Package: ggspavis (via r-universe)

September 22, 2024

Version 1.11.0

Title Visualization functions for spatial transcriptomics data

Description Visualization functions for spatial transcriptomics data.

Includes functions to generate several types of plots, including spot plots, feature (molecule) plots, reduced dimension plots, spot-level quality control (QC) plots, and feature-level QC plots, for datasets from the 10x Genomics Visium and other technological platforms. Datasets are assumed to be in either SpatialExperiment or SingleCellExperiment format.

URL https://github.com/lmweber/ggspavis

BugReports https://github.com/lmweber/ggspavis/issues

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Encoding UTF-8

biocViews Spatial, SingleCell, Transcriptomics, GeneExpression, QualityControl, DimensionReduction

Depends ggplot2

Imports SpatialExperiment, SingleCellExperiment, SummarizedExperiment, ggside, grid, ggrepel, RColorBrewer, scales, grDevices, methods, stats

VignetteBuilder knitr

Suggests BiocStyle, rmarkdown, knitr, STexampleData, BumpyMatrix, scater, scran, uwot, testthat, patchwork

RoxygenNote 7.3.1

Repository https://bioc.r-universe.dev

RemoteUrl https://github.com/bioc/ggspavis

RemoteRef HEAD

RemoteSha 57a47e6a3f2c2add0974d1f5d989e33c6cf1cd31

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plotDimRed

plotDimRed

Description

Plotting functions for spatial transcriptomics data.

Usage

```
plotDimRed(
   spe,
   plot_type = c("UMAP", "PCA"),
   annotate = NULL,
   feature_names = NULL,
   assay_name = "counts",
   update_dimnames = TRUE,
   pal = NULL,
   point_size = 0.3,
   legend_point_size = 3,
   text_by = NULL,
   text_by_size = 5,
   text_by_color = "black"
)
```

Arguments

spe	Input data, assumed to be a SpatialExperiment or SingleCellExperiment object.
plot_type	Type of reduced dimension plot. Possible options are "UMAP", "PCA", or any other set of reduced dimensions stored in the input object. Default = "UMAP".
annotate	Variable to show as annotations. This may be discrete or continuous. For a discrete variable (e.g. cluster labels), this should be the name of a column in colData containing a character vector or factor. For a continuous variable (e.g. a gene name), this should be an entry in feature_names. Default = NULL.

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feature_names	Name of column in rowData containing names of continuous features to plot
	(e.g. gene names). For example, set to feature_names = "gene_name" if gene

names are stored in a column named "gene_name". This argument is used if annotate is a continuous variable. Default = NULL, in which case the row

names of the input object will be used.

assay_name Name of assay in input object containing values to plot for a continuous vari-

able. Default = "counts".

update_dimnames

Whether to update column names of reducedDims to default values for plotting.

Default = TRUE.

pal Color palette for annotations. Options for discrete values are "libd_layer_colors",

"Okabe-Ito", or any vector of color names or hex values. For continuous values, provide a vector of length 2 for the low and high range, e.g. c("gray90", "navy").

point_size Point size. Default = 0.3.

legend_point_size

Legend point size for discrete annotations. Default = 3.

text_by Column name of annotation labels to display over each cluster of points. This

will usually be the same as annotate. Alternatively, another column may be used (e.g. with more readable classes or shorter strings). Only used for discrete

annotate. Default = NULL.

text_by_size Text size for annotation labels over each cluster. Default = 5.

Details

Function to create reduced dimension plot (e.g. PCA or UMAP) with additional optional annotations such as cluster labels, expression of a gene, or quality control metrics.

Value

Returns a ggplot object, which may be further modified using ggplot functions.

