

Package ‘SpaceTrooper’

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Type Package

Title SpaceTrooper performs Quality Control analysis of Image-Based spatial

Version 0.99.3

Description SpaceTrooper performs Quality Control analysis using data driven GLM models of Image-Based spatial data, providing exploration plots, QC metrics computation, outlier detection.

It implements a GLM strategy for the detection of low quality cells in imaging-based spatial data (Transcriptomics and Proteomics).

It additionally implements several plots for the visualization of imaging based polygons through the ggplot2 package.

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.addPolygonsToCD	<i>.addPolygonsToCD</i>
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Description

This function enriches a DataFrame (e.g., from colData) with matching polygon geometries.

Usage

```
.addPolygonsToCD(cd, polygons, polygonsCol = "polygons")
```

Arguments

cd	A DataFrame containing at least ‘fov’ and ‘cellID’ columns.
polygons	An sf object with matching ‘fov’ and ‘cellID’ columns.
polygonsCol	character indicating the name of the polygons column to add into the colData (default is ‘polygons’).

Value

A DataFrame identical to ‘cd’, but row-subset to cells present in ‘polygons’ and with a new ‘polygons’ list-column of sf geometries.

.centroid_image_theme	<i>.centroid_image_theme</i>
-----------------------	------------------------------

Description

internal function to setup the theme for the centroid plot background

Usage

```
.centroid_image_theme(backBorder = NA)
```

Arguments

backBorder	color for the borders of the background (default=NA)
------------	--

Value

a ggplot2 theme object

```
.checkPolygonsValidity
  .checkPolygonsValidity
```

Description

checks validity on a geometry of ‘sf’ object. It removes multipolygons when ‘keepMultiPol’ is ‘FALSE’

Usage

```
.checkPolygonsValidity(
  sf,
  geometry = NULL,
  keepMultiPol = TRUE,
  verbose = FALSE
)
```

Arguments

<code>sf</code>	An ‘sf’ class object containing the spatial data.
<code>geometry</code>	character for the geometry to check validity, if ‘NULL’ it checks the active geometry (default is ‘NULL’)
<code>keepMultiPol</code>	logical for keeping/removing multipolygons, if any (default is ‘TRUE’, so keeping the multipolygons)
<code>verbose</code>	logical to print verbose output (default is ‘FALSE’)

Details

In case geometry is NULL validity is checked on the active geometry, otherwise it is checked on the passed geometry without changing the active geometry of the sf object. In case of not valid polygons, these are removed. If keeMultiPol is FALSE, possible detected multipolygons are removed.

Value

An ‘sf’ object with valid geometries, possibly with multipolygons removed.

```
.computeBorderDistanceCosMx  
. computeBorderDistanceCosMx
```

Description

Calculates the minimum distance of each cell to the field-of-view border and adds it to ‘colData’.

Usage

```
.computeBorderDistanceCosMx(  
  spe,  
  xwindim = metadata(spe)$fov_dim[["xdim"]],  
  ywindim = metadata(spe)$fov_dim[["ydim"]]  
)
```

Arguments

spe	A ‘SpatialExperiment‘ object with CosMx data.
xwindim	Width of FOV in x (default from ‘metadata(spe)\$fov_dim’).
ywindim	Height of FOV in y (default from ‘metadata(spe)\$fov_dim’).

Value

A ‘SpatialExperiment‘ object with ‘dist_border‘ columns in ‘colData’.

```
.createPolygons      .createPolygons
```

Description

This internal function creates polygons from a data.frame or similar object.

Usage

```
.createPolygons(  
  spat_obj,  
  x = NULL,  
  y = NULL,  
  polygon_id = NULL,  
  geometry = "Geometry"  
)
```

Arguments

spat_obj	A data frame or similar object containing spatial data.
x	A character vector specifying the x-coordinates.
y	A character vector specifying the y-coordinates.
polygon_id	A character string specifying the polygon ID.

Value

An ‘sf’ object containing the created polygons.

.dark_theme*.dark_theme***Description**

internal function to setup the black background theme for the First Filter plot

Usage

```
.dark_theme(fillColor = "black", foreColor = "white")
```

Arguments

fillColor	color to fill the element_rect (default is "black")
foreColor	color for all the other elements (default is "white")

Value

a ggplot2 theme object

.fov_image_theme*.fov_image_theme***Description**

internal function to setup the theme for the fov background on the whole image

Usage

```
.fov_image_theme(backColor = "black", backBorder = NA, titleCol = "white")
```

Arguments

backColor	not used
backBorder	color for the borders of the background (default=NA)
titleCol	character indicating the color of the title

Value

a ggplot2 theme object

```
.getActiveGeometryName  
  .getActiveGeometryName
```

Description

.getActiveGeometryName

Usage

```
.getActiveGeometryName(sf)
```

Arguments

sf an sf object

Value

character with the name of the active geometry

Examples

```
example(readPolygonsCosmx)  
.getActiveGeometryName(polygons)
```

```
.light_theme                 .light_theme
```

Description

internal function to setup the white background theme for the First Filter plot

Usage

```
.light_theme(fillColor = "white", foreColor = "black")
```

Value

a ggplot2 theme object

`.negative_image_theme` *.negative_image_theme*

Description

internal function to setup the theme for the negative background for negative plots

Usage

```
.negative_image_theme(fillColor = "black", foreColor = "white")
```

Arguments

<code>fillColor</code>	color to fill the element_rect (default is "black")
<code>foreColor</code>	color for all the other elements (default is "white")

Value

a ggplot2 theme object

`.renameGeometry` *.renameGeometry*

Description

renames the ‘from’ to ‘to’ geometry of the ‘sf’ object. If ‘activate’ is ‘TRUE’ it set as the active geometry the new geometry name. Default behaviour is to check if the renamed geometry is already active and leave it as active with the new name.

Usage

```
.renameGeometry(sf, from, to, activate = FALSE)
```

Arguments

<code>sf</code>	an sf object with the ‘from’ geometry
<code>from</code>	character indicating the name of the geometry to change
<code>to</code>	character indicating the new name of the geometry
<code>activate</code>	logical indicating if the renamed geometry has to be activated

Value

an sf object

Examples

```
example(readPolygonsCosmx)
.renameGeometry(polygons, "global", "global1")
```

. setActiveGeometry .*setActiveGeometry*

Description

.setActiveGeometry

Usage

.setActiveGeometry(sf, name)

Arguments

sf	an sf object
name	character for the geometry to activate

Value

an sf object

Examples

```
example(readPolygonsCosmx)
.setActiveGeometry(polygons, "local")
```

addPolygonsToSPE *addPolygonsToSPE*

Description

This function adds polygon data to a ‘SpatialExperiment‘ object.

