

# Package ‘SpatialFeatureExperiment’

May 2, 2026

**Type** Package

**Title** Integrating SpatialExperiment with Simple Features in sf

**Version** 1.15.0

**Description** A new S4 class integrating Simple Features with the R package sf to bring geospatial data analysis methods based on vector data to spatial transcriptomics. Also implements management of spatial neighborhood graphs and geometric operations. This package builds upon SpatialExperiment and SingleCellExperiment, hence methods for these parent classes can still be used.

**Imports** Biobase, BiocGenerics (>= 0.51.2), BiocNeighbors, BiocParallel, data.table, DropletUtils, EBImage, grDevices, lifecycle, Matrix, methods, rjson, rlang, S4Vectors, sf, sfheaders, SingleCellExperiment, SpatialExperiment, spatialreg, spdep (>= 1.1-7), SummarizedExperiment, stats, terra, utils, zeallot

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---

addVisiumSpotPoly      *Add Visium spot polygons to colGeometry*

---

## Description

For adding the spot polygons to SFE objects converted from SPE.

## Usage

```
addVisiumSpotPoly(x, spotDiameter)
```

## Arguments

x	A SpatialFeatureExperiment object.
spotDiameter	Spot diameter for technologies with arrays of spots of fixed diameter per slide, such as Visium, ST, DBiT-seq, and slide-seq. The diameter must be in the same unit as the coordinates in the *Geometry arguments. Ignored for geometries that are not POINT or MULTIPOINT.

## Value

A SFE object with a new colGeometry called spotPoly, which has polygons of the spots.

## Examples

```
library(VisiumIO)
# From examples of TENxVisium()
sample_dir <- system.file(
  file.path("extdata", "10xVisium", "section1"),
  package = "VisiumIO"
)
## using spacerangerOut folder
tv <- TENxVisium(
  spacerangerOut = file.path(sample_dir, "outs"), processing = "raw", images = "lowres"
)
spe <- import(tv)
# There can't be suplicate barcodes
colnames(spe) <- make.unique(colnames(spe), sep = "-")
rownames(spatialCoords(spe)) <- colnames(spe)
sfe <- toSpatialFeatureExperiment(spe)
# A hypothetical spot diameter; check the scalefactors_json.json file for
# actual diameter in pixels in full resolution image.
sfe <- addVisiumSpotPoly(sfe, spotDiameter = 80)
```

---

affineImg	<i>Affine transformation of images</i>
-----------	--

---

### Description

This function performs affine transformation on images, with any matrix and translation vector.

### Usage

```
## S4 method for signature 'SpatRasterImage'  
affineImg(x, M, v, maxcell = 1e+07, ...)  
  
## S4 method for signature 'BioFormatsImage'  
affineImg(x, M, v, ...)  
  
## S4 method for signature 'ExtImage'  
affineImg(x, M, v, ...)
```

### Arguments

x	An object of class *Image as implemented in this package.
M	A 2x2 numeric matrix for the linear transformation in the xy plane.
v	A numeric vector of length 2 for translation in the xy plane.
maxcell	Max number of pixels to load SpatRasterImage into memory. The default 1e7 is chosen because this is the approximate number of pixels in the medium resolution image at resolution = 4L in Xenium OME-TIFF to make different methods of this function consistent.
...	Ignored. It's there so different methods can all be passed to the same lapply in the method for SFE objects. Some methods have extra arguments.

### Value

SpatRasterImage will be converted to ExtImage. Otherwise \*Image object of the same class. For BioFormatsImage, the transformation info is stored and will be applied when the image is loaded into memory as ExtImage.

### See Also

Other image methods: [SFE-image](#), [cropImg\(\)](#), [dim](#), [BioFormatsImage-method](#), [dim](#), [ExtImage-method](#), [ext\(\)](#), [imgRaster\(\)](#), [imgSource\(\)](#), [mirrorImg\(\)](#), [rotateImg\(\)](#), [scaleImg\(\)](#), [translateImg\(\)](#), [transposeImg\(\)](#)

---

 aggBboxes

*Aggregate bounding boxes*


---

### Description

To find the bounding box of multiple bounding boxes.

### Usage

```
aggBboxes(bboxes)
```

### Arguments

**bboxes** Either a matrix with 4 rows whose columns are the different bounding boxes, with row names "xmin", "xmax", "ymin", and "ymax" in any order, or a list of bounding boxes which are named numeric vectors.

### Value

A named numeric vector for the total bounding box.

### Examples

```
bboxes <- list(c(xmin = 5, xmax = 10, ymin = 2, ymax = 20),
              c(xmin = 8, xmax = 18, ymin = 0, ymax = 15))
bbox_all <- aggBboxes(bboxes)
```

---

 aggregate, SpatialFeatureExperiment-method

*Aggregate data in SFE using geometry*


---

### Description

Gene expression and numeric columns of colData will be aggregated with the function specified in FUN, according to another geometry supplied and a geometry predicate (such as st\_intersects). For example, when the predicate is st\_intersects and a spatial grid is used to aggregate, then the data associated with all cells that intersect with each grid cell will be aggregated with FUN, such as mean or sum. The categorical columns will be collected into list columns, and logical columns will be converted into numeric before applying FUN.

### Usage

```
## S4 method for signature 'SpatialFeatureExperiment'
aggregate(
  x,
  by = NULL,
  FUN = sum,
  sample_id = "all",
  colGeometryName = 1L,
  rowGeometryName = NULL,
```

```

    cellsize = NULL,
    square = TRUE,
    flat_topped = FALSE,
    new_geometry_name = "bins",
    join = st_intersects,
    sparse = FALSE,
    BPPARAM = SerialParam()
)

```

## Arguments

x	An SFE object to be aggregated.
by	A sf data frame whose geometry column is used for aggregation or sfc or for multiple samples a list of sfc whose names are the sample IDs. For multiple samples, the sf data frame must have a column sample_id to indicate which geometry for which sample. This argument is optional if cellsize is specified.
FUN	Function to aggregate the numerical columns in colData and the gene count matrix. This can be sum, mean, or any function that takes a numeric matrix as input and returns a numeric vector whose length is same as the number of rows in the input matrix, such as rowMedians. See package matrixStats. Depending on the function used for aggregation, numeric columns of colData may need to be interpreted differently after aggregation. Aggregation is not done when aggregating by transcript spots in rowGeometry. When it's sum or mean, matrix multiplication is used for aggregation rather than calling the sum or mean function itself; this is much faster than looping through the bins and calling the function on each of them.
sample_id	Which samples to aggregate, defaults to "all".
colGeometryName	Which colGeometry to spatially aggregate the data, by default the first one.
rowGeometryName	Which rowGeometry to spatially aggregate
cellsize	numeric of length 1 or 2 with target cellsize: for square or rectangular cells the width and height, for hexagonal cells the distance between opposite edges (edge length is cellsize/sqrt(3)). A length units object can be passed, or an area unit object with area size of the square or hexagonal cell.
square	logical; if FALSE, create hexagonal grid
flat_topped	logical; if TRUE generate flat topped hexagons, else generate pointy topped
new_geometry_name	Name to give to the new colGeometry in the output. Defaults to "bins".
join	logical spatial predicate function to use if by is a simple features object or geometry; see <a href="#">st_join</a>
sparse	Logical, whether the gene count matrix from aggregating transcript spots should be sparse. When the bins are large, the matrix will not be very sparse so using sparse matrix will not save memory, but when the bins are small, sparsity is worth it.
BPPARAM	A <a href="#">BiocParallelParam</a> object specifying parallel computing when aggregating data with functions other than sum and mean when aggregating cells. When aggregating transcript spots, this specifies parallel computing over genes. Defaults to SerialParam().

**Details**

For smFISH-based data where the transcript spots are available, the transcript spots can be used instead of cells to aggregate the gene count matrix, in which case all assays other than counts will be dropped and FUN only applies to colData because the transcript spots are simply counted.

What this function does is similar to **SEraster** but more general because any geometry and more aggregation function can be used, not just regular grids, and the aggregation can be performed on the transcript spots.

**Value**

An SFE object with colGeometry the same as the geometry specified in by or same as the grid specified in cellsize. rowGeometries and rowData remain the same as in the input x. reducedDims, localResults, colFeatureData (and its colGeometry, annotGeometry, and reducedDim counterparts), and spatialGraphs are dropped because those results no longer apply after aggregation.

**Note**

For developers: When debugging this function after calling devtools::load\_all("."), you may get an error that comes from S3 dispatch of aggregate.Vector from the S4Vectors package. When that happens, either restart the R session, or run setGeneric("aggregate", function(x, ...) standardGeneric("aggregate")) in the console to make an S4 generic as done in the terra package to prioritize S4 dispatch.

**Examples**

```
# example code
```

---

```
aggregateTx
```

```
Aggregate transcript spots from file
```

---

**Description**

This function reads the transcript spot file from the standard output of the commercial technologies (not GeoParquet) for spatial aggregation where the spots are assigned to polygons such as cells or spatial bins. Presets for Xenium, MERFISH, and CosMX are available. For Vizgen and Xenium, the images can be added when add\_images = TRUE.

**Usage**

```
aggregateTx(
  file,
  df = NULL,
  by = NULL,
  sample_id = "sample01",
  spatialCoordsNames = c("X", "Y", "Z"),
  gene_col = "gene",
  phred_col = "qv",
  min_phred = 20,
  flip_geometry = FALSE,
  cellsize = NULL,
```

```

square = TRUE,
flat_topped = FALSE,
new_geometry_name = "bins",
unit = "micron",
sparse = FALSE,
BPPARAM = SerialParam(),
save_memory = FALSE,
.orig_nrows = NULL
)

aggregateTxTech(
  data_dir,
  df = NULL,
  by = NULL,
  tech = c("Vizgen", "Xenium", "CosMX"),
  sample_id = "sample01",
  image = NULL,
  min_phred = 20,
  flip = c("geometry", "image", "none"),
  max_flip = "50 MB",
  cellsize = NULL,
  square = TRUE,
  flat_topped = FALSE,
  new_geometry_name = "bins",
  sparse = FALSE,
  BPPARAM = SerialParam(),
  save_memory = FALSE
)

```

### Arguments

file	File with the transcript spot coordinates. Should be one row per spot when read into R and should have columns for coordinates on each axis, gene the transcript is assigned to, and optionally cell the transcript is assigned to. Must be csv, tsv, or parquet.
df	If the file is already loaded into memory, a data frame (sf) with columns for the x, y, and optionally z coordinates and gene assignment of each transcript spot. If specified, then argument file will be ignored.
by	A sfc or sf object for spatial aggregation.
sample_id	Which sample in the SFE object the transcript spots should be added to.
spatialCoordsNames	Column names for the x, y, and optionally z coordinates of the spots. The defaults are for Vizgen.
gene_col	Column name for genes.
phred_col	Column name for Phred scores of the spots.
min_phred	Minimum Phred score to keep spot. By default 20, the conventional threshold indicating "acceptable", meaning that there's 1 chance that the spot was decoded in error.
flip_geometry	Logical, whether to flip the transcript spot geometries to match the images if added later.

cellsize	numeric of length 1 or 2 with target cellsize: for square or rectangular cells the width and height, for hexagonal cells the distance between opposite edges (edge length is $\text{cellsize}/\sqrt{3}$ ). A length units object can be passed, or an area unit object with area size of the square or hexagonal cell.
square	logical; if FALSE, create hexagonal grid
flat_topped	logical; if TRUE generate flat topped hexagons, else generate pointy topped
new_geometry_name	Name to give to the new colGeometry in the output. Defaults to "bins".
unit	Unit the coordinates are in, either microns or pixels in full resolution image.
sparse	Logical, whether the gene count matrix from aggregating transcript spots should be sparse. When the bins are large, the matrix will not be very sparse so using sparse matrix will not save memory, but when the bins are small, sparsity is worth it.
BPPARAM	bpparam object to specify parallel computing over genes. If a lot of memory is used, then stick to 'SerialParam()'. Using multicore in the R level when 'save_memory = TRUE' works and results into faster computation.
save_memory	Logical, if TRUE, then the transcript spots will not all be loaded into memory. <a href="#">open_dataset</a> is used to open a link to the data and then transcript spots of one gene is loaded into memory at a time.
.orig_nrows	Only used internally in the SFE method of aggregate
data_dir	Top level output directory.
tech	Which technology whose output to read, must be one of "Vizgen", "Xenium", or "CosMX" though more technologies may be added later.
image	String, which image(s) to add to the output SFE object. Not applicable to CosMX. See <a href="#">readVizgen</a> and <a href="#">readXenium</a> for options and multiple images can be specified. If NULL, then the default from the read function for the technology will be used.
flip	Logical, whether to flip the geometry to match image. Here the y coordinates are simply set to -y, so the original bounding box is not preserved. This is consistent with <a href="#">readVizgen</a> and <a href="#">readXenium</a> .
max_flip	Maximum size of the image allowed to flip the image. Because the image will be loaded into memory to be flipped. If the image is larger than this size then the coordinates will be flipped instead.

### Value

A SFE object with count matrix for number of spots of each gene in each geometry. Geometries with no spot are removed.

### Note

The resulting SFE object often includes geometries (e.g. grid cells) outside tissue, because there can be transcript spots detected outside the tissue. Also, bins at the edge of the tissue that don't fully overlap with the tissue will have lower transcript counts; this may have implications to downstream spatial analyses.

---

annotGeometries      *Annotation geometry methods*

---

## Description

"Annotation geometry" refers to Simple Feature (sf) geometries NOT associated with rows (features, genes) or columns (cells or spots) of the gene count matrix in the `SpatialFeatureExperiment` object. So there can be any number of rows in the sf data frame specifying the geometry. Examples of such geometries are tissue boundaries, pathologist annotation of histological regions, and objects not characterized by columns of the gene count matrix (e.g. nuclei segmentation in a Visium dataset where the columns are Visium spots). This page documents getters and setters for the annotation geometries. Internally, annotation geometries are stored in `int_metadata`.