Author(s)

Lukas M. Weber and Yixing E. Dong

```
library(STexampleData)
spe <- Visium_humanDLPFC()

# select spots over tissue
spe <- spe[, colData(spe)$in_tissue == 1]

# use small subset of data for this example
n <- 200
set.seed(123)
spe <- spe[, sample(seq_len(ncol(spe)), n)]</pre>
```

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```
# calculate logcounts
library(scran)
spe <- logNormCounts(spe)</pre>
# identify top highly variable genes (HVGs)
is_mito <- grepl("(^MT-)|(^mt-)", rowData(spe)$gene_name)</pre>
spe <- spe[!is_mito, ]</pre>
dec <- modelGeneVar(spe)</pre>
top_hvgs <- getTopHVGs(dec, prop = 0.1)</pre>
# run dimensionality reduction
library(scater)
set.seed(123)
spe <- runPCA(spe, subset_row = top_hvgs)</pre>
set.seed(123)
spe <- runUMAP(spe, dimred = "PCA")</pre>
colnames(reducedDim(spe, "UMAP")) <- paste0("UMAP", 1:2)</pre>
# generate plot
plotDimRed(spe, plot_type = "UMAP", annotate = "ground_truth")
```

plotFeatureQC

plotFeatureQC

Description

Plotting functions for spatial transcriptomics data.

Usage

```
plotFeatureQC(
   spe,
   plot_type = c("histogram", "violin"),
   x_metric = NULL,
   annotate = NULL,
   n_bins = 100,
   point_size = 0.1,
   scale_log1p = TRUE
)
```

Arguments

spe Input data, assumed to be a SpatialExperiment or SingleCellExperiment

object.

plot_type Type of QC plot. Options are "histogram" and "violin". See Details for addi-

tional details.

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x_metric	Name of column in rowData containing feature-level QC metric to plot on x-axis. Required for histograms and violin plots.
annotate	Name of column in rowData identifying selected features that do not meet QC filtering thresholds, which will be highlighted on a histogram or violin plot. Default = NULL. Optional argument used for histograms and violin plots.
n_bins	Number of bins for histograms. Default = 100. Optional argument used for histograms.
point_size	Point size. Default = 0.1 . Optional argument for violin plots.
scale_log1p	Whether to $log1p$ -scale axes. Default = TRUE.

Details

Function to create quality control (QC) plots for spatial transcriptomics data.

The following types of QC plots are available for feature-level QC (see plotSpotQC for spot-level or cell-level QC):

- Histogram (plot_type = "histogram") for a single QC metric, e.g. total UMI counts across all spots per feature. The histogram can optionally highlight selected features, e.g. low abundance features.
- Violin (plot_type = "violin") for a single QC metric, e.g. total UMI counts across all spots per feature. The violin plot can optionally highlight selected features, e.g. low abundance features.

Value

Returns a ggplot object, which may be further modified using ggplot functions.

Author(s)

Yixing E. Dong and Lukas M. Weber

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plotMol	ecules
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plotMolecules

Description

Plotting functions for spatial transcriptomics data.

Usage

```
plotMolecules(
   spe,
   molecule = NULL,
   x_coord = NULL,
   y_coord = NULL,
   sample_id = "sample_id",
   pal = c("gray90", "navy"),
   point_size = 0.3
)
```

Arguments

spe	(SpatialExperiment) Input data, assumed to be a SpatialExperiment object.
molecule	Name of mRNA molecule to plot (assumed to match one of the row names of $rowData$).
x_coord	Name of column in spatialCoords containing x coordinates. Default = NULL, which selects the first column of spatialCoords.
y_coord	Name of column in spatialCoords containing y coordinates. Default = $NULL$, which selects the second column of spatialCoords.
sample_id	Name of column in colData containing sample IDs. This argument is only required for datasets containing multiple samples (tissue sections). If provided, samples will be shown in multiple panels using facetting. Default = NULL.
pal	Color palette, provided as a vector of length 2 for the low and high range. Default = $c("gray90", "navy")$.
point_size	Point size. Default = 0.3 .

Details

Function to create spot plot for molecule-based datasets, showing spatial locations in x-y coordinates with optional annotations such as expression of a gene.

Value

Returns a ggplot object, which may be further modified using ggplot functions.

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Author(s)

Lukas M. Weber

Examples

```
library(STexampleData)
spe <- seqFISH_mouseEmbryo()
plotMolecules(spe, molecule = "Sox2")</pre>
```

plotSpotQC

plotSpotQC

Description

Plotting functions for spatial transcriptomics data.

Usage