Usage

addPolygonsToSPE(spe, polygons, polygonsCol = "polygons")

Arguments

spe	A ‘SpatialExperiment‘ object to which polygons will be added.
polygons	An ‘sf‘ object containing the polygon data.
polygonsCol	character indicating the name of the polygons column to add into the colData (default is ‘polygons‘).

Value

The ‘SpatialExperiment‘ object with polygons added to the ‘colData‘.

Examples

```
example(readCosmxSPE)
polygons <- readPolygonsCosmx(metadata(spe)$polygons)
spe <- addPolygonsToSPE(spe, polygons)
spe$polygons
```

computeAreaFromPolygons
computeAreaFromPolygons

Description

This function computes the area from polygon data.

Usage

```
computeAreaFromPolygons(polygons)
```

Arguments

polygons	An ‘sf‘ object containing polygon data.
----------	---

Value

A ‘numeric‘ vector with the area information.

Examples

```
example(readPolygonsMerfish)
area <- computeAreaFromPolygons(polygons)
area
```

computeAspectRatioFromPolygons
computeAspectRatioFromPolygons

Description

This function computes the aspect ratio from polygon data.

Usage

```
computeAspectRatioFromPolygons(polygons)
```

Arguments

polygons	An ‘sf‘ object containing polygon data.
----------	---

Value

A ‘numeric‘ vector with the aspect ratio information.

Examples

```
example(readPolygonsMerfish)
ar <- computeAspectRatioFromPolygons(polygons)
ar
```

computeCenterFromPolygons	<i>computeCenterFromPolygons</i>
---------------------------	----------------------------------

Description

This function computes the center coordinates on x and y axis from polygon data and adds it to the ‘colData’. It is necessary only for Merfish.

Usage

```
computeCenterFromPolygons(polygons, coldata)
```

Arguments

- | | |
|----------|---|
| polygons | An ‘sf’ object containing polygon data. |
| coldata | A ‘DataFrame‘ containing the ‘colData‘ to which center coordinates information will be added. |

Value

A ‘DataFrame‘ with the added center information.

Examples

```
example(readPolygonsMerfish)
coldata <- computeCenterFromPolygons(polygons, colData(spe))
colData(spe) <- coldata
```

computeLambda	<i>computeLambda</i>
---------------	----------------------

Description

Compute Optimal Ridge Regularization Parameter λ via Cross-Validation

computeLambda performs ridge (L2) logistic regression with cross-validation to identify the optimal regularization parameter λ for a binary response.

Usage

```
computeLambda(technology, trainDF)
```

Arguments

<code>technology</code>	\[character\] The name of the experimental technology. Passed to <code>getModelFormula()</code> to retrieve the corresponding model formula.
<code>trainDF</code>	\[data.frame\] A data frame for training that must include:
	Predictor columns All columns referenced in the formula returned by <code>getModelFormula()</code> . <code>qscore_train</code> A binary (0/1) response vector to be modeled.

Details

Internally, the function:

1. Calls `getModelFormula(technology)` to obtain a model formula as text,
2. Constructs the design matrix via `model.matrix()`,
3. Runs ridge logistic regression cross-validation using `cv.glmnet` with `alpha = 0`,
4. Extracts and returns `ridge_cv$lambda.min`.

Value

\[numeric\] The value of λ (i.e., `lambda.min`) from `cv.glmnet` that minimizes the cross-validation error.

See Also

[cv.glmnet](#)

Examples

```
example(spatialPerCellQC)
withr::with_seed(1998, trainDF <- computeTrainDF(spe))
best_lambda <- computeLambda(metadata(spe)$technology, trainDF)
print(best_lambda)
```

computeMissingMetricsMerfish
computeMissingMetricsMerfish

Description

‘computeMissingMetricsMerfish()’ takes cell metadata and boundary polygons, calculates per-cell area and aspect-ratio, and optionally appends the raw polygon geometries.

Usage

```
computeMissingMetricsMerfish(
  polFile,
  coldata,
  boundariesType = c("parquet", "HDF5"),
  keepPolygons = FALSE,
  polygonsCol = "polygons"
)
```

Arguments

polFile	path to the polygon file
colData	'DataFrame' or 'data.frame' Cell metadata with at least a 'cell_id' column.
boundariesType	'character(1)' One of "HDF5" or "parquet"—passed on to 'readPolygons-Merfish()'.
keepPolygons	'logical(1)' If 'TRUE', cbinds the raw polygon 'sf' columns onto 'colData'.
polygonsCol	character indicating the name of the polygons column to add into the colData (default is 'polygons').

Value

A 'DataFrame' (or 'data.frame') with: - all columns of 'colData' - 'um_area': area of each cell's polygon - 'AspectRatio': width/height aspect ratio - (optionally) the polygon geometries

Examples

```
example(readMerfishSPE)
cd <- computeMissingMetricsMerfish(metadata(spe)$polygons, colData(spe),
  boundariesType="parquet")
colData(spe) <- cd
cd
```

computeMissingMetricsXenium
computeMissingMetricsXenium

Description

Compute Missing Metrics for Xenium Data

This function computes missing metrics, such as the aspect ratio, from polygon data in a Xenium dataset and optionally appends the polygon data to the resulting 'colData'.

Usage

```
computeMissingMetricsXenium(
  polFile,
  colData,
  keepPolygons = FALSE,
  polygonsCol = "polygons"
)
```

Arguments

polFile	A character string specifying the file path to the polygon data.
colData	A 'DataFrame' containing the 'colData' for the Xenium dataset.
keepPolygons	A logical value indicating whether to keep the polygon data in the resulting 'colData'. Default is 'FALSE'.
polygonsCol	character indicating the name of the polygons column to add into the colData (default is 'polygons').

Details

The function reads the polygon data from the specified file, computes the aspect ratio for each polygon, and merges these metrics with the provided ‘colData’. Optionally, the polygon data can be kept in the returned ‘colData’.

Value

A ‘DataFrame‘ containing the updated ‘colData‘ with computed metrics. If ‘keepPolygons‘ is ‘TRUE‘, the polygon data is also included.

Examples

```
example(readXeniumSPE)
colData(spe) <- computeMissingMetricsXenium(metadata(spe)$polygons,
                                             colData(spe), keepPolygons=TRUE)
```

`computeQScore`

computeQScore

Description

Compute quality score and automatically define weights for quality score through glm training. This function computes quality score with a formula that depends on the technology.

Usage

```
computeQScore(spe, bestLambda = NULL, verbose = FALSE)
```

Arguments

- | | |
|------------|---|
| spe | A ‘SpatialExperiment‘ object with spatial transcriptomics data. |
| bestLambda | the best lambda typically computed using ‘computeLambda‘. |
| verbose | logical for having a verbose output. Default is FALSE. |

Details

For CosMx datasets, the Quality Score formula is defined as follows:

quality score ~ count density - aspect ratio - interaction term

count density is total counts-to-area ratio, aspect ratio represents border effect typical of CosMx datasets and the last one is the interaction term of the previous two terms.