## Usage

```
## S4 method for signature 'SpatialFeatureExperiment'
annotGeometries(x)

## S4 replacement method for signature 'SpatialFeatureExperiment'
annotGeometries(x, translate = TRUE, ...) <- value

## S4 method for signature 'SpatialFeatureExperiment'
annotGeometryNames(x)

## S4 replacement method for signature 'SpatialFeatureExperiment,character'
annotGeometryNames(x) <- value

## S4 method for signature 'SpatialFeatureExperiment'
annotGeometry(x, type = 1L, sample_id = NULL)

## S4 replacement method for signature 'SpatialFeatureExperiment'
annotGeometry(x, type = 1L, sample_id = NULL, translate = TRUE, ...) <- value

tissueBoundary(x, sample_id = 1L)

tissueBoundary(x, sample_id = 1L, translate = TRUE, ...) <- value
```

## Arguments

<code>x</code>	A <code>SpatialFeatureExperiment</code> object.
<code>translate</code>	Logical. Only used if <code>removeEmptySpace</code> has been run on the SFE object. If that's the case, this argument indicates whether the new value to be assigned to the geometry is in the coordinates prior to removal of empty space so it should be translated to match the new coordinates after removing empty space. Default to TRUE.
<code>...</code>	<code>spatialCoordsNames</code> , <code>spotDiameter</code> , <code>geometryType</code> passed to <code>df2sf</code> . Defaults are the same as in <code>df2sf</code> . For <code>dimGeometries&lt;-</code> only: <code>geometryType</code> can be a character vector of the geometry type of each data frame in the list of the same length as the list if the data frames specify different types of geometries.

value	Value to set. For <code>annotGeometry</code> , must be a <code>sf</code> data frame, or an ordinary data frame that can be converted to a <code>sf</code> data frame (see <a href="#">df2sf</a> ). For <code>annotGeometries</code> , must be a list of such <code>sf</code> or ordinary data frames. There must be a column <code>sample_id</code> to indicate the sample the geometries are for, and the <code>sample_id</code> must also appear in <code>colData</code> .
type	An integer specifying the index or string specifying the name of the <code>*Geometry</code> to query or replace. If missing, then the first item in the <code>*Geometries</code> will be returned or replaced.
sample_id	Sample ID to get or set geometries.

### Details

Wrapper for getter and setter of special geometry:

**tisseuBoundary** Boundary of the tissue of interest, including holes. This is usually of geometry type MULTIPOLYGON, though geometries in `annotGeometries` can have any type supported by `sf`.

### Value

Getters for multiple geometries return a named list. Getters for names return a character vector of the names. Getters for single geometries return an `sf` data frame. Setters return an SFE object.

### Examples

```
# Example dataset
library(SFEData)
sfe_small <- McKellarMuscleData(dataset = "small")

# Get all annotation geometries, returning a named list
annotGeometries(sfe_small)

# Set all annotation geometries, in a named list
toy <- readRDS(system.file("extdata/sfe_toy.rds",
  package = "SpatialFeatureExperiment"
))
ag <- readRDS(system.file("extdata/ag.rds",
  package = "SpatialFeatureExperiment"
))
annotGeometries(toy) <- list(hull = ag)

# Get names of annotation geometries
annotGeometryNames(sfe_small)

# Set names of annotation geometries
annotGeometryNames(toy) <- "foo"

# Get a specific annotation geometry by name
# sample_id is optional when there is only one sample present
nuclei <- annotGeometry(sfe_small, type = "nuclei", sample_id = "Vis5A")

# Get a specific annotation geometry by index
tb <- annotGeometry(sfe_small, type = 1L)

# Set a specific annotation geometry
```

```

annotGeometry(sfe_small, type = "nuclei2") <- nuclei

# Special convenience function for tissue boundaries
# Getter
tb <- tissueBoundary(sfe_small, sample_id = "Vis5A")
# Setter
tissueBoundary(sfe_small, sample_id = "Vis5A") <- tb

```

---

annotOp

*Binary operations for geometry of each cell/spot and annotation*


---

## Description

Just like [annotPred](#), but performs the operation rather than predicate. For example, this function would return the geometry of the intersections between each Visium spot and the tissue boundary for each sample, rather than whether each Visium spot intersects the tissue boundary. In case one cell/spot gets broken up into multiple geometries, the union of those geometries will be taken, so each cell/spot will only get one geometry.

## Usage

```

annotOp(
  sfe,
  colGeometryName = 1L,
  annotGeometryName = 1L,
  sample_id = "all",
  op = st_intersection
)

```

## Arguments

sfe	An SFE object.
colGeometryName	Name of column geometry for the predicate.
annotGeometryName	Name of annotation geometry for the predicate.
sample_id	Which sample(s) to operate on. Can be "all" to indicate all samples.
op	A binary operation function for the geometries. Defaults to <a href="#">st_intersection</a> .

## Value

A sf data frame with geometry column containing the geometries and corresponding column names of sfe as row names. There is no guarantee that the returned geometries are valid or preserve the geometry class (e.g. when the intersection of polygons result into a line of a point).

## See Also

[annotPred](#)

**Examples**

```

library(SFEData)
sfe <- McKellarMuscleData("small")
# Get the intersection of myofibers with each Visium spot
myofibers_on_spots <- annotOp(sfe, "spotPoly",
  annotGeometryName = "myofiber_simplified"
)

```

---

annotPred

*Binary predicates for geometry of each cell/spot and annotation*


---

**Description**

This function finds binary predicates for the geometry of each cell/spot (i.e. colGeometry) and an annotation geometry for each sample. For example, whether each Visium spot intersects with the tissue boundary in each sample.

**Usage**

```

annotPred(
  sfe,
  colGeometryName = 1L,
  annotGeometryName = 1L,
  sample_id = "all",
  pred = st_intersects,
  yx = FALSE
)

annotNPred(
  sfe,
  colGeometryName = 1L,
  annotGeometryName = 1L,
  sample_id = "all",
  pred = st_intersects
)

```

**Arguments**

sfe	An SFE object.
colGeometryName	Name of column geometry for the predicate.
annotGeometryName	Name of annotation geometry for the predicate.
sample_id	Which sample(s) to operate on. Can be "all" to indicate all samples.
pred	Predicate function to use, defaults to <a href="#">st_intersects</a> .
yx	Whether to do pred(y, x) instead of pred(x, y). For symmetric predicates, the results should be the same. When x has a large number of geometries and y has few, pred(y, x) is much faster than pred(x, y) for <a href="#">st_intersects</a> , <a href="#">st_disjoint</a> , and <a href="#">st_is_within_distance</a> .

**Value**

For `annotPred`, a logical vector of the same length as the number of columns in the sample(s) of interest, with barcodes (or corresponding column names of `sfe`) as names. For `annotNPred`, a numeric vector of the same length as the number of columns in the sample(s) of interest with barcodes as names, indicating the number of geometries in the `annotGeometry` of interest returns TRUE for the predicate for each each geometry in the `colGeometry` of interest.

**See Also**

`annotOp`

**Examples**

```
library(SFEData)
sfe <- McKellarMuscleData("small")
# Whether each spot is in tissue
in_tissue <- annotPred(sfe, "spotPoly", annotGeometryName = "tissueBoundary")
# How many nuclei are there in each Visium spot
n_nuclei <- annotNPred(sfe, "spotPoly", annotGeometryName = "nuclei")
```

---

annotSummary

*Summarize attributes of an annotGeometry for each cell/spot*

---

**Description**

In SFE objects, the annotation geometries don't have to correspond to the dimensions of the gene count matrix, so there generally is no one to one mapping between annotation geometries and cells/spots. However, it may be interesting to relate attributes of annotation geometries to cell/spots so the attributes can be related to gene expression. This function summarizes attributes of an `annotGeometry` for each cell/spot by a geometric predicate with a `colGeometry`.

**Usage**

```
annotSummary(
  sfe,
  colGeometryName = 1L,
  annotGeometryName = 1L,
  annotColNames = 1L,
  sample_id = "all",
  pred = st_intersects,
  summary_fun = mean
)
```

**Arguments**

`sfe` An SFE object.

`colGeometryName` Name of column geometry for the predicate.

`annotGeometryName` Name of annotation geometry for the predicate.

<code>annotColNames</code>	Character, column names of the <code>annotGeometry</code> of interest, to indicate the columns to summarize. Columns that are absent from the <code>annotGeometry</code> are removed. The column cannot be "geometry" or "barcode".
<code>sample_id</code>	Which sample(s) to operate on. Can be "all" to indicate all samples.
<code>pred</code>	Predicate function to use, defaults to <code>st_intersects</code> .
<code>summary_fun</code>	Function for the summary, defaults to mean.

**Value**

A data frame whose row names are the relevant column names of `sfe`, and each column of which is the summary of each column specified in `annotColName`.

**Examples**

```
library(SFData)
sfe <- McKellarMuscleData("small")
s <- annotSummary(sfe, "spotPoly", "myofiber_simplified",
  annotColNames = c("area", "convexity")
)
```

---

`bbox, SpatialFeatureExperiment-method`

*Find bounding box of SFE objects*

---

**Description**

Find bounding box of the union of all `colGeometries` and `annotGeometries` of each sample in the SFE object. This can be used to remove empty space so the tissue and geometries have one corner at the origin so all samples will be on comparable coordinates.

**Usage**

```
## S4 method for signature 'SpatialFeatureExperiment'
bbox(sfe, sample_id = "all", include_images = FALSE, include_row = TRUE)
```

**Arguments**

<code>sfe</code>	A <code>SpatialFeatureExperiment</code> object.
<code>sample_id</code>	Sample(s) whose bounding box(es) to find. The bounding box would be for the union of all <code>colGeometries</code> and <code>annotGeometries</code> associated with each sample.
<code>include_images</code>	Logical, whether the bounding boxes should include image extents. Defaults to FALSE because often the image has a lot of empty space surrounding the tissue.
<code>include_row</code>	Logical, whether the bounding boxes should include <code>rowGeometries</code> , defaults to TRUE.

**Value**

For one sample, then a named vector with names `xmin`, `ymin`, `xmax`, and `ymax` specifying the bounding box. For multiple samples, then a matrix whose columns are samples and whose rows delineate the bounding box.

**Examples**

```
library(SFEData)
sfe <- McKellarMuscleData("small")
bbox(sfe, sample_id = "Vis5A")
```

---

bbox_center	<i>Find center of bounding box</i>
-------------	------------------------------------

---

**Description**

Get x-y coordinates of the center of any bounding box

**Usage**

```
bbox_center(bbox)
```

**Arguments**

`bbox` A numeric vector of length 4 with names `xmin`, `xmax`, `ymin`, `ymax`, in any order.

**Value**

A numeric vector of length 2.

**Examples**

```
bbox <- c(xmin = 0, xmax = 100, ymin = 0, ymax = 80)
bbox_center(bbox)
```

---

BioFormatsImage	<i>On disk representation of BioFormats images in SFE object</i>
-----------------	--

---

**Description**

`'r lifecycle::badge("experimental")'` At present, the `BioFormatsImage` is designed for OME-TIFF from Xenium and has not been tested on other formats that can be read with `BioFormats`. The image is not loaded into memory, and when it is, the `BioFormatsImage` object is converted into `ExtImage` because the loaded image is of a class that inherits from `Image`. The `ExtImage` class is a thin wrapper inheriting from `VirtualSpatialImage` so it's compatible with `SpatialExperiment` from which `SFE` is derived. This class might drastically change as it matures, say to accommodate other formats supported by `BioFormats` and to store the transformation matrix rather than loading image into memory upon transform.

**Usage**

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

BioFormatsImage(
  path,
  ext = NULL,
  isFull = TRUE,
  origin = c(0, 0),
  transformation = list()
)
```

**Arguments**

object	A BioFormatsImage object.
path	Path to an OME-TIFF image file.
ext	Numeric vector with names "xmin", "xmax", "ymin", "ymax" in microns indicating the spatial extent covered by the image. If NULL, then the extent will be inferred from the metadata, from physical pixel size and the number of pixels.
isFull	Logical, if the extent specified in ext is the full extent. If ext = NULL so it will be inferred from metadata then isFull = TRUE will be set internally.
origin	Origin of the whole image in the x-y plane, defaults to c(0,0). This is shifted when the image is translated. This is not the same as xmin and xmax. For example, when the extent is only part of the whole image and the whole image itself can be spatially translated, the origin is needed to determine which part of the whole image this extent corresponds to.
transformation	Named list specifying affine transformation. The list can have names "name" and named parameter of the transformation, e.g. list(name = "mirror", direction = "vertical"), "rotate" and degrees = 90 (clockwise), and "scale" and factor = 2. The list can also have names "M" for a 2x2 linear transformation matrix in the xy plane and "v" for a translation vector of length 2 to specify general affine transformation.

**Details**

Spatial extent is inferred from OME-TIFF metadata if not specified. Physical pixel size from the metadata is used to make the extent in micron space. If physical pixel size is absent from metadata, then the extent will be in pixel space, which might mean that the image will not align with the geometries because often the geometry coordinates are in microns, so a warning is issued in this case.

Affine transformations can be specified in the transformation argument, either by name or by directly specifying the matrix. The transformations specified by name will always preserve the center of the image. When named transformations are chained, name and parameter will be converted to matrix and translation vector the second time a transformation is specified. If the subsequent transformation happens to restore the image to its original place, then transformation specifications will be removed.

**Value**

A BioFormatsImage object.

**See Also**

[isFull()], [origin()]

---

BioFormatsImage-getters

*Other BioFormatsImage getters*

---

**Description**

isFULL indicates if the extent is the full extent of the image. origin gets the x-y coordinates of the origin of the image, i.e. the smallest possible x-y coordinate values within the full image.

**Usage**

```
## S4 method for signature 'BioFormatsImage'
isFull(x)

## S4 method for signature 'BioFormatsImage'
origin(x)

## S4 method for signature 'BioFormatsImage'
transformation(x)
```

**Arguments**

x                    A [BioFormatsImage](#) object.

**Value**

For isFull: Logical scalar indicating whether the extent is the full extent. For origin: Numeric vector of length 2. For transformation, a list.

---

cbind, SpatialFeatureExperiment-method

*Concatenate SpatialFeatureExperiment objects*

---

**Description**

On top of the cbind method of SpatialExperiment, this method is needed to properly merge the spatialGraphs field in the different SFE objects. rowGeometries and annotGeometries also need to be combined properly.

**Usage**

```
## S4 method for signature 'SpatialFeatureExperiment'
cbind(..., deparse.level = 1)
```

**Arguments**

... SFE objects to cbind.  
 deparse.level See [?rbind](#).

**Value**

A combined SFE object.

**Examples**

```
library(SFEData)
sfe_small <- McKellarMuscleData(dataset = "small")
sfe_small2 <- McKellarMuscleData(dataset = "small2")
sfe2 <- cbind(sfe_small, sfe_small2)
```

---

changeSampleIDs	<i>Change sample IDs</i>
-----------------	--------------------------

---

**Description**

Change sample IDs in all fields of the SFE object where sample IDs are present, not just the colData.

**Usage**

```
changeSampleIDs(sfe, replacement)
```

**Arguments**

sfe A SpatialFeatureExperiment object.  
 replacement A named character vector whose names are the existing sample IDs to be changed and whose values are the corresponding replacements.

**Value**

An SFE object.

**Examples**

```
library(SFEData)
sfe <- McKellarMuscleData(dataset = "small")
sfe <- changeSampleIDs(sfe, c(Vis5A = "sample01"))
sampleIDs(sfe)
```

---

colFeatureData	<i>Get global spatial analysis results and metadata of colData, rowData, and geometries</i>
----------------	---

---

### Description

Results of spatial analyses on columns in colData, rowData, and geometries are stored in their metadata. The colFeatureData function allows the users to more directly access these results.

### Usage