```
plotSpotQC(
  spe,
  plot_type = c("histogram", "scatter", "spot", "violin"),
  x\_coord = NULL,
  y_coord = NULL,
  x_{metric} = NULL,
 y_metric = NULL,
  x_threshold = NULL,
 y_threshold = NULL,
  trend = TRUE,
 marginal = TRUE,
  annotate = NULL,
  in_tissue = NULL,
  legend_point_size = 3,
  n_bins = 100,
  point_size = 0.3,
 y_reverse = TRUE
)
plotQC(...)
```

Arguments

spe Input data, assumed to be a SpatialExperiment or SingleCellExperiment

object.

Type of QC plot. Options are "histogram", "scatter", "spot", and "violin". See

Details for additional details.

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x_coord	Name of column in spatialCoords (for a SpatialExperiment input object) or colData (for a SingleCellExperiment input object) containing x coordinates. Default = NULL (for a SpatialExperiment, the first column of spatialCoords will be selected in this case). Used for spot plots.
y_coord	Name of column in spatialCoords (for a SpatialExperiment input object) or colData (for a SingleCellExperiment input object) containing y coordinates. Default = NULL (for a SpatialExperiment, the second column of spatialCoords will be selected in this case). Used for spot plots.
x_metric	Name of column in colData containing QC metric to plot on x-axis. Required for histograms, scatter plots, and violin plots.
y_metric	Name of column in colData containing QC metric to plot on y-axis. Required for histograms, scatter plots, and violin plots.
x_threshold	QC filtering threshold on x-axis metric to highlight with vertical line. Default = NULL. Optional argument used for scatter plots.
y_threshold	QC filtering threshold on y-axis metric to highlight with horizontal line. Default = NULL. Optional argument used for scatter plots.
trend	Whether to show smoothed trend line (loess). Default = TRUE. Optional argument used for scatter plots.
marginal	Whether to show marginal histograms. Default = TRUE. Optional argument used for scatter plots.
annotate	Name of column in colData identifying selected spots that do not meet QC filtering thresholds, which will be highlighted on a histogram, spot plot, or violin plot. Default = NULL. Optional argument used for histograms, spot plots, and violin plots.
in_tissue	Name of column in colData identifying spots over tissue (e.g. "in_tissue" for 10x Genomics Visium datasets). If this argument is provided, only spots over tissue will be shown. Default = NULL. Optional argument used for spot plots.
legend_point_s	
	Legend point size. Default = 3. Optional argument used for spot plots.
n_bins	Number of bins for histograms. Default = 100. Optional argument used for histograms.
point_size	Point size. Default = 0.3. Optional argument for scatter plots, spot plots, and violin plots. Suggested values: 0.5 for scatter plots, 0.3 for spot plots, 0.1 for violin plots.
y_reverse	Whether to reverse y coordinates. This is usually required for 10x Genomics Visium datasets when using the default coordinate values. Default = TRUE. Set to FALSE if not needed, e.g. for other platforms. Optional argument used for spot plots.
• • •	Not used.

Details

Function to create quality control (QC) plots for spatial transcriptomics data.

The following types of QC plots are available for spot-level or cell-level QC (see plotFeatureQC for feature-level QC):

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• Histogram (plot_type = "histogram") for a single QC metric, e.g. number of UMI counts per spot. For number of counts per spot, the histogram can optionally highlight selected spots, e.g. spots with low library size.

- Scatter plot (plot_type = "scatter") comparing two QC metrics, e.g. number of detected features vs. number of cells per spot, with optional horizontal and vertical lines highlighting QC filtering thresholds.
- Spot plot (plot_type = "spot") showing spots in spatial x-y coordinates, e.g. highlighting selected spots that do not meet filtering thresholds.
- Violin plot (plot_type = "violin") for a single QC metric, e.g. number of UMI counts per spot. For number of counts per spot, the violin plot can optionally highlight selected spots, e.g. spots with low library size.

Value

Returns a ggplot object, which may be further modified using ggplot functions.

Author(s)

Lukas M. Weber and Yixing E. Dong

Examples