For Xenium and Merscope datasets, quality score depends solely on count density, as no border effect has been observed for these two technologies.

To automatically define the formula coefficient weights, model training is performed through ridge regression.

Value

The ‘SpatialExperiment‘ object with added quality score in ‘colData‘.

Examples

```
example(spatialPerCellQC)
set.seed(1998)
spe <- computeQScore(spe)
summary(spe$training_status)
summary(spe$quality_score)
```

`computeQScoreFlags` *computeQScoreFlags*

Description

Compute flagged cells based on a manually chosen threshold on quality score

This function Compute flagged cells based on a manually chosen threshold on quality score stored in ‘SpatialExperiment’ object.

Usage

```
computeQScoreFlags(spe, qsThreshold = 0.5, useQSQuantiles = FALSE)
```

Arguments

<code>spe</code>	A ‘SpatialExperiment‘ object with spatial transcriptomics data.
<code>qsThreshold</code>	Numeric threshold or quantile for quality score. Default ‘0.5‘.
<code>useQSQuantiles</code>	Logical; if ‘TRUE‘, treat ‘qsThreshold‘ as a percentile.

Value

The ‘SpatialExperiment‘ object with added filter flags in ‘colData‘.

Examples

```
example(computeQScore)
spe <- computeQScoreFlags(spe)
table(spe$low_qsore)
# if fixed filters are defined we have an additional column
spe <- computeThresholdFlags(spe)
spe <- computeQScoreFlags(spe)
table(spe$low_threshold_qsore)
```

`computeSpatialOutlier` *computeSpatialOutlier*

Description

Computes outliers based on the Area (in micron) of the experiment. It gives the possibility to choose between the medcouple (mc method argument) and the MADs (scuttle method argument).

Usage

```
computeSpatialOutlier(
  spe,
  computeBy = NULL,
  method = c("mc", "scuttle", "both"),
  mcDoScale = FALSE,
  scuttleType = c("both", "lower", "higher")
)
```

Arguments

<code>spe</code>	a SpatialExperiment object with target_counts, area in micron and log2 of the aspect ratio in the ‘colData’.
<code>computeBy</code>	character indicating a ‘colData’ column name on which compute the outlier.
<code>method</code>	one of ‘mc’, ‘scuttle’, ‘both’. Use ‘mc’ for medcouple, ‘scuttle’ for median absolute deviations as computed in ‘scuttle’, ‘both’ for computing both of them.
<code>mcDoScale</code>	logical indicating if the values to compute the medcouple for the outlier detection should be scaled (default is FALSE, as suggested by the original Medcouple authors.). See mc for further readings.
<code>scuttleType</code>	One of ““both”“, ““lower”“, ““higher”“ for scuttle method.

Details

The medcouple method is a measure for the skewness of univariate distribution as described in Hubert M. et al. (2008). In particular, the computed medcouple value must be in a range between -0.6 and 0.6 to compute adjusted boxplots and perform the outlier detection. For median absolute deviations (MADs) method we just wrap the `isOutlier` function in the `scuttle` package. Please see McCarthy DJ et al (2017) for further details.

Value

a SpatialExperiment object with additional column(s) (named as the column name indicated in ‘column_by’ followed by the outlier_sc/mc nomenclature) with the outlier detection as ‘outlier.filter’ logical class object. This allows to store the thresholds as attributes of the column. use `attr("thresholds")` to retrieve them.

Examples

```
example(spatialPerCellQC)
spe <- computeSpatialOutlier(spe, computeBy="log2CountArea", method="both")
table(spe$log2CountArea_outlier_mc)
table(spe$log2CountArea_outlier_sc)
```

computeThresholdFlags *computeThresholdFlags*

Description

Compute Flagged cells using fixed thresholds for SpatialExperiment.

This function calculates flagged cells only for total counts and control on total probe counts ratio using fixed thresholds for a ‘SpatialExperiment‘ object.

Usage

```
computeThresholdFlags(spe, totalThreshold = 0, ctrlTotRatioThreshold = 0.1)
```

Arguments

- `spe` A ‘SpatialExperiment‘ object with spatial transcriptomics data.
- `totalThreshold` A numeric value for the threshold of total counts to identify cells with low counts. Default is ‘0‘.
- `ctrlTotRatioThreshold` A numeric value for the threshold of control-to-total ratio to flag cells over a certain threshold. Default is ‘0.1‘.

Details

The function flags cells basing on zero counts and control-to-total ratio to identify junk cells. It also combines these flags into a single filter flag.

Value

The ‘SpatialExperiment‘ object with added filter flags in ‘colData‘.

Examples

```
example(readCosmxSPE)
spe <- spatialPerCellQC(spe)
spe <- computeThresholdFlags(spe)
table(spe$threshold_flags)
```

computeTrainDF *computeTrainDF*

Description

Build a Balanced Training Data Frame from a SpatialExperiment

`computeTrainDF` takes a **SpatialExperiment** object, flags spatial outliers on “log2CountArea”, then assembles a balanced training set of “good” vs “bad” cells for subsequent model fitting.

Usage

```
computeTrainDF(spe, verbose = FALSE)
```

Arguments

<code>spe</code>	<code>SpatialExperiment</code> A SpatialExperiment containing at least:
	<ul style="list-style-type: none"> • assay(s) with nonzero total counts, • <code>colData(spe)</code> columns including <code>log2CountArea</code>, <code>dist_border</code>, etc., • <code>metadata(spe)\$technology</code> indicating the platform.
<code>verbose</code>	<code>\[logical(1)\]</code> (default FALSE) If TRUE, prints the number of “bad” and “good” cells selected.

Details

Internally the function:

1. Filters out zero-count cells,
2. Calls `computeSpatialOutlier()` on “`log2CountArea`” to get fences,
3. Labels cells as “LOW”/“HIGH” outliers or “NO”,
4. Delegates to either `.computeCosmxTrainSet()` or `.computeXenMerTrainSet()` based on `metadata(spe)$technology`,
5. Deduplicates and down-samples “good” cells to match the number of “bad” cells.

Value

A `data.frame` with one row per cell, including:

- `qscore_train` (0/1) indicating “bad” vs “good”,
- relevant `colData` columns used for modeling.

Examples

```
example(spatialPerCellQC)
df_train <- computeTrainDF(spe, verbose = TRUE)
table(df_train$qscore_train)
```

`createPaletteFromColData`

createPaletteFromColData

Description

Create a Palette from `colData` in a `SpatialExperiment` Object

This function generates a palette mapping based on specified columns in the ‘`colData`’ of a ‘`SpatialExperiment`’ object.