```
colFeatureData(sfe)

rowFeatureData(sfe)

geometryFeatureData(sfe, type, MARGIN = 2L)

reducedDimFeatureData(sfe, dimred)
```

### Arguments

sfe	An SFE object.
type	Which geometry, can be name (character) or index (integer)
MARGIN	Integer, 1 means rowGeometry, 2 means colGeometry, and 3 means annotGeometry. Defaults to 2, colGeometry.
dimred	Name of a dimension reduction, can be seen in <a href="#">reducedDimNames</a> .

### Value

A DataFrame.

### See Also

`getParams`

### Examples

```
library(SpatialFeatureExperiment)
library(SingleCellExperiment)
library(SFEData)
library(Voyager)
sfe <- McKellarMuscleData("small")
colGraph(sfe, "visium") <- findVisiumGraph(sfe)
# Moran's I for colData
sfe <- colDataMoransI(sfe, "nCounts")
colFeatureData(sfe)
```

---

colGeometries                      *Column geometry getters and setters*

---

### Description

colGeometries are geometries that correspond to columns of the gene count matrix, such as Visium spots or cells. Same as `dimGeometry(x, MARGIN = 2L, ...)`, with convenience wrappers for getters and setters of special geometries:

**spotPoly** Polygons of spots from technologies such as Visium, ST, and slide-seq, which do not correspond to cells. Centroids of the polygons are stored in `spatialCoords` of the underlying `SpatialExperiment` object.

**ROI Poly** Polygons of regions of interest (ROIs) from technologies such as laser capture microdissection (LCM) and GeoMX DSP. These should correspond to columns of the gene count matrix.

**cellSeg** Cell segmentation polygons. If the columns of the gene count matrix are single cells, then this is stored in `colGeometries`. Otherwise, this is stored in [annotGeometries](#).

**nucSeg** Similar to `cellSeg`, but for nuclei rather than whole cell.

### Usage

```
colGeometry(x, type = 1L, sample_id = 1L, withDimnames = TRUE)
```

```
colGeometry(
  x,
  type = 1L,
  sample_id = 1L,
  withDimnames = TRUE,
  translate = TRUE
) <- value
```

```
colGeometries(x, withDimnames = TRUE)
```

```
colGeometries(x, withDimnames = TRUE, translate = TRUE) <- value
```

```
colGeometryNames(x)
```

```
colGeometryNames(x) <- value
```

```
spotPoly(x, sample_id = 1L, withDimnames = TRUE)
```

```
spotPoly(x, sample_id = 1L, withDimnames = TRUE, translate = TRUE) <- value
```

```
centroids(x, sample_id = 1L, withDimnames = TRUE)
```

```
centroids(x, sample_id = 1L, withDimnames = TRUE, translate = TRUE) <- value
```

```
ROI Poly(x, sample_id = 1L, withDimnames = TRUE)
```

```
ROI Poly(x, sample_id = 1L, withDimnames = TRUE, translate = TRUE) <- value
```

```

cellSeg(x, sample_id = 1L, withDimnames = TRUE)

cellSeg(x, sample_id = 1L, withDimnames = TRUE, translate = TRUE) <- value

nucSeg(x, sample_id = 1L, withDimnames = TRUE)

nucSeg(x, sample_id = 1L, withDimnames = TRUE, translate = TRUE) <- value

```

### Arguments

x	A SpatialFeatureExperiment object.
type	An integer specifying the index or string specifying the name of the *Geometry to query or replace. If missing, then the first item in the *Geometries will be returned or replaced.
sample_id	Sample ID to get or set geometries.
withDimnames	Logical. If TRUE, then the dimnames (colnames or rownames) of the gene count matrix should correspond to row names of the sf data frames of interest.
translate	Logical. Only used if <a href="#">removeEmptySpace</a> has been run of the SFE object. If that's the case, this argument indicates whether the new value to be assigned to the geometry is in the coordinates prior to removal of empty space so it should be translated to match the new coordinates after removing empty space. Default to TRUE.
value	Value to set. For dimGeometry, must be a sf data frame with the same number of rows as size in the dimension of interest, or an ordinary data frame that can be converted to such a sf data frame (see <a href="#">df2sf</a> ). For dimGeometries, must be a list of such sf or ordinary data frames.

### See Also

[dimGeometries()], [rowGeometries()]

### Examples

```

library(SFEData)
sfe <- McKellarMuscleData(dataset = "small")
cgs <- colGeometries(sfe)
spots <- spotPoly(sfe)

```

---

containsOutOfMemoryData, SpatialFeatureExperiment-method

*Whether an SFE object contains out of memory data*

---

### Description

Out of memory data, such as DelayedArray, some SpatRasterImage objects, and BioFormatsImage, will break if saved as RDS. This method of [containsOutOfMemoryData](#) checks if an SFE object has out of memory data, specifically the images. Having out of memory data will result into an error when saveRDS is called; we recommend using the `alabaster.sfe` package instead.

**Usage**

```
## S4 method for signature 'SpatialFeatureExperiment'
containsOutOfMemoryData(object)
```

**Arguments**

object            An SFE object

**Value**

TRUE or FALSE

**Examples**

```
outdir <- system.file("extdata", package = "SpatialFeatureExperiment")
samples <- file.path(outdir, paste0("sample0", 1:2))
sfe <- read10xVisiumSFE(samples, type = "sparse", data = "filtered")
containsOutOfMemoryData(sfe)
```

---

crop

*Crop an SFE object with a geometry*

---

**Description**

Returns an SFE object whose specified `colGeometry` returns TRUE with a geometric predicate function (usually intersects) with another geometry of interest. This can be used to subset an SFE object with a tissue boundary or histological region polygon, or crop away empty spaces. After cropping, not only will the cells/spots be subsetted, but also all geometries will be cropped.

**Usage**

```
crop(
  x,
  y = NULL,
  colGeometryName = 1L,
  sample_id = "all",
  op = st_intersection,
  keep_whole = "none",
  cover = FALSE
)
```

**Arguments**

x                    An SFE object.

y                    An object of class `sf`, `sfg`, `sfc` with which to crop the SFE object, or a bounding box with the format of the output of `bbox`, [SpatialFeatureExperiment-method](#).

colGeometryName    Column geometry to used to indicate which cells/spots to keep.

sample_id	Samples to crop. Optional when only one sample is present. Can be multiple samples, or "all", which means all samples. For multiple samples, sf data frame y may have column sample_id indicating which geometry subsets which sample or matrix y may indicate sample specific bounding boxes in its column names. Only samples included in the indicated sample IDs are subsetted. If sample is not indicated in y, then the same geometry or bounding box is used to subset all samples specified in the sample_id argument.
op	A geometric operation function to crop the geometries in the SFE object. Only <code>st_intersection</code> and <code>st_difference</code> are allowed. If "intersection", then only things inside y is kept after cropping. If "difference", then only things outside y is kept.
keep_whole	Character vector, can be one or more of "col" and "annot" to keep whole items from <code>colGeometries</code> or <code>annotGeometries</code> , keeping geometries that partially intersect with y whole. This can greatly speed up code while not breaking geometries into multiple pieces. Can also be "none" so all geometries are actually cropped.
cover	Logical, whether the geometries in x must be entirely covered by y if <code>op = st_intersection</code> or whether x must be entirely outside y if <code>op = st_difference</code> . Only relevant when <code>keep_whole != "none"</code> .

### Details

3D geometries are allowed, but geometric operations can only be performed in x and y but not z.

### Value

An SFE object. There is no guarantee that the geometries after cropping are still all valid or preserve the original geometry class.

### Examples

```
library(SFData)
sfe <- McKellarMuscleData("small")
# Subset sfe to only keep spots on tissue
sfe_on_tissue <- crop(sfe, tissueBoundary(sfe),
  colGeometryName = "spotPoly",
  sample_id = "Vis5A"
)
```

---

cropImg

*Crop images*

---

### Description

Crop images of class `*Image` in this package with a bounding box.

**Usage**

```
## S4 method for signature 'SpatRasterImage'
cropImg(x, bbox, filename = "")

## S4 method for signature 'BioFormatsImage'
cropImg(x, bbox)

## S4 method for signature 'ExtImage'
cropImg(x, bbox)
```

**Arguments**

x	An object of class *Image as implemented in this package.
bbox	Numeric vector with names "xmin", "xmax", "ymin", "ymax", in any order, to specify the bounding box.
filename	Output file name for transformed SpatRaster.

**Value**

Image of the same class as input but cropped. For BioFormatsImage, the image is not loaded into memory; only the extent is changed.

**See Also**

Other image methods: [SFE-image](#), [affineImg\(\)](#), [dim](#), [BioFormatsImage-method](#), [dim](#), [ExtImage-method](#), [ext\(\)](#), [imgRaster\(\)](#), [imgSource\(\)](#), [mirrorImg\(\)](#), [rotateImg\(\)](#), [scaleImg\(\)](#), [translateImg\(\)](#), [transposeImg\(\)](#)

---

df2sf

*From ordinary data frame to sf to construct SFE object*


---

**Description**

While the SpatialFeatureExperiment constructor and \*Geometry replacement methods can convert properly formatted ordinary data frames into sf objects which are used to store the geometries internally, the user might want to do the conversion, check if the geometry is valid, and inspect and fix any invalid geometries.

**Usage**

```
df2sf(
  df,
  spatialCoordsNames = c("x", "y"),
  spotDiameter = NA,
  geometryType = c("POINT", "LINESTRING", "POLYGON", "MULTIPOINT", "MULTILINESTRING",
    "MULTIPOLYGON"),
  group_col = "group",
  id_col = "ID",
  subid_col = "subID",
  check = TRUE,
  ...
)
```

**Arguments**

df	An ordinary data frame, i.e. not sf. Or a matrix that can be converted to a data frame.
spatialCoordsNames	Column names in df that specify spatial coordinates.
spotDiameter	Spot diameter for technologies with arrays of spots of fixed diameter per slide, such as Visium, ST, DBiT-seq, and slide-seq. The diameter must be in the same unit as the coordinates in the *Geometry arguments. Ignored for geometries that are not POINT or MULTIPOINT.
geometryType	Type of geometry to convert the ordinary data frame to. If the geometry in df is de facto points, then this argument will be ignored and the returned sf will have geometry type POINT.
group_col	Column to indicate which coordinates for which MULTI geometry, such as to identify which MULTIPOLYGON or MULTIPOINT.
id_col	Column to indicate coordinates for which geometry, within a MULTI geometry if applicable, such as to identify which POLYGON or which polygon within a MULTIPOLYGON.
subid_col	Column to indicate coordinates for holes in polygons.
check	Logical, whether to check the input data frame for issues related to constructing the geometry of interest such as number of vertices per geometry. If FALSE, it will save a bit of time, which is useful when the input is already known to be good.
...	Other arguments passed to 'sf::st_buffer', mainly to make polygon shapes, eg Visium spot 'endCapStyle = "ROUND"' and VisiumHD bin 'endCapStyle = "SQUARE"'

**Value**

An sf object.

**Examples**

```
# Points, use spotDiameter to convert to circle polygons
# This is done to Visium spots
pts_df <- readRDS(system.file("extdata/pts_df.rds",
  package = "SpatialFeatureExperiment"
))
sf_use <- df2sf(pts_df, geometryType = "POINT", spotDiameter = 0.1)
# Linestring
ls_df <- readRDS(system.file("extdata/ls_df.rds",
  package = "SpatialFeatureExperiment"
))
sf_use <- df2sf(ls_df, geometryType = "LINESTRING")
# Polygon
pol_df <- readRDS(system.file("extdata/pol_df.rds",
  package = "SpatialFeatureExperiment"
))
sf_use <- df2sf(pol_df,
  geometryType = "POLYGON",
  spatialCoordsNames = c("V1", "V2")
)
# Multipolygon
```

```

mpol_df <- readRDS(system.file("extdata/mpol_df.rds",
  package = "SpatialFeatureExperiment"
))
sf_use <- df2sf(mpol_df,
  geometryType = "MULTIPOLYGON",
  spatialCoordsNames = c("V1", "V2")
)
# Multiple sample_ids present
multipts_df <- readRDS(system.file("extdata/multipts_df.rds",
  package = "SpatialFeatureExperiment"
))
sf_use <- df2sf(multipts_df, geometryType = "MULTIPOINT")

```

---

dim,BioFormatsImage-method

*Find dimension of BioFormatsImage*

---

## Description

This is different from other classes. The metadata is read where the dimensions in pixels can be found. The image itself is not read into memory here.

## Usage

```

## S4 method for signature 'BioFormatsImage'
dim(x)

```

## Arguments

x                    A [BioFormatsImage](#) object.

## Value

An integer vector of length 5 showing the number of rows and columns in the full resolution image. The 5 dimensions are in the order of XYCZT: x, y, channel, z, and time. This is not changed by transformations. Use [ext](#) to see the extent after transformation.

## See Also

Other image methods: [SFE-image](#), [affineImg\(\)](#), [cropImg\(\)](#), [dim,ExtImage-method](#), [ext\(\)](#), [imgRaster\(\)](#), [imgSource\(\)](#), [mirrorImg\(\)](#), [rotateImg\(\)](#), [scaleImg\(\)](#), [translateImg\(\)](#), [transposeImg\(\)](#)

---

dim,ExtImage-method     *Find dimensions of ExtImage*

---

### Description

This method exists to make the output of dim() for ExtImage consistent with that of Image which ExtImage inherits from, overriding the VirtualSpatialImage method.

### Usage

```
## S4 method for signature 'ExtImage'
dim(x)
```

### Arguments

x                     A [ExtImage](#) object.

### Value

An integer vector. As in EImage, the first element indicates number of pixels in the x direction, or number of columns in the image, and the second element indicates the number of pixels in the y direction. This is unlike array indexing.

### See Also

Other image methods: [SFE-image](#), [affineImg\(\)](#), [cropImg\(\)](#), [dim,BioFormatsImage-method](#), [ext\(\)](#), [imgRaster\(\)](#), [imgSource\(\)](#), [mirrorImg\(\)](#), [rotateImg\(\)](#), [scaleImg\(\)](#), [translateImg\(\)](#), [transposeImg\(\)](#)

---

dimGeometries             *Dimension geometry methods*

---

### Description

"Dimension geometry" refers to Simple Feature (sf) geometries associated with rows (features, genes) or columns (cells or spots) of the gene count matrix in the SpatialFeatureExperiment object. For each dimension, the number of rows in the sf data frame specifying the geometries must match the size of the dimension of interest. For example, there must be the same number of rows in the sf data frame describing cells as there are cells in the gene count matrix. This page documents getters and setters for the dimension geometries. The getters and setters are implemented in a way similar to those of reducedDims in SingleCellExperiment.

### Usage

```
## S4 method for signature 'SpatialFeatureExperiment'
dimGeometries(x, MARGIN = 2, withDimnames = TRUE)

## S4 replacement method for signature 'SpatialFeatureExperiment'
dimGeometries(x, MARGIN, withDimnames = TRUE, translate = TRUE, ...) <- value
```