```
library(STexampleData)
spe <- Visium_humanDLPFC()

colData(spe)$sum <- colSums(counts(spe))
colData(spe)$low_libsize <- colData(spe)$sum < 400

plotSpotQC(spe, plot_type = "histogram", x_metric = "sum", annotate = "low_libsize")
plotSpotQC(spe, plot_type = "scatter", x_metric = "sum", y_metric = "cell_count")
plotSpotQC(spe, plot_type = "spot", annotate = "low_libsize", in_tissue = "in_tissue")
plotSpotQC(spe, plot_type = "violin", x_metric = "sum", annotate = "low_libsize")</pre>
```

plotSpots

plotSpots

Description

Plotting functions for spatial transcriptomics data.

Usage

```
plotSpots(
   spe,
   x_coord = NULL,
   y_coord = NULL,
   sample_id = NULL,
```

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```
in_tissue = "in_tissue",
  annotate = NULL,
  feature_names = NULL,
  assay_name = "counts",
  pal = NULL,
  point_size = 0.3,
  legend_position = "right",
  legend_point_size = 3,
  show_axes = FALSE,
 y_reverse = TRUE,
  text_by = NULL,
  text_by_size = 5,
  text_by_color = "black"
)
```

Arguments

spe Input data, assumed to be a SpatialExperiment or SingleCellExperiment

object.

 x_{coord} Name of column in spatialCoords (for a SpatialExperiment input object) or

colData (for a SingleCellExperiment input object) containing x coordinates. Default = NULL (for a SpatialExperiment, the first column of spatialCoords

will be selected in this case).

y_coord Name of column in spatialCoords (for a SpatialExperiment input object)

> or colData (for a SingleCellExperiment input object) containing y coordinates. Default = NULL (for a SpatialExperiment, the second column of

spatialCoords will be selected in this case).

sample_id Name of column in colData containing sample IDs. This argument is only

required for datasets containing multiple samples (tissue sections). If provided,

samples will be shown in multiple panels using facetting. Default = NULL.

Name of column in colData identifying spots over tissue (e.g. "in_tissue" for in_tissue

10x Genomics Visium datasets). If this argument is provided, only spots over tissue will be shown. Default = "in_tissue". Set to NULL to display all spots.

annotate Variable to show as annotations. This may be discrete or continuous. For a

> discrete variable (e.g. cluster labels), this should be the name of a column in colData containing a character vector or factor. For a continuous variable (e.g.

a gene name), this should be an entry in feature_names. Default = NULL.

Name of column in rowData containing names of continuous features to plot feature_names

> (e.g. gene names). For example, set to feature_names = "gene_name" if gene names are stored in a column named "gene_name". This argument is used if annotate is a continuous variable. Default = NULL, in which case the row

names of the input object will be used.

Name of assay in input object containing values to plot for a continuous variassay_name

able. Default = "counts".

Color palette for annotations. Options for discrete values are "libd_layer_colors", pal

> "Okabe-Ito", or any vector of color names or hex values. For continuous values, provide a vector of length 2 for the low and high range, e.g. c("gray90", "navy").

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Point size. Default = 0.3. point_size legend_position Legend position for discrete annotations. Options are "left", "right", "top", "bottom", and "none". Default = "right". legend_point_size Legend point size for discrete annotations. Default = 3. show_axes Whether to show axis titles, text, and ticks. Default = FALSE. Whether to reverse y coordinates. This is usually required for 10x Genomics y_reverse Visium datasets when using the default coordinate values. Default = TRUE. Set to FALSE if not needed, e.g. for other platforms. text_by Column name of annotation labels to display over each cluster of points. This will usually be the same as annotate. Alternatively, another column may be used (e.g. with more readable classes or shorter strings). Only used for discrete annotate. Default = NULL. text_by_size Text size for annotation labels over each cluster. Default = 5. Color name or hex code for annotation labels. Default = "black". text_by_color

Details

Function to create spot plot showing spatial locations in x-y coordinates with optional annotations such as cluster labels, expression of a gene, or quality control metrics.