Usage

```
createPaletteFromColData(spe, paletteNames, paletteColors)
```

Arguments

<code>spe</code>	A ‘SpatialExperiment‘ object with spatial transcriptomics data.
<code>paletteNames</code>	A character string specifying the column in ‘colData(spe)‘ to be used for the names in the palette.
<code>paletteColors</code>	A character string specifying the column in ‘colData(spe)‘ to be used for the colors in the palette.

Details

The function creates a new palette based on the unique combinations of values in the specified ‘`paletteNames`‘ and ‘`paletteColors`‘ columns in ‘`colData(spe)`‘.

Value

A character vector representing the palette mapping, where each element is a string in the format ‘“name=color”‘.

`dot-addFovFromTx` *.addFovFromTx*

Description

Add FOV information from transcript file to cell metadata.

This function retrieves FOV information from transcript file and appends the data to the resulting ‘`colData`‘.

Usage

```
.addFovFromTx(txFile, colData)
```

Arguments

<code>txFile</code>	‘character(1)‘ path to a Xenium Output tx file.
<code>colData</code>	A ‘ <code>DataFrame</code> ‘ containing the ‘ <code>colData</code> ‘ for the Xenium dataset.

Details

The function reads the transcript file then groups it by `cell_id` and merges the FOV information to the cell metadata in ‘`colData`‘. Only parquet file is supported for this operation

Value

A ‘`DataFrame`‘ containing the updated ‘`colData`‘ with FOV information.

```
dot-computeCosmxProteinTrainSet  
.computeCosmxProteinTrainSet
```

Description

Internal: Build Training Set for CosMx-Protein Splits a SpatialExperiment into “bad” vs “good” cells based on outliers in aspect ratio near tissue border or low count area.

Usage

```
.computeCosmxProteinTrainSet(spe)
```

Arguments

spe [SpatialExperiment](#)

Value

A list with elements bad and good, each a data.frame with qscore_train and (for “good”) an is_a_bad_boy flag.

```
dot-computeCosmxTrainSet  
.computeCosmxTrainSet
```

Description

Internal: Build Training Set for CosMx Splits a SpatialExperiment into “bad” vs “good” cells based on outliers in aspect ratio near tissue border or low count area.

Usage

```
.computeCosmxTrainSet(spe)
```

Arguments

spe [SpatialExperiment](#)

Value

A list with elements bad and good, each a data.frame with qscore_train and (for “good”) an is_a_bad_boy flag.

```
dot-computeXenMerTrainSet  
.computeXenMerTrainSet
```

Description

Internal: Build Training Set for Xenium & MERFISH Splits a SpatialExperiment into “bad” vs “good” cells based on pre-computed outlier labels on log2CountArea.

Usage

```
.computeXenMerTrainSet(spe)
```

Arguments

spe	SpatialExperiment
-----	-------------------

Value

A list with elements bad and good, each a data.frame with qscore_train and (for “good”) an is_a_bad_boy flag.

```
firstFlagPalette      firstFlagPalette
```

Description

neon color palette for firstFlagPlot

Usage

```
firstFlagPalette
```

Format

An object of class character of length 6.

Value

a palette for firstFlagPlot

<code>getFencesOutlier</code>	<i>getFencesOutlier</i>
-------------------------------	-------------------------

Description

Retrieve Threshold (Fence) Values from a SpatialExperiment Object

This function extracts the threshold values, also known as fences, from a specified column in the ‘colData‘ of a ‘SpatialExperiment‘ object.

Usage

```
getFencesOutlier(
  spe,
  fencesOf,
  highLow = c("both", "lower", "higher"),
  decimalRound = NULL
)
```

Arguments

<code>spe</code>	A ‘SpatialExperiment‘ object containing spatial transcriptomics data.
<code>fencesOf</code>	A character string specifying the name of the column in ‘colData(spe)‘ from which to extract the fence values. This column should contain an ‘outlier.filter‘ object (see ‘computeSpatialOutlier““).
<code>highLow</code>	character indicating which fence to get if "higher", "lower" or "both" (default is "both").
<code>decimalRound</code>	An optional integer specifying the number of decimal places to which the fence values should be rounded. If ‘NULL‘, no rounding is applied. Default is ‘NULL‘.

Value

A numeric vector containing the lower and upper threshold values extracted from the specified column.

Examples

```
example(computeSpatialOutlier)
getFencesOutlier(spe, fencesOf="log2CountArea_outlier_mc")
```

<code>getModelFormula</code>	<i>getModelFormula</i>
------------------------------	------------------------

Description

Returns the right-hand side of a model formula string based on technology.

Usage

```
getModelFormula(technology)
```

Arguments

`technology` `\[character\]` Technology name to decide which predictors to include.

Value

`\[character\]` A one-sided formula as a string (e.g. " $\sim \log2CountArea + \dots$ ").

Examples

```
example(spatialPerCellQC)
getModelFormula(metadata(spe)$technology)
```

<code>plotCellsFovs</code>	<i>plotCellsFovs</i>
----------------------------	----------------------

Description

Plot cell centroids in FoVs Creates a scatter plot with cell centroids arranged in their FoVs as an overlapping grid.

Usage

```
plotCellsFovs(
  spe,
  sampleId = unique(spe$sample_id),
  pointCol = "firebrick",
  numbersCol = "black",
  alphaNumbers = 0.8,
  fovDim = metadata(spe)$fov_dim
)
```

Arguments

<code>spe</code>	A ‘SpatialExperiment‘ object with ‘fov‘ in ‘colData‘.
<code>sampleId</code>	Character string identifying which sample to plot. Default: ‘unique(spe\$sample_id)‘.
<code>pointCol</code>	Color for the cell centroids. Default: ‘"firebrick"‘.
<code>numbersCol</code>	Color for the FoV labels. Default: ‘"black"‘.
<code>alphaNumbers</code>	Numeric transparency for FoV labels. Default: ‘0.8‘.
<code>fovDim</code>	numeric with two named dimensions xdim, ydim. (Default is <code>metadata(spe)\$fov_dim</code>)

Value

A ‘ggplot‘ object showing cell centroids and FoV boundaries.

Examples

```
example(readCosmxSPE)
g <- plotCellsFovs(spe)
print(g)
```

plotCentroids

plotCentroids

Description

Plot Spatial Coordinates for a SpatialExperiment Object This function generates a ggplot of spatial coordinates from a ‘SpatialExperiment‘ object, optionally coloring the points by a specified column in ‘colData‘.