```

## S4 method for signature 'SpatialFeatureExperiment'
dimGeometryNames(x, MARGIN)

## S4 replacement method for signature 'SpatialFeatureExperiment,numeric,character'
dimGeometryNames(x, MARGIN) <- value

## S4 method for signature 'SpatialFeatureExperiment'
dimGeometry(x, type = 1L, MARGIN, sample_id = 1L, withDimnames = TRUE)

## S4 replacement method for signature 'SpatialFeatureExperiment'
dimGeometry(
  x,
  type = 1L,
  MARGIN,
  sample_id = 1L,
  withDimnames = TRUE,
  translate = TRUE,
  ...
) <- value

```

### Arguments

x	A <code>SpatialFeatureExperiment</code> object.
MARGIN	As in <code>apply</code> . 1 stands for rows and 2 stands for columns.
withDimnames	Logical. If TRUE, then the dimnames (colnames or rownames) of the gene count matrix should correspond to row names of the sf data frames of interest.
translate	Logical. Only used if <code>removeEmptySpace</code> has been run of the SFE object. If that's the case, this argument indicates whether the new value to be assigned to the geometry is in the coordinates prior to removal of empty space so it should be translated to match the new coordinates after removing empty space. Default to TRUE.
...	<code>spatialCoordsNames</code> , <code>spotDiameter</code> , <code>geometryType</code> passed to <code>df2sf</code> . Defaults are the same as in <code>df2sf</code> . For <code>dimGeometries&lt;-</code> only: <code>geometryType</code> can be a character vector of the geometry type of each data frame in the list of the same length as the list if the data frames specify different types of geometries.
value	Value to set. For <code>dimGeometry</code> , must be a sf data frame with the same number of rows as size in the dimension of interest, or an ordinary data frame that can be converted to such a sf data frame (see <code>df2sf</code> ). For <code>dimGeometries</code> , must be a list of such sf or ordinary data frames.
type	An integer specifying the index or string specifying the name of the <code>*Geometry</code> to query or replace. If missing, then the first item in the <code>*Geometries</code> will be returned or replaced.
sample_id	Sample ID to get or set geometries.

### Value

Getters for multiple geometries return a named list. Getters for names return a character vector of the names. Getters for single geometries return an sf data frame. Setters return an SFE object.

**See Also**

[colGeometries()], [rowGeometries()]

**Examples**

```

library(SFEData)
sfe <- McKellarMuscleData(dataset = "small")

# Get all column geometries as a named list
# Use MARGIN = 1 or rowGeometry/ies for rowGeometries
cgs <- dimGeometries(sfe, MARGIN = 2)
# Or equivalently
cgs <- colGeometries(sfe)

# Set all column geometries with a named list
dimGeometries(sfe, MARGIN = 2) <- cgs
# Or equivalently
colGeometries(sfe) <- cgs

# Get names of column geometries
cgns <- dimGeometryNames(sfe, MARGIN = 2)
cgns <- colGeometryNames(sfe)

# Set column geometry names
dimGeometryNames(sfe, MARGIN = 2) <- cgns
colGeometryNames(sfe) <- cgns

# Get a specific column geometry by name
spots <- dimGeometry(sfe, "spotPoly", MARGIN = 2)
spots <- colGeometry(sfe, "spotPoly")
# Or equivalently, the wrapper specifically for Visium spot polygons,
# for the name "spotPoly"
spots <- spotPoly(sfe)
# Other colGeometry wrappers for specific names:
# ROIpoly (for LCM and GeoMX DSP), cellSeg and nucSeg (for MERFISH; would
# query annotGeometries for Visium)
# rowGeometry wrappers for specific names: txSpots (MERFISH transcript spots)
# By index
spots <- colGeometry(sfe, 1L)

# Multiple samples, only get geometries for one sample
sfe2 <- McKellarMuscleData("small2")
sfe_combined <- cbind(sfe, sfe2)
spots1 <- colGeometry(sfe, "spotPoly", sample_id = "Vis5A")
spots2 <- spotPoly(sfe_combined, sample_id = "sample02")
# Get geometries for multiple samples
spots3 <- spotPoly(sfe_combined, sample_id = c("Vis5A", "sample02"))
# All samples
spots3 <- spotPoly(sfe_combined, sample_id = "all")

# Set specific column geometry by name
colGeometry(sfe, "foobar") <- spots
# Or use wrapper
spotPoly(sfe) <- spots
# Specify sample_id
colGeometry(sfe_combined, "foobar", sample_id = "Vis5A") <- spots1

```

```
# Only entries for the specified sample are set.
foobar <- colGeometry(sfe_combined, "foobar", sample_id = "sample02")
```

---

 ext

*Get and set extent of image objects*


---

## Description

Unlike in `SpatialExperiment`, images in SFE have extents which are used to align them to the geometries and in geometric operations on SFE objects. These functions get or set the extent for S4 image classes inheriting from `VirtualSpatialImage` implemented in the SFE package.

## Usage

```
## S4 method for signature 'BioFormatsImage'
ext(x)

## S4 method for signature 'ExtImage'
ext(x)

## S4 method for signature 'SpatRasterImage'
ext(x)

## S4 replacement method for signature 'BioFormatsImage,numeric'
ext(x) <- value

## S4 replacement method for signature 'ExtImage,numeric'
ext(x) <- value

## S4 replacement method for signature 'SpatRasterImage,numeric'
ext(x) <- value
```

## Arguments

x	A <code>*Image</code> object.
value	A numeric vector with names "xmin", "xmax", "ymin", "ymax" specifying the extent to use.

## Value

Getters return a numeric vector specifying the extent. Setters return a `*Image` object of the same class as the input.

## Note

For `SpatRasterImage`, the image may be may not be loaded into memory. You can check if the image is loaded into memory with `terra::inMemory(x)`, and check the original file path with [imgSource](#). If the image is not loaded into memory, then the original file must be present at the path indicated by [imgSource](#) in order for any code using the image to work, which includes this function `ext`.

For `BioFormatsImage`, internally only the pre-transform extent is stored. The `ext` getter will apply the transformation on the fly. The setter sets the pre-transformation extent.

**See Also**

Other image methods: [SFE-image](#), [affineImg\(\)](#), [cropImg\(\)](#), [dim,BioFormatsImage-method](#), [dim,ExtImage-method](#), [imgRaster\(\)](#), [imgSource\(\)](#), [mirrorImg\(\)](#), [rotateImg\(\)](#), [scaleImg\(\)](#), [translateImg\(\)](#), [transposeImg\(\)](#)

---

 ExtImage

*Use the EImage Image class in SFE objects*


---

**Description**

This is a thin wrapper around the [Image](#) class in the EImage package so it inherits from `VirtualSpatialImage` to be compatible with `SpatialExperiment` from which SFE inherits. An `ext` field is added to specify the spatial extent of the image in microns to facilitate geometric operations on the SFE object (including the images) and plotting with `Voyager`.

**Usage**

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

ExtImage(img, ext = NULL)
```

**Arguments**

<code>object</code>	An ExtImage object.
<code>img</code>	An Image object or anything that inherits from Image such as <code>AnnotatedImage</code> in <code>RBioFormats</code> .
<code>ext</code>	Numeric vector with names "xmin", "xmax", "ymin", "ymax" in microns indicating the spatial extent covered by the image. If NULL, then the extent will be inferred from the metadata, from physical pixel size and the number of pixels.

**Value**

An ExtImage object.

---

 findDebrisCells

*Identify cells in small bits outside the main piece of tissue*


---

**Description**

This is used for quality control (QC), as the small bits are likely to be low quality technical artifacts and are not informative to spatial analyses. Please confirm the quality of those cells by checking the histology image with ImageJ or QuPath.

**Usage**

```

## S4 method for signature 'matrix'
findDebrisCells(
  x,
  max_cells = 5,
  distance_cutoff = 50,
  BNPARAM = NULL,
  BPPARAM = SerialParam()
)

## S4 method for signature 'sfc'
findDebrisCells(
  x,
  max_cells = 5,
  distance_cutoff = 50,
  BNPARAM = NULL,
  BPPARAM = SerialParam()
)

## S4 method for signature 'sf'
findDebrisCells(
  x,
  max_cells = 5,
  distance_cutoff = 50,
  BNPARAM = NULL,
  BPPARAM = SerialParam()
)

## S4 method for signature 'SpatialExperiment'
findDebrisCells(
  x,
  max_cells = 5,
  distance_cutoff = 50,
  BNPARAM = NULL,
  BPPARAM = SerialParam()
)

```

**Arguments**

x	Object with spatial coordinates of cells. Can be a <code>SpatialExperiment</code> or <code>SpatialFeatureExperiment</code> object, a matrix with 2 columns for x and y coordinates of cells, or a <code>sf</code> or <code>sfc</code> object with cell geometries.
max_cells	Maximum number of cells for a clump of cells to be considered debris.
distance_cutoff	Minimum distance of cell to the tissue for it to be considered debris, in the same unit as in x.
BNPARAM	A <code>BiocNeighborParam</code> object specifying the algorithm to find k nearest neighbors and distance based neighbors with <code>nn_method = "bioc"</code> . For distance based neighbors, only <code>KmknParam</code> and <code>VptreeParam</code> are applicable.
BPPARAM	A <code>BiocParallelParam</code> object for multithreading. Only used for k nearest neighbor and distance based neighbor with <code>nn_method = "bioc"</code> .

## Details

How this function works: A distance-based spatial neighborhood graph is computed, with `distance_cutoff` as the distance cutoff. Then disjoint connected subgraphs are found. Cells in subgraphs with `max_cells` or fewer cells are considered debris.

## Value

Depends on the method:

**Spatial(Feature)Experiment** The same object with a logical column "is\_debris" added to `colData`.

**Matrix and sf(c)** A logical vector indicating whether each cell is debris.

---

findSpatialNeighbors *Find spatial neighborhood graph*

---

## Description

This function wraps all spatial neighborhood graphs implemented in the package `spdep` for the `SpatialFeatureExperiment` (SFE) class, to find spatial neighborhood graphs for the entities represented by columns or rows of the gene count matrix in the SFE object or spatial entities in the `annotGeometries` field of the SFE object. Results are stored as `listw` objects in the `spatialGraphs` field of the SFE object, as `listw` is used in many methods that facilitate the spatial neighborhood graph in the `spdep`, `spatialreg`, and `adespatial`. The edge weights of the graph in the `listw` object are by default style `W` (see [nb2listw](#)) and the unweighted neighbor list is in the `neighbours` field of the `listw` object.

## Usage

```
## S4 method for signature 'SpatialFeatureExperiment'
findSpatialNeighbors(
  x,
  sample_id = "all",
  type = "spatialCoords",
  MARGIN = 2,
  method = c("tri2nb", "knearneigh", "dnearneigh", "gabrielneigh", "relativeneigh",
    "soi.graph", "poly2nb"),
  dist_type = c("none", "idw", "exp", "dpd"),
  glist = NULL,
  style = c("raw", "W", "B", "C", "U", "minmax", "S"),
  nn_method = c("bioc", "spdep"),
  alpha = 1,
  dmax = NULL,
  BPPARAM = SerialParam(),
  BNPARAM = KmknnParam(),
  zero.policy = TRUE,
  ...
)
```

**Arguments**

x	A <a href="#">SpatialFeatureExperiment</a> object.
sample_id	Which sample(s) in the SFE object to use for the graph. Can also be "all", which means this function will compute the graph for all samples independently.
type	Name of the geometry associated with the MARGIN of interest for which to compute the graph.
MARGIN	Just like in <a href="#">apply</a> , where 1 stands for row, 2 stands for column. Here, in addition, 3 stands for annotation, to query the <a href="#">annotGeometries</a> , such as nuclei segmentation in a Visium data
method	Name of function in the package <a href="#">spdep</a> to use to find the spatial neighborhood graph.
dist_type	Type of distance-based weight. "none" means not using distance-based weights; the edge weights of the spatial neighborhood graph will be entirely determined by the <code>style</code> argument. "idw" means inverse distance weighting. "exp" means exponential decay. "dpd" means double-power distance weights. See <a href="#">nb2listwdist</a> for details.
glist	list of general weights corresponding to neighbours
style	style can take values "W", "B", "C", "U", "minmax" and "S"
nn_method	Method to find k nearest neighbors and distance based neighbors. Can be either "bioc" or "spdep". For "bioc", methods from <a href="#">BiocNeighbors</a> are used. For "spdep", methods from the <a href="#">spdep</a> package are used. The "bioc" option is more scalable to larger datasets and supports multithreading.
alpha	Only relevant when <code>dist_type = "dpd"</code> .
dmax	Only relevant when <code>dist_type = "dpd"</code> .
BPPARAM	A <a href="#">BiocParallelParam</a> object for multithreading. Only used for k nearest neighbor and distance based neighbor with <code>nn_method = "bioc"</code> .
BNPARAM	A <a href="#">BiocNeighborParam</a> object specifying the algorithm to find k nearest neighbors and distance based neighbors with <code>nn_method = "bioc"</code> . For distance based neighbors, only <a href="#">KmknnParam</a> and <a href="#">VptreeParam</a> are applicable.
zero.policy	default NULL, use global option value; if FALSE stop with error for any empty neighbour sets, if TRUE permit the weights list to be formed with zero-length weights vectors
...	Extra arguments passed to the <a href="#">spdep</a> function stated in the <code>method</code> argument, such as <code>k</code> , <code>use_kd_tree</code> , <code>d1</code> , <code>d2</code> , <code>nnmult</code> , <code>sym</code> , and <code>quadsegs</code> . Note that any arguments about using longitude and latitude, which are irrelevant, are ignored.

**Value**

For one sample, then a `listw` object representing the graph, with an attribute "method" recording the function used to build the graph, its arguments, and information about the geometry for which the graph was built. The attribute is used to reconstruct the graphs when the SFE object is subsetted since some nodes in the graph will no longer be present. If `sample_id = "all"` or has length > 1, then a named list of `listw` objects, whose names are the `sample_ids`. To add the list for multiple samples to a SFE object, specify the name argument in the [spatialGraphs](#) replacement method, so graph of the same name will be added to the SFE object for each sample.

**Note**

style = "raw" is only applicable when dist\_type is not "none". If dist\_type = "none" and style = "raw", then style will default to "W". Using distance based weights does not supplant finding a spatial neighborhood graph. The spatial neighborhood graph is first found and then its edges weighted based on distance in this function.

**Examples**

```
library(SFEData)
sfe <- McKellarMuscleData(dataset = "small")
# sample_id is optional when only one sample is present
g <- findSpatialNeighbors(sfe, sample_id = "Vis5A")
attr(g, "method")
# Returns named list for multiple samples
sfe2 <- McKellarMuscleData(dataset = "small2")
sfe_combined <- cbind(sfe, sfe2)
gs <- findSpatialNeighbors(sfe, sample_id = "all")
```

---

findVisiumGraph

*Find spatial neighborhood graphs for Visium spots*


---

**Description**

Visium spots are arranged in a hexagonal grid. This function uses the known locations of the Visium barcodes to construct a neighborhood graph, so adjacent spots are connected by edges. Since the known rows and columns of the spots are used, the unit the spot centroid coordinates are in does not matter.