Value

Returns a ggplot object, which may be further modified using ggplot functions.

Author(s)

Lukas M. Weber and Yixing E. Dong

```
library(STexampleData)

# discrete annotations
spe <- Visium_humanDLPFC()
plotSpots(spe, annotate = "ground_truth")

# continuous annotations
spe <- Visium_mouseCoronal()
plotSpots(spe, annotate = "Gapdh", feature_names = "gene_name")</pre>
```

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plotVisium plotVisium

Description

Plots for spatially resolved transcriptomics data from the 10x Genomics Visium platform

Usage

```
plotVisium(
  spe,
  spots = TRUE,
  annotate = NULL,
  highlight = NULL,
  facets = "sample_id",
  image = TRUE,
  zoom = FALSE,
  show_axes = FALSE,
  assay = "counts",
  trans = "identity",
  point_size = 1,
  legend_position = "right",
  x\_coord = NULL,
 y_{coord} = NULL,
 y_reverse = TRUE,
  sample_ids = NULL,
  image_ids = NULL,
 pal = NULL
)
```

Arguments

spe	(SpatialExperiment) Input data object.
spots	(logical) Whether to display spots (spatial barcodes) as points. Default = TRUE.
annotate	(character) Column in colData to use to fill points by color. If annotate contains a numeric column (e.g. total UMI counts), a continuous color scale will be used. If annotate contains a factor (e.g. cluster labels), a discrete color scale will be used. Default = $NULL$.
highlight	(character) Column in colData to use to highlight points by outlining them. For example, in_tissue will highlight spots overlapping with tissue. Default = NULL.
facets	(character) Column in colData to use to facet plots, i.e. show multiple panels of plots. Default = "sample_id". Set to NULL to disable.
image	(logical) Whether to show histology image as background. Default = TRUE.
zoom	(logical) Whether to zoom to area of tissue containing spots. Default = FALSE

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(logical) Whether to show axes and coordinates. Default = FALSE
(character) Name of assay data to use when annotate is in rownames(spe). Should be one of assayNames(spe).
Transformation to apply for continuous scales. Ignored unless annotate is numeric, e.g. feature expression. (See ggplot2{continuous_scale} for valid options.)
(numeric) Point size. Default = 1.
n
Legend position for annotations. Options are "left", "right", "top", "bottom", and "none". Default = "right".
(character) Column in spatialCoords containing x-coordinates. Default = $NULL$, which selects the first column.
(character) Column in spatialCoords containing y-coordinates. Default = NULL, which selects the second column.
(logical) Whether to reverse y coordinates, which is often required for Visium data, depending on the orientation of the raw data. Default = TRUE.
(character) Samples to show, if multiple samples are available. Default = NULL (show all samples).
(character) Images to show, if multiple images are available. Default = NULL (show all images).
(character) Color palette for points. Options for discrete labels are "libd_layer_colors", "Okabe-Ito", or a custom vector of hex color codes. Options for continuous values are "viridis", a single color name (e.g. "red", "navy", etc), or a vector of length two containing color names for each end of the scale. Default = "libd_layer_colors" for discrete data, and "viridis" for continuous data.

Details

Function to generate plots for spatially resolved transcriptomics datasets from the 10x Genomics Visium spatially platform.

This function generates a plot for spot-based spatially resolved transcriptomics data from the 10x Genomics Visium platform, with several options available to adjust the plot type and style.

Value

Returns a ggplot object. Additional plot elements can be added as ggplot elements (e.g. title, customized formatting, etc).

Author(s)

Helena L. Crowell, with modifications by Lukas M. Weber and Yixing E. Dong

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```
library(STexampleData)

spe <- Visium_mouseCoronal()

# color by x coordinate, highlight in-tissue spots
plotVisium(spe, annotate = "pxl_col_in_fullres", highlight = "in_tissue")

# subset in-tissue spots
sub <- spe[, as.logical(colData(spe)$in_tissue)]

# color by feature counts, don't include image
rownames(sub) <- make.names(rowData(sub)$gene_name)
plotVisium(sub, annotate = "Gad2", assay = "counts")</pre>
```

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