Usage

```
plotCentroids(
  spe,
  colourBy = NULL,
  colourLog = FALSE,
  sampleId = unique(spe$sample_id),
  isNegativeProbe = FALSE,
  palette = NULL,
  pointCol = "darkmagenta",
  size = 0.05,
  alpha = 0.8,
  aspectRatio = 1
)
```

Arguments

<code>spe</code>	A ‘SpatialExperiment‘ object containing spatial transcriptomics data.
<code>colourBy</code>	An optional character string specifying the column in ‘colData(spe)‘ to use for coloring the points. If ‘NULL‘, all points will be colored the same.
<code>colourLog</code>	Logical to log-transform the data to enhance visualization (Default is FALSE).
<code>sampleId</code>	A character string specifying the sample identifier to be used as the plot title. (Default is the unique sample ID from ‘spe‘)
<code>isNegativeProbe</code>	A logical value indicating whether to apply a custom color gradient for negative probe data. (Default is ‘FALSE‘)
<code>palette</code>	A vector of colors to be used as a custom palette. For categorical data, this should be a vector of colors with the same length as the number of levels in ‘colourBy‘. For continuous data, this should be a vector of colors used to create a gradient.
<code>pointCol</code>	A character string specifying the color of the points when ‘colourBy‘ is ‘NULL‘. (Default is “darkmagenta”)

size	A numeric value specifying the size of the points. (Default is ‘0.05’)
alpha	A numeric value specifying the transparency level of the points. (Default is ‘0.2’)
aspectRatio	A numeric value specifying the aspect ratio of the plot. (Default is ‘1’)

Value

A ‘ggplot‘ object representing the spatial coordinates plot of polygon centroids.

Examples

```
example(readCosmxSPE)
g <- plotCentroids(spe, colourBy="Mean.DAPI")
print(g)
```

plotMetricHist

plotMetricHist

Description

Plot a Histogram for a Given Metric in a SpatialExperiment Object

This function generates a histogram for a specified metric in a ‘SpatialExperiment‘ object.

Usage

```
plotMetricHist(
  spe,
  metric,
  fillColor = "#c0c8cf",
  useFences = NULL,
  fencesColors = c(lower = "purple4", higher = "tomato"),
  bins = 30,
  binWidth = NULL
)
```

Arguments

spe	A ‘SpatialExperiment‘ object containing spatial transcriptomics data.
metric	A character string specifying the name of the metric (column in ‘colData(spe)‘) to plot.
fillColor	A character string specifying the fill color of the histogram bars. (Default is ‘#69b3a2’)
useFences	A character string specifying the name of the column in ‘colData(spe)‘ that contains the fence thresholds (typically from an outlier filter). If ‘NULL‘, no fences will be plotted. (Default is ‘NULL‘)
fencesColors	A named character vector specifying the colors to use for the lower and higher fences. The names should be “lower” and “higher”. (Default is ‘c("lower"="purple4", "higher"="tomato")’)
bins	An integer specifying the number of bins to use in the histogram. (Default is ‘30’)
binWidth	A numeric value specifying the width of the bins. If ‘NULL‘, the bin width will be automatically determined based on the ‘bins‘ parameter. (Default is ‘NULL‘)

Value

A ‘ggplot‘ object representing the histogram of the specified metric.

Examples

```
example(readCosmxSPE)
g <- plotMetricHist(spe, metric="Mean.DAPI")
print(g)
```

plotPolygons

plotPolygons

Description

Plot polygons from a ‘SpatialExperiment‘ object using ggplot2.

Usage

```
plotPolygons(
  spe,
  colourBy = "darkgrey",
  colourLog = FALSE,
  polyColumn = "polygons.global",
  sampleId = unique(spe$sample_id),
  bgColor = "white",
  fillAlpha = 1,
  palette = NULL,
  borderCol = NA,
  borderAlpha = 1,
  borderLineWidth = 0.1,
  drawBorders = TRUE
)
```

Arguments

<code>spe</code>	A ‘SpatialExperiment‘ object with polygon data as an ‘sf‘ object.
<code>colourBy</code>	A column in ‘colData(spe)‘ for coloring the polygons or a string color in colors(). (Default is "darkgrey")
<code>colourLog</code>	Logical to log-transform the data to enhance visualization (Default is FALSE).
<code>polyColumn</code>	character for the name of the column where the polygons sf are stored (default is "polygons.global")
<code>sampleId</code>	Sample ID for plot title. Default is the unique sample ID.
<code>bgColor</code>	character indicating color for the background (default is "white")
<code>fillAlpha</code>	Transparency level for polygon fill. Default is '1'.
<code>palette</code>	Colors to use if ‘colourBy‘ is a factor. Default is ‘NULL‘.
<code>borderCol</code>	Color of polygon borders. Default is ‘"black"‘.
<code>borderAlpha</code>	Transparency level for borders. Default is '1'.
<code>borderLineWidth</code>	Width of polygon borders. Default is '0.1'.
<code>drawBorders</code>	Logical; whether to draw borders. Default is ‘TRUE‘.

Value

A ‘ggplot‘ object representing the polygon plot of the spatial data.

Examples

```
example(readAndAddPolygonsToSPE)
plotPolygons(spe, colourBy="Mean.DAPI")
```

plotQScoreTerms

*plotQScoreTerms***Description**

Plots the individual terms that combine into the quality score formula, allowing assessment of each term’s impact on the final score.

Usage

```
plotQScoreTerms(
  spe,
  sampleId = unique(spe$sample_id),
  size = 0.05,
  alpha = 0.8,
  aspectRatio = 1,
  custom = FALSE
)
```

Arguments

<code>spe</code>	A ‘SpatialExperiment‘ object with ‘quality_score‘ and term columns in ‘colData‘.
<code>sampleId</code>	Character string for plot title. Must match values in the ‘fov‘ column of ‘colData(spe)‘. Default: ‘unique(spe\$sample_id)‘.
<code>size</code>	Numeric point size for the scatter plots. Default: ‘0.05‘.
<code>alpha</code>	Numeric transparency for the scatter plots. Default: ‘0.2‘.
<code>aspectRatio</code>	Numeric aspect ratio of the plots. Default: ‘1‘.
<code>custom</code>	Logical; if ‘TRUE‘, use custom polygon-derived metrics.

Value

A combined plot (via ‘cowplot::plot_grid‘) showing spatial maps of each QS term.

Examples

```
example(readAndAddPolygonsToSPE)
example(spatialPerCellQC)
p <- plotQScoreTerms(spe)
print(p)
```

`plotZoomFovsMap` *plotZoomFovsMap*

Description

Plot Zoomed-in FOVs with Map and Polygons

This function generates a plot that shows a map of all fields of view (FOVs) within a ‘SpatialExperiment‘ object, alongside a zoomed-in view of the specified FOVs with an overlay of polygons and optional coloring.