**Usage**

```
findVisiumGraph(x, sample_id = "all", style = "W", zero.policy = NULL)
```

**Arguments**

x	A SpatialFeatureExperiment object with Visium data. Column names of the gene count matrix must be Visium barcodes, which may have a numeric suffix to distinguish between samples (e.g. "AAACAACGAATAGTTC-1").
sample_id	Which sample(s) in the SFE object to use for the graph. Can also be "all", which means this function will compute the graph for all samples independently.
style	style can take values "W", "B", "C", "U", "minmax" and "S"
zero.policy	default NULL, use global option value; if FALSE stop with error for any empty neighbour sets, if TRUE permit the weights list to be formed with zero-length weights vectors

**Value**

For one sample, then a listw object representing the graph, with an attribute "method" recording the function used to build the graph, its arguments, and information about the geometry for which the graph was built. The attribute is used to reconstruct the graphs when the SFE object is subsetted since some nodes in the graph will no longer be present. If sample\_id = "all" or has length > 1, then a named list of listw objects, whose names are the sample\_ids. To add the list for multiple samples to a SFE object, specify the name argument in the [spatialGraphs](#) replacement method, so graph of the same name will be added to the SFE object for each sample.

**Examples**

```

library(SFEData)
sfe <- McKellarMuscleData(dataset = "small")
g <- findVisiumGraph(sfe)
# For multiple samples, returns named list
sfe2 <- McKellarMuscleData(dataset = "small2")
sfe_combined <- cbind(sfe, sfe2)
gs <- findVisiumGraph(sfe, sample_id = "all")

```

---

findVisiumHDGraph	<i>Find Visium HD spatial neighborhood graph</i>
-------------------	--

---

**Description**

Visium HD spots are arranged in a square grid. This function finds either a rook or a queen spatial neighborhood graph for the spots. colData of the SFE object must have columns array\_row and array\_col.

**Usage**

```
findVisiumHDGraph(x, style = "W", queen = FALSE, zero.policy = TRUE)
```

**Arguments**

x	An SFE object with Visium HD data with one sample with the required information in its colData.
style	style can take values "W", "B", "C", "U", "minmax" and "S"
queen	Logical. Default is FALSE, using rook neighbors.
zero.policy	default NULL, use global option value; if FALSE stop with error for any empty neighbour sets, if TRUE permit the weights list to be formed with zero-length weights vectors

**Value**

A listw object for the graph.

---

formatTxSpots	<i>Read and process transcript spots geometry for SFE</i>
---------------	---

---

**Description**

The function 'formatTxSpots' reads the transcript spot coordinates of smFISH-based data and formats the data. The data is not added to an SFE object. If the file specified in 'file\_out' already exists, then this file will be read instead of the original file in the 'file' argument, so the processing is not run multiple times. The function 'addTxSpots' adds the data read and processed in 'formatTxSpots' to the SFE object, and reads all transcript spot data. To only read a subset of transcript spot data, first use 'formatTxSpots' to write the re-formatted data to disk. Then read the specific subset and add them separately to the SFE object with the setter functions.

**Usage**

```

formatTxSpots(
  file,
  dest = c("rowGeometry", "colGeometry"),
  spatialCoordsNames = c("global_x", "global_y", "global_z"),
  gene_col = "gene",
  cell_col = "cell_id",
  z = "all",
  phred_col = "qv",
  min_phred = 20,
  split_col = NULL,
  not_in_cell_id = c("-1", "UNASSIGNED"),
  z_option = c("3d", "split"),
  flip = FALSE,
  file_out = NULL,
  BPPARAM = SerialParam(),
  return = TRUE,
  save_memory = FALSE,
  partition = FALSE
)

addTxSpots(
  sfe,
  file,
  sample_id = 1L,
  spatialCoordsNames = c("global_x", "global_y", "global_z"),
  gene_col = "gene",
  z = "all",
  phred_col = "qv",
  min_phred = 20,
  split_col = NULL,
  z_option = c("3d", "split"),
  flip = FALSE,
  file_out = NULL,
  BPPARAM = SerialParam()
)

```

**Arguments**

file	File with the transcript spot coordinates. Should be one row per spot when read into R and should have columns for coordinates on each axis, gene the transcript is assigned to, and optionally cell the transcript is assigned to. Must be csv, tsv, or parquet.
dest	Where in the SFE object to store the spot geometries. This affects how the data is processed. Options: <ul style="list-style-type: none"> <li><b>rowGeometry</b> All spots for each gene will be a ‘MULTIPOINT’ geometry, regardless of whether they are in cells or which cells they are assigned to.</li> <li><b>colGeometry</b> The spots for each gene assigned to a cell of interest will be a ‘MULTIPOINT’ geometry; since the gene count matrix is sparse, the geometries are NOT returned to memory.</li> </ul>

spatialCoordsNames	Column names for the x, y, and optionally z coordinates of the spots. The defaults are for Vizgen.
gene_col	Column name for genes.
cell_col	Column name for cell IDs, ignored if 'dest = "rowGeometry"'. Can have length > 1 when multiple columns are needed to uniquely identify cells, in which case the contents of the columns will be concatenated, such as in CosMX data where cell ID is only unique within the same FOV. Default "cell_id" is for Vizgen MERFISH. Should be 'c("cell_ID", "fov")' for CosMX.
z	Index of z plane to read. Can be "all" to read all z-planes into MULTIPOINT geometries with XYZ coordinates. If z values are not integer, then spots with all z values will be read.
phred_col	Column name for Phred scores of the spots.
min_phred	Minimum Phred score to keep spot. By default 20, the conventional threshold indicating "acceptable", meaning that there's 1 chance that the spot was decoded in error.
split_col	Categorical column to split the geometries, such as cell compartment the spots are assigned to as in the "CellComp" column in CosMX output.
not_in_cell_id	Value of cell ID indicating that the spot is not assigned to any cell, such as "-1" in Vizgen MERFISH and "0" in CosMX. When there're multiple columns for 'cell_col', the first column is used to identify spots that are not in cells.
z_option	What to do with z coordinates. "3d" is to construct 3D geometries. "split" is to create a separate 2D geometry for each z-plane so geometric operations are fully supported but some data wrangling is required to perform 3D analyses. When the z coordinates are not integers, 3D geometries will always be constructed since there are no z-planes to speak of. This argument does not apply when 'spatialCoordsNames' has length 2.
flip	Logical, whether to flip the geometry to match image. Here the y coordinates are simply set to -y, so the original bounding box is not preserved. This is consistent with readVizgen and readXenium.
file_out	Name of file to save the geometry or raster to disk. Especially when the geometries are so large that it's unwieldy to load everything into memory. If this file (or directory for multiple files) already exists, then the existing file(s) will be read, skipping the processing. When writing the file, extensions supplied are ignored and extensions are determined based on 'dest'.
BPPARAM	<a href="#">BiocParallelParam</a> object to specify multithreading to convert raw char in some parquet files to R objects. Not used otherwise.
return	Logical, whether to return the geometries in memory. This does not depend on whether the geometries are written to file. Always 'FALSE' when 'dest = "colGeometry"'.
save_memory	Don't load transcript spots of all genes at once. Not fully implemented here.
partition	Whether to partition the output by gene.
sfe	A 'SpatialFeatureExperiment' object.
sample_id	Which sample in the SFE object the transcript spots should be added to.

**Value**

A sf data frame for vector geometries if 'file\_out' is not set. 'SpatRaster' for raster. If there are multiple files written, such as when splitting by cell compartment or when 'dest = "colGeometry"', then a directory with the same name as 'file\_out' will be created (but without the extension) and the files are written to that directory with informative names. 'parquet' files that can be read with 'st\_read' is written for vector geometries. When 'return = FALSE', the file name or directory (when there're multiple files) is returned.

The 'sf' data frame, or path to file where geometries are written if 'return = FALSE'.

**Note**

When 'dest = "colGeometry"', the geometries are always written to disk and not returned in memory, because this is essentially the gene count matrix, which is sparse. This kind of reformatting is implemented so users can read in MULTIPOINT geometries with transcript spots for each gene assigned to each cell for spatial point process analyses, where not all genes are loaded at once.

**Examples**

```
# Default arguments are for MERFISH
fp <- tempfile()
dir_use <- SFEData::VizgenOutput(file_path = fp)
g <- formatTxSpots(file.path(dir_use, "detected_transcripts.csv"))
unlink(dir_use, recursive = TRUE)

# For CosMX, note the colnames, also dest = "colGeometry"
# Results are written to the tx_spots directory
dir_use <- SFEData::CosMXOutput(file_path = fp)
cg <- formatTxSpots(file.path(dir_use, "Run5642_S3_Quarter_tx_file.csv"),
  dest = "colGeometry", z = "all",
  cell_col = c("cell_ID", "fov"),
  gene_col = "target", not_in_cell_id = "0",
  spatialCoordsNames = c("x_global_px", "y_global_px", "z"),
  file_out = file.path(dir_use, "tx_spots"))
# Cleanup
unlink(dir_use, recursive = TRUE)
```

---

formatTxTech

*Read and process transcript spots for specific commercial technologies*


---

**Description**

To preset parameters such as spatialCoordsNames, gene\_col, cell\_col, and phred\_col that are standard for the output of the technology.

**Usage**

```
formatTxTech(
  data_dir,
  tech = c("Vizgen", "Xenium", "CosMX"),
  dest = c("rowGeometry", "colGeometry"),
  z = "all",
```

```

    min_phred = 20,
    split_cell_comps = FALSE,
    z_option = c("3d", "split"),
    flip = FALSE,
    file_out = NULL,
    BPPARAM = SerialParam(),
    return = TRUE
)

addTxTech(
  sfe,
  data_dir,
  sample_id = 1L,
  tech = c("Vizgen", "Xenium", "CosMX"),
  z = "all",
  min_phred = 20,
  split_cell_comps = FALSE,
  z_option = c("3d", "split"),
  flip = FALSE,
  file_out = NULL,
  BPPARAM = SerialParam()
)

```

### Arguments

<code>data_dir</code>	Top level output directory.
<code>tech</code>	Which technology whose output to read, must be one of "Vizgen", "Xenium", or "CosMX" though more technologies may be added later.
<code>dest</code>	Where in the SFE object to store the spot geometries. This affects how the data is processed. Options:  <b>rowGeometry</b> All spots for each gene will be a 'MULTIPOINT' geometry, regardless of whether they are in cells or which cells they are assigned to. <b>colGeometry</b> The spots for each gene assigned to a cell of interest will be a 'MULTIPOINT' geometry; since the gene count matrix is sparse, the geometries are NOT returned to memory.
<code>z</code>	Which z-planes to read. Always "all" for Xenium where the z coordinates are not discrete.
<code>min_phred</code>	Minimum Phred score to keep spot. By default 20, the conventional threshold indicating "acceptable", meaning that there's 1 chance that the spot was decoded in error.
<code>split_cell_comps</code>	Only relevant to CosMX whose transcript spot file assigns the spots to cell components. Setting this argument to TRUE
<code>z_option</code>	What to do with z coordinates. "3d" is to construct 3D geometries. "split" is to create a separate 2D geometry for each z-plane so geometric operations are fully supported but some data wrangling is required to perform 3D analyses. When the z coordinates are not integers, 3D geometries will always be constructed since there are no z-planes to speak of. This argument does not apply when 'spatialCoordsNames' has length 2.

flip	Logical, whether to flip the geometry to match image. Here the y coordinates are simply set to -y, so the original bounding box is not preserved. This is consistent with readVizgen and readXenium.
file_out	Name of file to save the geometry or raster to disk. Especially when the geometries are so large that it's unwieldy to load everything into memory. If this file (or directory for multiple files) already exists, then the existing file(s) will be read, skipping the processing. When writing the file, extensions supplied are ignored and extensions are determined based on 'dest'.
BPPARAM	<a href="#">BiocParallelParam</a> object to specify multithreading to convert raw char in some parquet files to R objects. Not used otherwise.
return	Logical, whether to return the geometries in memory. This does not depend on whether the geometries are written to file. Always 'FALSE' when 'dest = "colGeometry"'.
sfe	A 'SpatialFeatureExperiment' object.
sample_id	Which sample in the SFE object the transcript spots should be added to.

### Value

The 'sf' data frame, or path to file where geometries are written if 'return = FALSE'.

### Examples

```
library(SFData)
fp <- tempfile()
dir_use <- XeniumOutput("v2", file_path = fp)
fn_tx <- formatTxTech(dir_use, tech = "Xenium", flip = TRUE, return = FALSE,
                      file_out = file.path(dir_use, "tx_spots.parquet"))
```

---

gdalParquetAvailable *Check if Parquet GDAL driver is available*

---

### Description

The GeoParquet files for geometries are typically written and read with the sfarrow package, but to add only a select few genes to the SFE object say for visualization purposes, the Parquet GDAL driver is required in order to use GDAL's SQL to query the GeoParquet file to only load the few genes requested. The transcript spots from a large dataset can take up a lot of memory if all loaded.

### Usage

```
gdalParquetAvailable()
```

### Details

The Parquet driver has been supported since GDAL 3.5.0. The arrow C++ library must be installed in order to make the Parquet driver available. When arrow is installed, newer versions of GDAL installed from Homebrew (Mac) should have the Parquet driver. For Linux, the binary from apt-get's default repo is 3.4.1 (as of April 2024). To use the Parquet driver, GDAL may need to be installed from source. See script from the [geospatial rocker](#). A Voyager docker container with the Parquet driver will soon be provided.

**Value**

Logical, indicating whether the Parquet driver is present.

**Examples**

```
gdalParquetAvailable()
```

---

getParams

*Get parameters used in spatial methods*

---

**Description**

The getParams function allows users to access the parameters used to compute the results that may be stored in [colFeatureData](#).

**Usage**

```
getParams(
  sfe,
  name,
  local = FALSE,
  colData = FALSE,
  colGeometryName = NULL,
  annotGeometryName = NULL,
  reducedDimName = NULL
)
```

**Arguments**

sfe	A SpatialFeatureExperiment object.
name	Name used to store the results.
local	Logical, whether the results of interest come from a local spatial method.
colData	Logical, whether the results were computed for a column of colData(sfe).
colGeometryName	To get results for a colGeometry.
annotGeometryName	To get results for an annotGeometry; colGeometry has precedence so this argument is ignored if colGeometryName is specified.
reducedDimName	Name of a dimension reduction, can be seen in <a href="#">reducedDimNames</a> . colGeometryName and annotGeometryName have precedence over reducedDimName.

