103, 105, 192, 193, 196,	showPattern3D, 232
201, 203, 205, 206, 208, 210	showPatternGenes, 233
plotPCA_3D, 100, 103, 105, 192, 194, 195,	showProcessingSteps, 234
201, 203, 205, 206, 208, 210	showSaveParameters, 89, 92, 93, 96, 98, 99,
plotRankSpatvsExpr, 196	102, 104, 182, 188, 190, 192, 194,
plotRecovery, 197	195, 201, 203, 204, 206, 208, 209,
plotRecovery_sub, 198	226, 227, 235, 241, 244, 247, 251,
plotStatDelaunayNetwork, 199	253, 256, 259, 261, 266, 269, 270,
plotTSNE, 100, 103, 105, 192, 194, 196, 200,	273, 275, 282, 286, 288, 302
203, 205, 206, 208, 210	showSpatialCorGenes, 87, 235
plotTSNE_2D, 100, 103, 105, 192, 194, 196,	signPCA, 236
201, 202, 205, 206, 208, 210	silhouetteRank, 238
plotTSNE_3D, 100, 103, 105, 192, 194, 196,	sNN, 65
201, 203, 204, 206, 208, 210	sNNclust, <i>122</i>
plotUMAP, 100, 103, 105, 192, 194, 196, 201,	spat_fish_func, 288
203, 205, 205, 208, 210	spat_OR_func, 289
, , ,,,	

INDEX 349

spatCellCellcom, <i>171</i> , <i>172</i> , 239	visGenePlot_2D_ggplot, 318
spatCellPlot, 240, 245	visGenePlot_3D_plotly, 319
spatCellPlot2D, 240, 241, 242	visPlot, 298, 321
spatDimCellPlot, 245, 252	visPlot_2D_ggplot, 323
spatDimCellPlot2D, 245, 247, 247	visPlot_2D_plotly, 326
spatDimGenePlot, 252, 257, 259	visPlot_3D_plotly, 327
spatDimGenePlot2D, 252, 253, 254, 259	visSpatDimGenePlot, 329
spatDimGenePlot3D, 253, 257, 257	visSpatDimGenePlot_2D, 332
spatDimPlot, 259, 266, 269	visSpatDimGenePlot_3D, 334
spatDimPlot2D, 260, 261, 262, 269	visSpatDimPlot, 336
spatDimPlot3D, 261, 266, 266	visSpatDimPlot_2D, 338
spatGenePlot, 269, 273, 275	visSpatDimPlot_3D, 341
spatGenePlot2D, <i>81</i> , <i>270</i> , <i>271</i> , <i>271</i> , <i>275</i>	V135patb1iii 10t_5b, 541
	<pre>write_giotto_viewer_annotation, 343</pre>
spatGenePlot3D, 81, 271, 273, 274	write_giotto_viewer_dim_reduction, 344
spatialAEH, 276	write_giotto_viewer_numeric_annotation,
SpatialDE, 276	345
spatialDE, 277	writeHMRFresults, 343
spatNetwDistributions, 278	wiltering results, 545
spatNetwDistributionsDistance, 279	zlm, <i>143</i>
spatNetwDistributionsKneighbors, 280	,
spatPlot, 281, 286, 288	
spatPlot2D, 281, 282, 283, 288, 299	
spatPlot3D, 282, 286, 286, 300	
specificCellCellcommunicationScores,	
289	
split_dendrogram_in_two, 290	
standardise_giotto, 291	
stitchFieldCoordinates, 60, 291	
stitchTileCoordinates, 292	
subClusterCells, 293	
subsetGiotto, 295	
subsetGiottoLocs, 295	
trendSceek, 296	
trendsceek_test, 296	
triangulate, 66, 67, 200	
umap, 221	
updateGiottoImage, 297	
viewHMRFresults, 298	
viewHMRFresults2D, 299	
viewHMRFresults3D, 300	
violinPlot, 301	
visDimGenePlot, 302	
,	
visDimGenePlot_2D_ggplot, 304	
visDimGenePlot_3D_plotly, 305	
visDimPlot, 307	
visDimPlot_2D_ggplot, 309	
visDimPlot_2D_plotly, 311	
visDimPlot_3D_plotly, 313	
visForceLayoutPlot, 314	
visGenePlot, 316	