Usage

```
plotZoomFovsMap(
  spe,
  fovs = NULL,
  mapPointCol = "darkmagenta",
  mapNumbersCol = "black",
  mapAlphaNumbers = 0.8,
  title = NULL,
  ...
)
```

Arguments

<code>spe</code>	A ‘SpatialExperiment‘ object containing spatial transcriptomics data.
<code>fovs</code>	A character vector specifying the FOVs to be zoomed in and plotted. Must match values in the ‘fov‘ column of ‘colData(spe)‘.
<code>mapPointCol</code>	A character string specifying the color of the points in the map. Default is “darkmagenta”.
<code>mapNumbersCol</code>	A character string specifying the color of the numbers on the map. Default is “black”.
<code>mapAlphaNumbers</code>	A numeric value specifying the transparency of the numbers on the map. Default is ‘0.8’.
<code>title</code>	An optional character string specifying the title of the final plot. If ‘NULL‘, no title is added. Default is ‘NULL‘.
<code>...</code>	Additional arguments passed to ‘plotPolygons‘.

Details

The function first filters the ‘SpatialExperiment‘ object to the specified FOVs, generates a plot of the cells for the entire map, then creates a detailed polygon plot of the selected FOVs, and finally combines these into a single side-by-side visualization. If ‘title‘ is not ‘NULL‘, it adds a title to the combined plot.

Value

A combined plot showing a map of all FOVs with zoomed-in views of the specified FOVs and their associated polygons.

Examples

```
example(readAndAddPolygonsToSPE)
plotZoomFovsMap(spe, fovs=16, title="FOV 16")
```

`qcFlagPlots`

qcFlagPlots

Description

Plots the flagged cells identified with first filter, based on control count on total count ratio, area in um and DAPI signal.

This function generates a plot that shows selected (FOVs) within a ‘SpatialExperiment‘ object, with cells flagged in different colors over a light or dark layout chosen by the user.

Usage

```
qcFlagPlots(
  spe,
  fov = unique(spe$fov),
  theme = c("light", "dark"),
  custom = FALSE
)
```

Arguments

<code>spe</code>	A ‘SpatialExperiment‘ object containing spatial transcriptomics data.
<code>fov</code>	An integer or numeric vector specifying the FOVs to be plotted Must match values in the ‘fov‘ column of ‘colData(spe)‘.
<code>theme</code>	A character string among "light" or "dark".
<code>custom</code>	A boolean value. If TRUE, custom polygons derived metrics will be used.

Value

A panel with multiple plots showing flagged cells for different variables.

Examples

```
example(readAndAddPolygonsToSPE)
spe <- spatialPerCellQC(spe)
spe <- computeThresholdFlags(spe)
p <- qcFlagPlots(spe, fov=16, theme="dark")
print(p)
```

readAndAddPolygonsToSPE
readAndAddPolygonsToSPE

Description

Read and Add Polygons to a SpatialExperiment Object

This function reads polygon boundary data based on the technology associated with the provided SpatialExperiment (SPE) object and adds the polygons to the SPE.

Usage

```
readAndAddPolygonsToSPE(
  spe,
  polygonsCol = "polygons",
  keepMultiPol = TRUE,
  boundariesType = c("csv", "HDF5", "parquet")
)
```

Arguments

<code>spe</code>	A SpatialExperiment object. The object should contain metadata with the field "technology", specifying the technology used (e.g., "Nanostring_CosMx", "Vizgen_MERFISH", or "10X_Xenium").
<code>polygonsCol</code>	character indicating the name of the polygons column to add into the colData (default is 'polygons').
<code>keepMultiPol</code>	Logical. If TRUE, multi-polygon features will be kept when reading the boundary data. Defaults to TRUE.
<code>boundariesType</code>	Character. Specifies the type of boundary file format to read. Options are "HDF5" or "parquet". Defaults to "HDF5".

Value

A SpatialExperiment object with the added polygon data.

Examples

```
example(readCosmxSPE)
spe <- readAndAddPolygonsToSPE(spe)
colData(spe)
```

readCosmxSPE

readCosmxSPE

Description

Read and Construct a SpatialExperiment Object from CosMx Data

This function reads in data from Nanostring CosMx files and constructs a ‘SpatialExperiment‘ object, optionally including polygon data.

Usage

```
readCosmxSPE(
  dirName,
  sampleName = "sample01",
  coordNames = c("CenterX_global_px", "CenterY_global_px"),
  countMatFPattern = "exprMat_file.csv",
  metadataFPattern = "metadata_file.csv",
  polygonsFPattern = "polygons.csv",
  fovPosFPattern = "fov_positions_file.csv",
  fovdims = c(xdim = 4256, ydim = 4256)
)
```

Arguments

<code>dirName</code>	A character string specifying the directory containing the CosMx data files.
<code>sampleName</code>	A character string specifying the sample name. Default is “sample01”.
<code>coordNames</code>	A character vector specifying the names of the spatial coordinate columns in the data. Default is ‘c("CenterX_global_px", "CenterY_global_px")’.
<code>countMatFPattern</code>	A character string specifying the pattern to match the count matrix file. Default is “exprMat_file.csv”.
<code>metadataFPattern</code>	A character string specifying the pattern to match the metadata file. Default is “metadata_file.csv”.
<code>polygonsFPattern</code>	A character string specifying the pattern to match the polygons file. Default is “polygons.csv”.
<code>fovPosFPattern</code>	A character string specifying the pattern to match the FOV positions file. Default is “fov_positions_file.csv”.
<code>fovdims</code>	A named numeric vector specifying the dimensions of the FOV in pixels. Default is ‘c(xdim=4256, ydim=4256)’.

Details

The function reads in the specified files for count matrices, metadata, and FOV positions, and constructs a ‘SpatialExperiment‘ object. Optionally, polygon data can be read and added to the object.

`readCosmxProteinSPE` is a wrapper of `readCosmxSPE`, it only changes the technology metadata in `Nanostring_CosMx_Protein`.

Value

A ‘SpatialExperiment’ object containing the read CosMx data, including count matrices, metadata, and optionally polygons.

Author(s)

Dario Righelli, Benedetta Banzi

Examples

```
cospath <- system.file(file.path("extdata", "CosMx_DBKero_Tiny"),
  package="SpaceTrooper")
spe <- readCosmxSPE(cospath, sampleName="DBKero_Tiny")
```

<code>readh5polygons</code>	<i>readh5polygons</i>
-----------------------------	-----------------------

Description

This function reads polygon data from an HDF5 file.

Usage

```
readh5polygons(polFile)
```

Arguments

<code>polFile</code>	A character string specifying the file path to the HDF5 polygon data.
----------------------	---

Value

A list containing the polygon geometries and their associated cell_id

Author(s)

Lambda Moses

<code>readMerfishSPE</code>	<i>readMerfishSPE</i>
-----------------------------	-----------------------

Description

‘readMerfishSPE()’ imports MERFISH/Merscore outputs (counts, metadata, and optionally cell boundary polygons) from a directory and builds a SpatialExperiment object.