**Value**

A named list showing the parameters

**Examples**

```
library(SFEData)
library(scater)
library(Voyager)
sfe <- McKellarMuscleData("small")
colGraph(sfe, "visium") <- findVisiumGraph(sfe)
sfe <- colDataMoransI(sfe, "nCounts")
getParams(sfe, "morán", colData = TRUE)
```

---

getPixelSize	<i>Get physical size of pixels</i>
--------------	------------------------------------

---

**Description**

This function gets physical size of pixels in each resolution of a OME-TIFF pyramid in [BioFormatsImage](#).

**Usage**

```
getPixelSize(file, resolution = 1L)
```

**Arguments**

file	Path to an OME-TIFF file.
resolution	Which resolution to query; 1 means the highest resolution. The pixels will be larger for the lower resolutions.

**Value**

Numeric vector of length 2 of pixel size in x and y. Usually they're the same.

**Examples**

```
library(SFEData)
fp <- tempfile()
dir_use <- XeniumOutput("v1", file_path = fp)
# RBioFormats null pointer error
try(getPixelSize(file.path(dir_use, "morphology_focus.ome.tif")))
getPixelSize(file.path(dir_use, "morphology_focus.ome.tif"))
unlink(dir_use, recursive = TRUE)
```

---

getTechTxFields      *Get relevant fields and file paths for transcript spots*

---

### Description

Get column names for x, y, and z coordinates, gene IDs, and cell IDs from the transcript file and get file paths for transcript spot coordinates given technology.

### Usage

```
getTechTxFields(tech, data_dir = NULL)
```

### Arguments

tech	Name of the commercial technology, must be one of Vizgen, Xenium, and CosMX.
data_dir	Top level directory of the output.

### Value

A named list with elements:

spatialCoordsNames	A character vector for column names for the xyz coordinates of the transcript spots.
gene_col	Column name for gene IDs.
cell_col	Column name for cell IDs.
fn	File path of the transcript spot file.

---

getTissueBoundaryConcave  
*Get tissue boundary from concave hull of cell geometries*

---

### Description

The concave hull will be a smoothed outline, and the smoothness can be adjusted with the ratio parameter. Run [findDebrisCells](#) to remove small bits outside the main piece tissue before running this function.

### Usage

```
## S4 method for signature 'SpatialFeatureExperiment'
getTissueBoundaryConcave(
  x,
  sample_id = NULL,
  colGeometryName = 1L,
  ratio = 0.01,
  allow_holes = TRUE,
  multiple_pieces = FALSE,
  min_cells = 100,
```

```

    distance_cutoff = 50,
    BNPARAM = NULL,
    BPPARAM = SerialParam()
)

## S4 method for signature 'sfc'
getTissueBoundaryConcave(
  x,
  ratio = 0.01,
  allow_holes = TRUE,
  multiple_pieces = FALSE,
  min_cells = 100,
  distance_cutoff = 50,
  BNPARAM = NULL,
  BPPARAM = SerialParam()
)

## S4 method for signature 'sf'
getTissueBoundaryConcave(
  x,
  ratio = 0.01,
  allow_holes = TRUE,
  multiple_pieces = FALSE,
  min_cells = 100,
  distance_cutoff = 50,
  BNPARAM = NULL,
  BPPARAM = SerialParam()
)

```

### Arguments

x	object of class <code>sfg</code> , <code>sfc</code> or <code>sf</code>
sample_id	Sample id(s) whose tissue boundaries are to be found.
colGeometryName	Name of the <code>colGeometry</code> to use to infer the concave hull.
ratio	numeric; fraction convex: 1 returns the convex hulls, 0 maximally concave hulls
allow_holes	logical; if TRUE, the resulting concave hull may have holes
multiple_pieces	Logical, whether there are multiple pieces of tissue in the same sample.
min_cells	Minimum number of cells per graph component; components with fewer than this number of cells are considered debris and removed.
distance_cutoff	For the <code>sf</code> and <code>sfc</code> methods, the distance cutoff used to construct a distance-based spatial neighborhood graph, in the same unit as the spatial coordinates. Anything within this distance is considered a neighbor.
BNPARAM	A <code>BiocNeighborParam</code> object specifying the algorithm to find <code>k</code> nearest neighbors and distance based neighbors with <code>nn_method = "bioc"</code> . For distance based neighbors, only <code>KmknParam</code> and <code>VptreeParam</code> are applicable.
BPPARAM	A <code>BiocParallelParam</code> object for multithreading. Only used for <code>k</code> nearest neighbor and distance based neighbor with <code>nn_method = "bioc"</code> .

**Value**

A sf data frame with columns `sample_id` and `geometry`.

---

`getTissueBoundaryImg` *Get tissue boundary from histology image*

---

**Description**

This function gets the tissue boundary from image and makes sure that it is properly aligned with the geometries in the SFE object. Note that it will not work on fluorescent images when the fluorescence looks sparse. In that case, use [getTissueBoundaryConcave](#).

**Usage**

```
getTissueBoundaryImg(
  sfe,
  sample_id = NULL,
  image_id = NULL,
  image_type = c("brightfield", "fluorescent"),
  channel = NULL,
  n_pieces = 1,
  resolution = 4,
  maxcell = 1e+07,
  fill_holes = FALSE,
  simplify = TRUE,
  dTolerance = 0
)
```

**Arguments**

<code>sfe</code>	An SFE object with images
<code>sample_id</code>	Sample id(s) whose tissue boundaries are to be found.
<code>image_id</code>	ID of image to use to get boundary.
<code>image_type</code>	Character, either "brightfield" or "fluorescent"
<code>channel</code>	Channel to use for tissue segmentation. If NULL use average of all channels.
<code>n_pieces</code>	Number of pieces; only this number of largest pieces are kept. Smaller pieces will be considered debris and removed. Can be a vector of the same length as <code>sample_id</code> , or if <code>sample_id</code> is "all" then same length as the number of samples in the SFE object to specify different number of pieces in different samples. If <code>n_pieces</code> is length 1 while there are multiple samples, then the same number is applied to all samples.
<code>resolution</code>	Integer, which resolution to use for tissue boundary in a pyramidal OME-TIFF stack. Only applicable to <code>BioFormatsImage</code> . Note that the image will be loaded into memory and you usually don't need the highest resolution for the tissue boundary.
<code>maxcell</code>	Max number of pixels when loading <code>SpatRasterImage</code> into memory.
<code>fill_holes</code>	Logical, whether to fill holes in the tissue, to only get the outer outline.
<code>simplify</code>	Logical, whether to simplify the output polygon.
<code>dTolerance</code>	Distance tolerance when simplifying the polygon, in the same unit as the geometries in the SFE object.

**Value**

A sf data frame with columns `sample_id` and `geometry`.

---

imageIDs	<i>Show all image_ids in the SFE object</i>
----------	---

---

**Description**

The title is self-explanatory. Some functions require `image_id` to get or set images.

**Usage**

```
imageIDs(sfe)
```

**Arguments**

`sfe` A `SpatialFeatureExperiment` object.

**Value**

A character vector of `image_ids`.

**Examples**

```
fp <- system.file(file.path("extdata", "sample01"),
  package = "SpatialFeatureExperiment")
sfe <- read10xVisiumSFE(fp, type = "sparse")
imageIDs(sfe)
```

---

<code>Img&lt;-</code> , <code>SpatialExperiment</code> -method	<i>Image setter</i>
--	---------------------

---

**Description**

Modify or replace images stored in a `SpatialExperiment` object. This is different from `addImg` which adds the image from files and can't replace existing images, which is there to be consistent with `SpatialExperiment`. This setter here can replace existing images with another object that inherits from `VirtualSpatialImage`, including `SpatRasterImage`, `BioFormatsImage`, and `ExtImage`.

**Usage**

```
## S4 replacement method for signature 'SpatialExperiment'
Img(x, sample_id = 1L, image_id, scale_fct = 1) <- value
```

**Arguments**

x	A SpatialExperiment object, which includes SFE.
sample_id	Which sample the image is associated with. Use <a href="#">sampleIDs</a> to get sample IDs present in the SFE object.
image_id	Image ID, such as "lowres" and "hires" for Visium data and "DAPI" and "PolyT" for Vizgen MERFISH data.
scale_fct	Scale factor to convert pixels in lower resolution to those in the full resolution. Only relevant to image classes implemented in SpatialExperiment but not SpatialFeatureExperiment because the spatial extent of images in SFE takes precedence.
value	New version of image to add, must inherit from VirtualSpatialImage.

**Value**

SFE object with the new image added.

**Examples**

```
library(EBImage)
library(SFEData)
library(RBioFormats)
fp <- tempfile()
fn <- XeniumOutput("v2", file_path = fp)
# Weirdly the first time I get the null pointer error
try(sfe <- readXenium(fn))
sfe <- readXenium(fn)
img <- getImg(sfe) |> toExtImage(resolution = 1L)
img <- img[, ,1] > 500
Img(sfe, image_id = "mask") <- img
imageIDs(sfe)
unlink(fn, recursive = TRUE)
```

---

imgRaster

*Get the image from \*Image class*


---

**Description**

In SFE, S4 classes inheriting from VirtualSpatialImage have been implemented to make these image classes compatible with SpatialExperiment.

**Usage**

```
## S4 method for signature 'SpatRasterImage'
imgRaster(x, maxcell = 1e+07, col = terra::map.pal("viridis", 100))

## S4 method for signature 'BioFormatsImage'
imgRaster(x, resolution = 4L)

## S4 method for signature 'ExtImage'
imgRaster(x)
```

**Arguments**

x	An object of class *Image as implemented in this package.
maxcell	positive integer. Maximum number of cells to use for the plot
col	vector of colors. The default is <code>map.pal("viridis", 100)</code>
resolution	Resolution to read in from OME-TIFF, defaults to 4, which is a medium resolution in Xenium.

**Value**

Since version 1.9.8, `imgRaster` will return an array of hex colors, or the raster object, as required by `SpatialExperiment`. This will break older SFE code calling `imgRaster`.

**See Also**

Other image methods: [SFE-image](#), [affineImg\(\)](#), [cropImg\(\)](#), [dim,BioFormatsImage-method](#), [dim,ExtImage-method](#), [ext\(\)](#), [imgSource\(\)](#), [mirrorImg\(\)](#), [rotateImg\(\)](#), [scaleImg\(\)](#), [translateImg\(\)](#), [transposeImg\(\)](#)

---

imgSource

*Source of images that are on disk*


---

**Description**

Get the file path of images that are on disk and not read into memory. Only applies to `SpatRasterImage` and `BioFormatsImage`.

**Usage**

```
## S4 method for signature 'SpatRasterImage'
imgSource(x)
```

```
## S4 method for signature 'BioFormatsImage'
imgSource(x)
```

```
## S4 method for signature 'ExtImage'
imgSource(x)
```

**Arguments**

x	An object of class *Image as implemented in this package.
---	---

**Value**

String, file path to the original image on disk. For `SpatRasterImage`, if the image is loaded into memory, then NULL.

**See Also**

Other image methods: [SFE-image](#), [affineImg\(\)](#), [cropImg\(\)](#), [dim,BioFormatsImage-method](#), [dim,ExtImage-method](#), [ext\(\)](#), [imgRaster\(\)](#), [mirrorImg\(\)](#), [rotateImg\(\)](#), [scaleImg\(\)](#), [translateImg\(\)](#), [transposeImg\(\)](#)

---

internal-Voyager      *Internal functions also used in Voyager and Wayfarer*

---

### Description

Not meant for the user, but exporting to be used internally in Voyager. But one day I may clean these up and remove the internal note for people building on top of SFE.

### Usage

```
.value2df(value, use_geometry, feature = NULL)

.check_features(x, features, colGeometryName = NULL, swap_rownames = NULL)

.warn_symbol_duplicate(x, symbols, swap_rownames = "symbol")

.symbol2id(x, features, swap_rownames)

.check_sample_id(x, sample_id, one = TRUE, mustWork = TRUE)

.rm_empty_geometries(g, MARGIN)

.check_rg(type, x, sample_id)

.ext_(x)

.check_tx_file(
  file,
  spatialCoordsNames,
  gene_col,
  phred_col,
  min_phred,
  flip,
  BPPARAM = SerialParam(),
  save_memory = FALSE
)

.check_tx_file_mem(
  file,
  spatialCoordsNames,
  gene_col,
  phred_col,
  min_phred,
  flip,
  BPPARAM = SerialParam()
)
```

### Value

Internal

---

 localResults

*Get and set results from local spatial statistics*


---

### Description

Local spatial statistics like local Moran's I, local Geary's C, Getis-Ord  $G_i^*$ , and geographically weighted summary statistics return values at each spatial location. Just like dimension reductions, these results are clearly associated with the broader SFE object, so they should have a place within the object. However, a separate field is needed because these analyses are conceptually distinct from dimension reduction. Also, each feature (e.g. gene) can have its own results with values at each location. The localResults field in the SFE object stores these results that has a value for each spatial location.

### Usage

```
## S4 method for signature 'SpatialFeatureExperiment'
localResults(
  x,
  sample_id = "all",
  name = "all",
  features = NULL,
  colGeometryName = NULL,
  annotGeometryName = NULL,
  withDimnames = TRUE,
  swap_rownames = NULL,
  ...
)

## S4 replacement method for signature 'SpatialFeatureExperiment'
localResults(
  x,
  sample_id = "all",
  name = "all",
  features = NULL,
  colGeometryName = NULL,
  annotGeometryName = NULL,
  withDimnames = TRUE,
  swap_rownames = NULL,
  ...
) <- value

## S4 method for signature 'SpatialFeatureExperiment'
localResultNames(x)

## S4 replacement method for signature 'SpatialFeatureExperiment,character'
localResultNames(x) <- value

## S4 method for signature 'SpatialFeatureExperiment'
localResultFeatures(
  x,
  type = 1L,
```

```

    colGeometryName = NULL,
    annotGeometryName = NULL,
    swap_rownames = NULL
  )

## S4 method for signature 'SpatialFeatureExperiment'
localResultAttrs(
  x,
  type = 1L,
  feature,
  colGeometryName = NULL,
  annotGeometryName = NULL,
  swap_rownames = NULL
)

## S4 method for signature 'SpatialFeatureExperiment'
localResult(
  x,
  type = 1L,
  feature,
  colGeometryName = NULL,
  annotGeometryName = NULL,
  sample_id = 1L,
  withDimnames = TRUE,
  simplify = TRUE,
  swap_rownames = NULL
)

## S4 replacement method for signature 'SpatialFeatureExperiment'
localResult(
  x,
  type = 1L,
  feature,
  colGeometryName = NULL,
  annotGeometryName = NULL,
  sample_id = 1L,
  withDimnames = TRUE
) <- value

```

### Arguments

<code>x</code>	A <code>SpatialFeatureExperiment</code> object.
<code>sample_id</code>	Sample ID to get or set geometries.
<code>name</code>	Name of the spatial method used, such as "localmoran".
<code>features</code>	Features whose local results to get or set, for <code>localResults</code> getter and setter for multiple features at a time.
<code>colGeometryName</code>	Which <code>colGeometry</code> to get or set local results.
<code>annotGeometryName</code>	Which <code>annotGeometry</code> to get or set local results.
<code>withDimnames</code>	Logical. If TRUE, then the <code>dimnames</code> ( <code>colnames</code> or <code>rownames</code> ) of the gene count matrix should correspond to row names of the <code>sf</code> data frames of interest.

swap_rownames	Name of a column in rowData to identify features instead of the row names of the SFE object. For example, if the row names of the SFE object are Ensembl IDs and gene symbols are in the "symbol" column in rowData, then putting "symbol" for this argument will use the gene symbols to identify which gene's local results to get or set.
...	Ignored
value	Values to set, should be either a matrix or a data frame.
type	Name or index of the spatial method used, such as "localmoran".
feature	Feature whose local results to get or set, for localResult getter and setter for one feature at a time.
simplify	Basically whether to return the content of the list rather than a list when the list only has one element, such as results for one type and one feature.

### Value

localResults returns a named list each element of which is a set of local results of interest. localResult returns a matrix or a data frame, whichever the original is when it's set. localResultNames returns a character vector. Setters return an SFE object with the desired field set. For genes and colData columns, the local results are stored in the localResults field in int\_colData, whereas for colGeometries and annotGeometries, the local results are stored as columns in the same sf data frames. localResultFeatures returns a character vector of names of features for which local results are available. localResultAttrs returns a character vector of the column names of the local results of one type for one feature. It returns NULL if the results are a vector.

### Examples

```
# Toy example
sfe <- readRDS(system.file("extdata/sfe_toy.rds",
  package = "SpatialFeatureExperiment"
))
# localResults functions are written for organizing results from local
# spatial statistics (see the Voyager package). But for the examples here,
# random toy matrices are used. The real results are often matrices, with a
# matrix for each feature.
library(S4Vectors)
set.seed(29)
toy_res1 <- matrix(rnorm(10),
  nrow = 5, ncol = 2,
  dimnames = list(colnames(sfe), c("meow", "purr")))
)
toy_res1b <- matrix(rgamma(10, shape = 2),
  nrow = 5, ncol = 2,
  dimnames = list(colnames(sfe), c("meow", "purr")))
)
toy_df1 <- DataFrame(gene1 = I(toy_res1), gene2 = I(toy_res1b))

toy_res2 <- matrix(rpois(10, lambda = 2),
  nrow = 5, ncol = 2,
  dimnames = list(colnames(sfe), c("sassy", "tortitude")))
)
toy_df2 <- DataFrame(gene1 = I(toy_res2))
# Set all local results
localResults(sfe) <- list(localmoran = toy_df1, Gistar = toy_df2)
# Get all local results
```

```

lrs <- localResults(sfe)

# Set results of the same type for multiple genes
localResults(sfe, name = "localmoran") <- toy_df1
# Can also use a list
localResults(sfe, name = "localmoran") <- as.list(toy_df1)
# Get results of the same type for multiple genes
lrs <- localResults(sfe, name = "localmoran", features = c("gene1", "gene2"))

# Set results for one type and one gene
localResult(sfe, "localmoran", feature = "gene1") <- toy_res1
# Get results for one type and one gene
lr <- localResult(sfe, "localmoran", feature = "gene1")

# Set results for a feature in colGeometries
cg_toy <- readRDS(system.file("extdata/cg_toy.rds",
  package = "SpatialFeatureExperiment"
))
colGeometry(sfe, "cg") <- cg_toy
localResult(sfe, "localmoran",
  feature = "gene1",
  colGeometryName = "cg"
) <- toy_res1
# Get results for a feature in colGeometries
lr <- localResult(sfe, "localmoran", "gene1", colGeometryName = "cg")

```

---

mirrorImg

*Mirror/flip images*


---

## Description

Flip images along the middle horizontal or vertical axis.

## Usage

```

## S4 method for signature 'SpatRasterImage'
mirrorImg(
  x,
  direction = c("vertical", "horizontal"),
  filename = "",
  maxcell = NULL,
  ...
)

## S4 method for signature 'BioFormatsImage'
mirrorImg(x, direction = c("vertical", "horizontal"), ...)

## S4 method for signature 'ExtImage'
mirrorImg(x, direction = c("vertical", "horizontal"), ...)

```

## Arguments

x                   SpatRaster or SpatVector

direction	character. Should (partially) match "vertical" to flip by rows, or "horizontal" to flip by columns
filename	character. Output filename
maxcell	Max number of pixels to load <code>SpatRasterImage</code> into memory. The default <code>1e7</code> is chosen because this is the approximate number of pixels in the medium resolution image at resolution = 4L in Xenium OME-TIFF to make different methods of this function consistent.
...	additional arguments for writing files as in <code>writeRaster</code>

**Value**

\*Image object of the same class.

**See Also**

Other image methods: [SFE-image](#), [affineImg\(\)](#), [cropImg\(\)](#), [dim,BioFormatsImage-method](#), [dim,ExtImage-method](#), [ext\(\)](#), [imgRaster\(\)](#), [imgSource\(\)](#), [rotateImg\(\)](#), [scaleImg\(\)](#), [translateImg\(\)](#), [transposeImg\(\)](#)

---

multi_listw2sparse	<i>Convert multiple listw graphs into a single sparse adjacency matrix</i>
--------------------	--

---

**Description**

Each sample in the SFE object has a separate spatial neighborhood graph. Spatial analyses performed jointly on multiple samples require a combined spatial neighborhood graph from the different samples, where the different samples would be disconnected components of the graph. This combined adjacency matrix can be used in MULTISPATI PCA.

**Usage**

```
multi_listw2sparse(listws)
```

**Arguments**

listws            A list of listw objects.

**Value**

A sparse `dgCMatrix` of the combined spatial neighborhood graph, with the original spatial neighborhood graphs of the samples on the diagonal. When the input is an SFE object, the rows and columns will match the column names of the SFE object.

---

read10xVisiumSFE

*Read 10X Visium data as SpatialFeatureExperiment*


---

### Description

Read Space Ranger output from Visium v1 (not HD) as a SpatialFeatureExperiment object, where spots are represented with polygons in the colGeometry called "spotPoly". Other geometries can be added later after the dataset is read. If data = "filtered", then spatial neighborhood graphs of the spots are also computed and stored in the colGraph called "visium" in all samples for downstream spatial analyses.

### Usage

```
read10xVisiumSFE(
  dirs,
  sample_id = paste0("sample", sprintf("%02d", seq_along(dirs))),
  type = c("HDF5", "sparse"),
  data = c("filtered", "raw"),
  images = c("lowres", "hires"),
  unit = c("full_res_image_pixel", "micron"),
  style = "W",
  zero.policy = NULL,
  row.names = c("id", "symbol"),
  flip = c("geometry", "image", "none"),
  read_spatial_enrichment = TRUE
)
```

### Arguments

dirs	Directory for each sample that contains the spatial and raw/filtered_feats_bc_matrix directories.
sample_id	Which sample(s) in the SFE object to use for the graph. Can also be "all", which means this function will compute the graph for all samples independently.
type	Either "HDF5", and the matrix will be represented as TENxMatrix, or "sparse", and the matrix will be read as a dgCMatix.
data	character string specifying whether to read in filtered (spots mapped to tissue) or raw data (all spots).
images	character vector specifying which images to include. Valid values are "lowres", "hires", "fullres", "detected", "aligned"
unit	Whether to use pixels in full resolution image or microns as the unit. If using microns, then spacing between spots in pixels will be used to convert the coordinates into microns, as the spacing is known to be 100 microns. This is used to plot scale bar.
style	style can take values "W", "B", "C", "U", "minmax" and "S"
zero.policy	default NULL, use global option value; if FALSE stop with error for any empty neighbour sets, if TRUE permit the weights list to be formed with zero-length weights vectors

row.names	String specifying whether to use Ensembl IDs ("ID") or gene symbols ("Symbol") as row names. For symbols, the Ensembl ID will be appended to disambiguate rows where the same symbol corresponds to multiple Ensembl IDs.
flip	Whether to flip the geometries or the images, because in <code>sf</code> and <code>terra</code> , the geometries use the Cartesian coordinates greater y coordinates going up, while in images, greater y values go down. Originally the Visium spots are in pixels in full res image. Either the image or the geometry needs to be flipped for them match in the Cartesian coordinate system.
read_spatial_enrichment	Logical, whether to read the 'spatial_enrichment.csv' file from Visium output if the file is present and add its contents to 'rowData'.

**Value**

A `SpatialFeatureExperiment` object. The images might need to be manually transposed and/or mirrored to match the spots in this version of this package.

**Note**

It is assumed that the images have not been cropped. Otherwise the images might not align with the spots.

**Examples**

```
dir <- system.file("extdata", package = "SpatialFeatureExperiment")

sample_ids <- c("sample01", "sample02")
samples <- file.path(dir, sample_ids, "outs")

list.files(samples[1])
list.files(file.path(samples[1], "spatial"))
(sfe <- read10xVisiumSFE(dirs = samples, sample_id = sample_ids,
  type = "sparse", data = "filtered"
))
```

---

readCosMX

*Read CosMX data into SFE*


---

**Description**

This function reads the standard CosMX output into an SFE object, as in "Basic Data Files" on the Nanostring website. For new version of CosMX, these files are the flat files in the AtoMX output.

**Usage**

```
readCosMX(
  data_dir,
  z = "all",
  sample_id = "sample01",
  min_area = NULL,
  add_molecules = FALSE,
```

```

split_cell_comps = FALSE,
BPPARAM = SerialParam(),
file_out = file.path(data_dir, "tx_spots.parquet"),
z_option = c("3d", "split")
)

```

## Arguments

<code>data_dir</code>	Top level output directory.
<code>z</code>	Integer z index or "all" to indicate which z-planes to read for the transcript spots.
<code>sample_id</code>	A character sample identifier, which matches the <code>sample_id</code> in <code>imgData</code> . The <code>sample_id</code> will also be stored in a new column in <code>colData</code> , if not already present. Default = <code>sample01</code> .
<code>min_area</code>	Minimum cell area in square microns or pixel units (eg for CosMX). Anything smaller will be considered artifact or debris and removed. Default to 'NULL', ie no filtering of polygons.
<code>add_molecules</code>	Logical, whether to add transcripts coordinates to an object.
<code>split_cell_comps</code>	Logical, whether to split transcript spot geometries by cell compartment. Only relevant when <code>'add_molecules = TRUE'</code> .
<code>BPPARAM</code>	A <code>BiocParallelParam</code> object specifying parallel processing backend and number of threads to use for parallelizable tasks: <ol style="list-style-type: none"> <li>1. To load cell segmentation from HDF5 files from different fields of view (FOVs) with multiple cores. A progress bar can be configured in the <code>BiocParallelParam</code> object. When there are numerous FOVs, reading in the geometries can be time consuming, so we recommend using a server and larger number of threads. This argument is not used if <code>use_cellpose = TRUE</code> and the parquet file is present.</li> <li>2. To get the largest piece and see if it's larger than <code>min_area</code> when there are multiple pieces in the cell segmentation for one cell.</li> </ol>
<code>file_out</code>	Name of file to save the geometry or raster to disk. Especially when the geometries are so large that it's unwieldy to load everything into memory. If this file (or directory for multiple files) already exists, then the existing file(s) will be read, skipping the processing. When writing the file, extensions supplied are ignored and extensions are determined based on <code>'dest'</code> .
<code>z_option</code>	What to do with z coordinates. "3d" is to construct 3D geometries. "split" is to create a separate 2D geometry for each z-plane so geometric operations are fully supported but some data wrangling is required to perform 3D analyses. When the z coordinates are not integers, 3D geometries will always be constructed since there are no z-planes to speak of. This argument does not apply when <code>'spatialCoordsNames'</code> has length 2.

## Value

An SFE object. Cell polygons are written to `'cell_boundaries_sf.parquet'` in `'data_dir'`. If reading transcript spots (`'add_molecules = TRUE'`), then the reformatted transcript spots are saved to file specified in the `'file_out'` argument, which is by default `'tx_spots.parquet'` in the same directory as the rest of the data.

**Examples**

```
fp <- tempfile()
dir_use <- SFEData::CosMXOutput(file_path = fp)
sfe <- readCosMX(dir_use, z = "all", add_molecules = TRUE)
# Clean up
unlink(dir_use, recursive = TRUE)
```

---

readSelectTx	<i>Read transcript spots of select genes</i>
--------------	--

---

**Description**

I speculate that in practice, the most common use of the transcript spots is visualization, and only a few genes can be visualized at a time or the spots will overcrowd. Then it doesn't make sense to load the transcript spots of all genes into memory as they can take up a lot of memory. The function `readSelectTx` reads transcript spots of select genes into R, and the function `addSelectTx` adds them to `rowGeometries` of the SFE object.

**Usage**

```
readSelectTx(file, gene_select, z = "all", z_option = c("3d", "split"))

addSelectTx(
  sfe,
  file,
  gene_select,
  sample_id = 1L,
  z = "all",
  z_option = c("3d", "split"),
  swap_rownames = NULL
)
```

**Arguments**

file	File path of a GeoParquet file (e.g. already reformatted with the <a href="#">formatTxSpots</a> or <a href="#">addTxSpots</a> function, should have already flipped to match image if necessary).
gene_select	Character vector of a subset of genes. If NULL, then all genes that have transcript spots are added. Only relevant when reading data from formatted files on disk. If specified, then <code>return = TRUE</code> .
z	Index of z plane to read. Can be "all" to read all z-planes into MULTIPOINT geometries with XYZ coordinates. If z values are not integer, then spots with all z values will be read.
z_option	What to do with z coordinates. "3d" is to construct 3D geometries. "split" is to create a separate 2D geometry for each z-plane so geometric operations are fully supported but some data wrangling is required to perform 3D analyses. When the z coordinates are not integers, 3D geometries will always be constructed since there are no z-planes to speak of. This argument does not apply when <code>'spatialCoordsNames'</code> has length 2.
sfe	A <code>'SpatialFeatureExperiment'</code> object.

`sample_id` Which sample in the SFE object the transcript spots should be added to.

`swap_rownames` Name of a column in `rowData(sfe)` to use as gene identifiers in place of the actual row names. In some cases this may be needed to match each transcript spot MULTIPOINT geometry to rows of `sfe`.

### Value

When there are multipel parquet files to be read, a list of sf data frames with MULTIPOINT geometry for genes selected. When there is only one file, then one sf data frame. For `addSelectTx`, an SFE object with the transcript spots of the selected genes added.

### Note

The GDAL Parquet driver is required for this function, though not for other functions that work with GeoParquet files. GDAL Parquet driver has been supported since GDAL 3.5.0, but is not part of the default installation. The `z` and `z_option` arguments are there since the file names contain z-plane information when relevant. See the [GDAL documentation page for the Parquet driver](#).