Usage

```
readMerfishSPE(
  dirName,
  sampleName = "sample01",
  computeMissingMetrics = TRUE,
  keepPolygons = FALSE,
  boundariesType = c("HDF5", "parquet"),
  countmatFPattern = "cell_by_gene.csv",
  metadataFPattern = "cell_metadata.csv",
  polygonsFPattern = "cell_boundaries.parquet",
  coordNames = c("center_x", "center_y"),
  polygonsCol = "polygons"
)
```

Arguments

dirName	'character(1)' Path to a folder containing MERFISH output files.
sampleName	'character(1)' Identifier to assign to the 'sample_id' field in the returned object. Default: "sample01".
computeMissingMetrics	'logical(1)' If 'TRUE', compute area and aspect-ratio metrics from the cell boundary polygons. Default: 'TRUE'.
keepPolygons	'logical(1)' If 'TRUE', attach raw polygon geometries as extra columns in 'colData'. Default: 'FALSE'.
boundariesType	'character(1)' One of "HDF5" or "parquet". If "HDF5", uses a folder of HDF5 polygon files; if "parquet", reads a single Parquet file of boundaries.
countmatFPattern	'character(1)' Regex passed to 'list.files()' to find the count matrix CSV. Default: "cell_by_gene.csv".
metadataFPattern	'character(1)' Pattern to find the cell metadata CSV. Default: "cell_metadata.csv".
polygonsFPattern	'character(1)' Pattern to find the cell boundaries file. Default: "cell_boundaries.parquet".
coordNames	'character(2)' Names of the columns in 'colData' that store X/Y spatial coordinates. Default: 'c("center_x", "center_y")'.
polygonsCol	character indicating the name of the polygons column to add into the colData (default is 'polygons').

Value

A 'SpatialExperiment' object with:

- 'assays\$counts': gene × cell count matrix
- 'colData': per-cell metadata (including computed metrics)
- spatial coordinates named by 'coordNames'
- 'metadata\$polygons': path to the boundaries file
- 'metadata\$technology': "Vizgen_MERFISH"

Author(s)

Dario Righelli, Benedetta Banzi

Examples

```
path <- system.file("extdata", "Merfish_Tiny",
                     package = "SpaceTrooper")
spe <- readMerfishSPE(
  dirName = path,
  sampleName = "Patient2",
  keepPolygons = TRUE,
  boundariesType = "parquet"
)
spe
```

readPolygons

readPolygons

Description

Read and Validate Polygons from a File

This function reads polygon data from a specified file, validates the polygons, and returns them as an ‘sf’ object. It supports multiple file formats and can handle both global and local coordinates.

Usage

```
readPolygons(
  polygonsFile,
  type = c("csv", "parquet", "h5"),
  x = c("x_global_px", "vertex_x"),
  y = c("y_global_px", "vertex_y"),
  xloc = "x_local_px",
  yloc = "y_local_px",
  keepMultiPol = TRUE,
  verbose = FALSE
)
```

Arguments

<code>polygonsFile</code>	A character string specifying the path to the polygon file.
<code>type</code>	A character string specifying the file type. Supported types are ““csv”“, ““parquet”“, and ““h5”“. Default is ““csv”“.
<code>x</code>	A character vector specifying the column names for the x-coordinates in the polygon data. Default is ‘c(“x_global_px”, “vertex_x”)’.
<code>y</code>	A character vector specifying the column names for the y-coordinates in the polygon data. Default is ‘c(“y_global_px”, “vertex_y”)’.
<code>xloc</code>	A character string specifying the column name for the local x-coordinates. Default is ““x_local_px”“.
<code>yloc</code>	A character string specifying the column name for the local y-coordinates. Default is ““y_local_px”“.
<code>keepMultiPol</code>	A logical value indicating whether to keep multipolygons during validation. Default is ‘TRUE’.
<code>verbose</code>	A logical value indicating whether to print additional information during processing. Default is ‘FALSE’.

Details

The function reads polygon data from the specified file and formats. It validates the polygons and handles both global and local coordinates if provided. If the file type is "h5", the function currently does not handle the data, as this part of the code is not implemented.

Value

An 'sf' object with the loaded and validated polygons.

Examples

```
example(readCosmxSPE)
polygons <- readPolygons(metadata(spe)$polygons)
polygons
```

`readPolygonsCosmx` *readPolygonsCosmx*

Description

This function reads polygon data specific to CosMx technology.

Usage

```
readPolygonsCosmx(
  polygonsFile,
  type = c("csv", "parquet"),
  x = "x_global_px",
  y = "y_global_px",
  xloc = "x_local_px",
  yloc = "y_local_px",
  keepMultiPol = TRUE,
  verbose = FALSE
)
```

Arguments

<code>polygonsFile</code>	A character string specifying the file path to the polygon data.
<code>type</code>	A character string specifying the file type ("csv" or "parquet").
<code>x</code>	A character string specifying the x-coordinate column.
<code>y</code>	A character string specifying the y-coordinate column.
<code>xloc</code>	A character string specifying the local x-coordinate column.
<code>yloc</code>	A character string specifying the local y-coordinate column.
<code>keepMultiPol</code>	A logical value indicating whether to keep multipolygons.
<code>verbose</code>	A logical value indicating whether to print additional information.

Value

An 'sf' object containing the CosMx polygon data.

Examples

```
example(readCosmxSPE)
polygons <- readPolygonsCosmx(metadata(spe)$polygons)
polygons
```

readPolygonsMerfish *readPolygonsMerfish*

Description

This function reads polygon data specific to MERFISH technology.

Usage

```
readPolygonsMerfish(
  polygons,
  type = c("parquet", "HDF5"),
  keepMultiPol = TRUE,
  hdf5pattern = "hdf5",
  zLev = 3L,
  zcolumn = "ZIndex",
  geometry = "Geometry",
  verbose = FALSE
)
```

Arguments

<code>polygons</code>	A character string specifying the folder containing the polygon data files in case of HDF5, or a path to a parquet file (see ‘type’).
<code>type</code>	A character string specifying the file type("HDF5" or "parquet"). Default is parquet.
<code>keepMultiPol</code>	A logical value indicating whether to keep multipolygons.
<code>hdf5pattern</code>	A character string specifying the pattern to match HDF5 files.
<code>zLev</code>	An integer specifying the Z level to filter the data. Default is ‘3L’.
<code>zcolumn</code>	A character string specifying the column name for the Z index.
<code>geometry</code>	A character string specifying the geometry column name.
<code>verbose</code>	A logical value indicating whether to print additional information.

Value

An ‘sf’ object containing the MERFISH polygon data.

Examples

```
example(readMerfishSPE)
polygons <- readPolygonsMerfish(metadata(spe)$polygons, type="parquet")
polygons
```

<code>readPolygonsXenium</code>	<i>readPolygonsXenium</i>
---------------------------------	---------------------------

Description

This function reads polygon data specific to Xenium technology.

Usage

```
readPolygonsXenium(
  polygonsFile,
  type = c("parquet", "csv"),
  x = "vertex_x",
  y = "vertex_y",
  keepMultiPol = TRUE,
  verbose = FALSE
)
```

Arguments

<code>polygonsFile</code>	A character string specifying the file path to the polygon data.
<code>type</code>	A character string specifying the file type ("parquet" or "csv"). Default is parquet.
<code>x</code>	A character string specifying the x-coordinate column.
<code>y</code>	A character string specifying the y-coordinate column.
<code>keepMultiPol</code>	A logical value indicating whether to keep multipolygons.
<code>verbose</code>	A logical value indicating whether to print additional information.

Value

An ‘sf’ object containing the Xenium polygon data.

Examples

```
example(readXeniumSPE)
polygons <- readPolygonsXenium(metadata(spe)$polygons, type="parquet")
polygons
```

<code>readXeniumSPE</code>	<i>Load data from a 10x Genomics Xenium experiment</i>
----------------------------	--

Description

Creates a [‘SpatialExperiment’] from an unzipped Xenium Output Bundle directory containing spatial gene expression data.

Usage

```
readXeniumSPE(
  dirName,
  sampleName = "sample01",
  type = c("HDF5", "sparse"),
  coordNames = c("x_centroid", "y_centroid"),
  boundariesType = c("parquet", "csv"),
  computeMissingMetrics = TRUE,
  keepPolygons = FALSE,
  countsFilePattern = "cell_feature_matrix",
  metadataFPattern = "cells.csv.gz",
  polygonsFPattern = "cell_boundaries",
  polygonsCol = "polygons",
  txPattern = "transcripts",
  addFOVs = FALSE
)
```

Arguments

dirName	'character(1)'	Path to a Xenium Output Bundle directory.
sampleName	'character(1)'	Sample identifier to assign to 'sample_id'. Default: "sample01".
type	'character(1)'	One of "HDF5" or "sparse"; method to read the feature matrix.
coordNames	'character(2)'	Names of X/Y spatial coordinate columns. Default: 'c("x_centroid", "y_centroid")'.
boundariesType	'character(1)'	One of "parquet" or "csv"; format of the polygon file.
computeMissingMetrics	'logical(1)'	If 'TRUE', compute area and aspect-ratio from boundary polygons.
keepPolygons	'logical(1)'	If 'TRUE', append raw polygon geometries to 'colData'.
countsFilePattern	'character(1)'	Pattern to locate the feature matrix file. Default: "cell_feature_matrix".
metadataFPattern	'character(1)'	Pattern to locate the cell metadata file. Default: "cells".
polygonsFPattern	'character(1)'	Pattern to locate the cell boundaries file. Default: "cell_boundaries".
polygonsCol	'character(1)'	Name of the polygons column to add to 'colData'. Default: "polygons".
txPattern	'character(1)'	Pattern (base filename, without extension) to locate the transcript file (usually a '.parquet' file) from which to extract Field-Of-View (FOV) information for each cell. Default: "transcripts".
addFOVs	'logical(1)'	If 'TRUE', extract Field-Of-View (FOV) information from the transcript file (as located by 'txPattern') and append it to cell metadata ('colData'). Default: 'FALSE'.

Details

readXeniumSPE

Expects the unzipped bundle to contain an 'outs/' folder with:

- 'cell_feature_matrix.h5' or 'cell_feature_matrix/'
- 'cells.csv.gz'

Value

A [‘SpatialExperiment’] object with assays, ‘colData’, spatial coordinates, and ‘metadata\$polygons’ & ‘metadata\$technology’.

Author(s)

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Examples

```
xpath <- system.file(
  "extdata", "Xenium_small", package = "SpaceTrooper"
)
(spe <- readXeniumSPE(
  dirName = xpath,
  keepPolygons = TRUE
))
```

`spatialPerCellQC` *spatialPerCellQC*

Description

Computes quality-control metrics for each cell and adds them to ‘colData’.

Usage

```
spatialPerCellQC(
  spe,
  micronConvFact = 0.12,
  rmZeros = TRUE,
  negProbList = c("NegPrb", "Negative", "SystemControl", "Ms IgG1", "Rb IgG", "BLANK_",
    "NegControlProbe", "NegControlCodeword", "UnassignedCodeword", "Blank"),
  use_altexps = NULL
)
```

Arguments

<code>spe</code>	A ‘SpatialExperiment’ object containing spatial data.
<code>micronConvFact</code>	Numeric factor to convert pixels to microns. Default ‘0.12’.
<code>rmZeros</code>	logical for removing zero counts cells (default is TRUE).
<code>negProbList</code>	Character vector of patterns to identify negative probes. Defaults include: - Nanostring CosMx: “NegPrb”, “Negative”, “SystemControl” - Xenium: “NegControlProbe”, “NegControlCodeword”, “UnassignedCodeword” - MER-FISH: “Blank”
<code>use_altexps</code>	logical for ‘use_altexps’ in ‘scuttle’ package. If TRUE uses the altexps for computing some metrics on it. Useful for interoperability with ‘SpatialExperimentIO’. (See addPerCellQC for additional details).

Details

Calculates sums and detected counts for control and target probes, computes ratio and count-area metrics, converts coords to microns for CosMx, and drops zero-count cells.

Value

A ‘SpatialExperiment‘ object with added QC metrics in ‘colData‘.

Examples

```
example(readCosmxSPE)
spe <- spatialPerCellQC(spe)
```

trainModel

trainModel

Description

Fit a Ridge Logistic Regression Model

`trainModel` fits an L2-regularized (ridge) logistic regression using `glmnet`, given a design matrix and a training data frame.

Usage

```
trainModel(modelMatrix, trainDF)
```

Arguments

- | | |
|--------------------------|---|
| <code>modelMatrix</code> | a matrix describing the model variables, typically created with ‘ <code>getModelFormula</code> ‘ and ‘ <code>model.matrix</code> ‘ functions. |
| <code>trainDF</code> | \[<code>data.frame</code> \] A data frame containing at least the response column <code>qscore_train</code> , coded as 0/1. |

Value

A `glmnet` model object fitted with `family="binomial"`, `alpha=0` (ridge), and a sequence of λ values.

Examples

```
example(computeTrainDF)
model_formula <- getModelFormula(metadata(spe)$technology)
model_matrix <- model.matrix(as.formula(model_formula), data=df_train)
fit <- trainModel(model_matrix, df_train)
coef(fit, s = 0.01)
```

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