### Examples

```
library(SFEData)
if (gdalParquetAvailable()) {
  fp <- tempfile()
  dir_use <- XeniumOutput("v2", file_path = fp)
  fn_tx <- formatTxTech(dir_use, tech = "Xenium", flip = TRUE, return = FALSE,
                       file_out = file.path(dir_use, "tx_spots.parquet"))
  gene_select <- c("ACE2", "BMX")
  df <- readSelectTx(fn_tx, gene_select)
  # RBioFormats null pointer error the first time
  try(sfe <- readXenium(dir_use))
  sfe <- readXenium(dir_use)
  sfe <- addSelectTx(sfe, fn_tx, head(rownames(sfe), 5), swap_rownames = "Symbol")
  unlink(dir_use, recursive = TRUE)
}
```

---

readVisiumHD

*Read Visium HD data*

---

### Description

This function reads Visium HD Space Ranger output into R.

### Usage

```
readVisiumHD(
  data_dir,
  bin_size = 8L,
  sample_id = "sample01",
  use_cellseg = TRUE,
  type = c("HDF5", "sparse"),
  data = c("filtered", "raw"),
  images = c("lowres", "hires"),
  unit = c("full_res_image_pixel", "micron"),
```

```

    style = "W",
    zero.policy = TRUE,
    row.names = c("id", "symbol"),
    flip = c("geometry", "image"),
    add_graph = FALSE
  )

```

### Arguments

data_dir	Directory with Visium HD output
bin_size	One or more resolutions to load, must be 2, 8, or 16. Can be either integer or character.
sample_id	Which sample(s) in the SFE object to use for the graph. Can also be "all", which means this function will compute the graph for all samples independently.
use_cellseg	Logical, whether to use cell segmentation if present. Only applicable when the segmented_outputs directory is present. If TRUE and the directory is present, then bin_size will be ignored and cell segmentations will be used instead.
type	Either "HDF5", and the matrix will be represented as <code>TENxMatrix</code> , or "sparse", and the matrix will be read as a <code>dgCMatrix</code> .
data	character string specifying whether to read in filtered (spots mapped to tissue) or raw data (all spots).
images	character vector specifying which images to include. Valid values are "lowres", "hires", "fullres", "detected", "aligned"
unit	Whether to use pixels in full resolution image or microns as the unit. If using microns, then spacing between spots in pixels will be used to convert the coordinates into microns, as the spacing is known to be 100 microns. This is used to plot scale bar.
style	style can take values "W", "B", "C", "U", "minmax" and "S"
zero.policy	default NULL, use global option value; if FALSE stop with error for any empty neighbour sets, if TRUE permit the weights list to be formed with zero-length weights vectors
row.names	String specifying whether to use Ensembl IDs ("ID") or gene symbols ("Symbol") as row names. For symbols, the Ensembl ID will be appended to disambiguate rows where the same symbol corresponds to multiple Ensembl IDs.
flip	Whether to flip the geometries or the images, because in <code>sf</code> and <code>terra</code> , the geometries use the Cartesian coordinates greater y coordinates going up, while in images, greater y values go down. Originally the Visium spots are in pixels in full res image. Either the image or the geometry needs to be flipped for them match in the Cartesian coordinate system.
add_graph	Logical, if to add spatial neighborhood graph for spots and only if <code>c(data = "filtered")</code> . Default is FALSE. This is optional because for larger datasets, the graph can take a while to compute.

### Value

An SFE object if `length(bin_size) == 1L`, otherwise a list of SFE objects each element of which is for one bin size. They're not concatenated since it might not make sense to perform joint analyses on the different resolutions that benefit from having them in the same SFE object, unlike different biological replica. Here unlike in `read10xVisiumSFE`, the centroids geometry is also added because it will greatly facilitate plotting when there are many spots when not zooming in. See the `scattermore` argument in `plotSpatialFeature`.

## Examples

```
library(OSTA.data)
id <- "VisiumHD_HumanColon_Oliveira"
pa <- OSTA.data_load(id)
dir.create(dir <- tempfile())
unzip(pa, exdir=dir)

# Use binned output
sfe1 <- readVisiumHD(dir, bin_size = 16, use_cellseg = FALSE)

# Use cell segmentations
sfe <- readVisiumHD(dir, use_cellseg = TRUE, unit = "micron")

unlink(dir, recursive = TRUE)
```

---

readVizgen

*Read Vizgen MERFISH output as SpatialFeatureExperiment*


---

## Description

This function reads the standard Vizgen MERFISH output into an SFE object. The coordinates are in microns. Cell centroids are read into `colGeometry` "centroids", and cell segmentations are read into `colGeometry` "cellSeg". The image(s) (polyT, DAPI, and cell boundaries) are also read as `SpatRaster` objects so they are not loaded into memory unless necessary. Because the image's origin is the top left while the geometry's origin is bottom left, either the image or the geometry needs to be flipped. Because the image accompanying MERFISH datasets are usually very large, the coordinates will be flipped so the flipping operation won't load the entire image into memory. Large datasets with hundreds of thousands of cells can take a while to read if reading transcript spots as it takes a while to convert the spots to MULTIPOINT geometries.

## Usage

```
readVizgen(
  data_dir,
  z = "all",
  sample_id = "sample01",
  min_area = NULL,
  image = c("DAPI", "PolyT", "Cellbound"),
  flip = c("geometry", "image", "none"),
  max_flip = "50 MB",
  filter_counts = FALSE,
  add_molecules = FALSE,
  use_bboxes = FALSE,
  use_cellpose = TRUE,
  BPPARAM = SerialParam(),
  file_out = file.path(data_dir, "detected_transcripts.parquet"),
  z_option = c("3d", "split")
)
```

**Arguments**

data_dir	Top level output directory.
z	Integer, z index to read, or "all", indicating z-planes of the images and transcript spots to read. While cell segmentation seems to have multiple z-planes, the segmentation in all z-planes are the same so in effect the cell segmentation is only in 2D.
sample_id	A character sample identifier, which matches the sample_id in <code>imgData</code> . The sample_id will also be stored in a new column in <code>colData</code> , if not already present. Default = sample01.
min_area	Minimum cell area in square microns or pixel units (eg for CosMX). Anything smaller will be considered artifact or debris and removed. Default to 'NULL', ie no filtering of polygons.
image	Which image(s) to load, can be "DAPI", "PolyT", "Cellbound" or any combination of them.
flip	To flip the image, geometry coordinates, or none. Because the image has the origin at the top left while the geometry has origin at the bottom left, one of them needs to be flipped for them to match. If one of them is already flipped, then use "none". The image will not be flipped if it's GeoTIFF.
max_flip	Maximum size of the image allowed to flip the image. Because the image will be loaded into memory to be flipped. If the image is larger than this size then the coordinates will be flipped instead.
filter_counts	Logical, whether to keep cells with counts > 0.
add_molecules	Logical, whether to add transcripts coordinates to an object.
use_bboxes	If no segmentation output is present, use cell_metadata to make bounding boxes instead.
use_cellpose	Whether to read the parquet files from CellPose cell segmentation. If FALSE, cell segmentation will be read from the HDF5 files. Note that reading HDF5 files for numerous FOVs is very slow.
BPPARAM	A <code>BiocParallelParam</code> object specifying parallel processing backend and number of threads to use for parallelizable tasks: <ol style="list-style-type: none"> <li>1. To load cell segmentation from HDF5 files from different fields of view (FOVs) with multiple cores. A progress bar can be configured in the <code>BiocParallelParam</code> object. When there are numerous FOVs, reading in the geometries can be time consuming, so we recommend using a server and larger number of threads. This argument is not used if use_cellpose = TRUE and the parquet file is present.</li> <li>2. To get the largest piece and see if it's larger than min_area when there are multiple pieces in the cell segmentation for one cell.</li> </ol>
file_out	Name of file to save the geometry or raster to disk. Especially when the geometries are so large that it's unwieldy to load everything into memory. If this file (or directory for multiple files) already exists, then the existing file(s) will be read, skipping the processing. When writing the file, extensions supplied are ignored and extensions are determined based on 'dest'.
z_option	What to do with z coordinates. "3d" is to construct 3D geometries. "split" is to create a separate 2D geometry for each z-plane so geometric operations are fully supported but some data wrangling is required to perform 3D analyses. When the z coordinates are not integers, 3D geometries will always be constructed since there are no z-planes to speak of. This argument does not apply when 'spatialCoordsNames' has length 2.

**Value**

A SpatialFeatureExperiment object.

**Note**

Since the transcript spots file is often very large, we recommend only using `add_molecules = TRUE` on servers with a lot of memory. If reading all z-planes, conversion of transcript spot geometry to parquet file might fail due to arrow data length limit. In a future version, when the transcript spot geometry is large, it will be written to multiple separate parquet files which are then concatenated with DuckDB. Also, in a future version, the transcript spot processing function might be rewritten in C++ to stream the original CSV file so it's not entirely loaded into memory.

**Examples**

```
fp <- tempfile()
dir_use <- SFEData::VizgenOutput(file_path = fp)
sfe <- readVizgen(dir_use, z = 3L, image = "PolyT",
  flip = "geometry")

## Filtering of counts, and addition of molecule coordinates..
sfe <- readVizgen(dir_use, z = 3L, image = "PolyT", filter_counts = TRUE,
  add_molecules = TRUE, flip = "geometry")

unlink(dir_use, recursive = TRUE)
```

---

readXenium

*Read 10X Xenium output as SpatialFeatureExperiment*


---

**Description**

This function reads the standard 10X Xenium output into an SFE object.

**Usage**

```
readXenium(
  data_dir,
  sample_id = "sample01",
  min_area = NULL,
  image = c("morphology_focus", "morphology_mip"),
  segmentations = c("cell", "nucleus"),
  row.names = c("id", "symbol"),
  flip = c("geometry", "image", "none"),
  max_flip = "50 MB",
  filter_counts = FALSE,
  add_molecules = FALSE,
  min_phred = 20,
  BPPARAM = SerialParam(),
  file_out = file.path(data_dir, "tx_spots.parquet")
)
```

**Arguments**

data_dir	Top level output directory.
sample_id	A character sample identifier, which matches the <code>sample_id</code> in <code>imgData</code> . The <code>sample_id</code> will also be stored in a new column in <code>colData</code> , if not already present. Default = <code>sample01</code> .
min_area	Minimum cell area in square microns or pixel units (eg for CosMX). Anything smaller will be considered artifact or debris and removed. Default to 'NULL', ie no filtering of polygons.
image	Which image(s) to load, can be "morphology_mip", "morphology_focus" or both. Note that in Xenium Onboarding Analysis (XOA) v2, there is no longer "morphology_mip" and "morphology_focus" is a directory with 4 images corresponding to 4 channels: DAPI, "Cadherin", 18S, and Vimentin. So this argument is ignored for XOA v2.
segmentations	Which segmentation outputs to read, can be "cell", "nucleus", or both.
row.names	String specifying whether to use Ensembl IDs ("id") or gene symbols ("symbol") as row names. If using symbols, the Ensembl ID will be appended to disambiguate in case the same symbol corresponds to multiple Ensembl IDs. Always "symbol" if 'add_molecules = TRUE' because only gene symbols are used in the transcript spot files.
flip	To flip the image, geometry coordinates, or none. Because the image has the origin at the top left while the geometry has origin at the bottom left, one of them needs to be flipped for them to match. If one of them is already flipped, then use "none". The image will not be flipped if it's GeoTIFF.
max_flip	Maximum size of the image allowed to flip the image. Because the image will be loaded into memory to be flipped. If the image is larger than this size then the coordinates will be flipped instead.
filter_counts	Logical, whether to keep cells with counts > 0.
add_molecules	Logical, whether to add transcripts coordinates to an object.
min_phred	Minimum Phred score to keep spot. By default 20, the conventional threshold indicating "acceptable", meaning that there's 1 chance that the spot was decoded in error.
BPPARAM	A <code>BiocParallelParam</code> object specifying parallel processing backend and number of threads to use for parallelizable tasks: <ol style="list-style-type: none"> <li>1. To load cell segmentation from HDF5 files from different fields of view (FOVs) with multiple cores. A progress bar can be configured in the <code>BiocParallelParam</code> object. When there are numerous FOVs, reading in the geometries can be time consuming, so we recommend using a server and larger number of threads. This argument is not used if <code>use_cellpose = TRUE</code> and the parquet file is present.</li> <li>2. To get the largest piece and see if it's larger than <code>min_area</code> when there are multiple pieces in the cell segmentation for one cell.</li> </ol>
file_out	Name of file to save the geometry or raster to disk. Especially when the geometries are so large that it's unwieldy to load everything into memory. If this file (or directory for multiple files) already exists, then the existing file(s) will be read, skipping the processing. When writing the file, extensions supplied are ignored and extensions are determined based on 'dest'.

**Value**

An SFE object. If reading segmentations, the cell or nuclei segmentation will be saved to ‘cell\_boundaries\_sf.parquet’ and ‘nucleus\_boundaries\_sf.parquet’ respectively in ‘data.dir’ so next time the boundaries can be read much more quickly. If reading transcript spots (‘add\_molecules = TRUE’), then the reformatted transcript spots are saved to file specified in the ‘file\_out’ argument, which is by default ‘tx\_spots.parquet’ in the same directory as the rest of the data. If images are present, then the images will be of the BioFormatsImage class and not loaded into memory until necessary in later operations.

**Note**

Sometimes when reading images, you will see this error the first time: ‘java.lang.NullPointerException: Cannot invoke "loci.formats.DimensionSwapper.setMetadataFiltered(boolean)" because "RBioFormats.reader" is null’. See this issue <https://github.com/aoles/RBioFormats/issues/42> Rerun the code and it should work the second time.

**Examples**

```
library(SFEData)
library(RBioFormats)
fp <- tempfile()
dir_use <- XeniumOutput("v2", file_path = fp)
# RBioFormats issue
try(sfe <- readXenium(dir_use, add_molecules = TRUE))
sfe <- readXenium(dir_use, add_molecules = TRUE)
unlink(dir_use, recursive = TRUE)
```

---

reexports

*Functions re-exported from other packages*


---

**Description**

These are some commonly used getters and setters of classes that SFE inherits so you don’t have to separately attach those packages to use these functions.

**Usage**

```
colData(x, ...)

rowData(x, use.names = TRUE, ...)

colData(x, ...) <- value

spatialCoords(x, ...)

spatialCoords(x) <- value

spatialCoordsNames(x)

getImg(x, ...)
```

```
imgData(x)
rmvImg(x, ...)
counts(object, ...)
logcounts(object, ...)
reducedDim(x, type, ...)
```

### Arguments

x	A SummarizedExperiment object or derivative.
...	For assay, arguments in ... are forwarded to assays. For rbind, cbind, ... contains SummarizedExperiment objects (or derivatives) to be combined. For other accessors, ignored.
use.names	For rowData: Like <code>mcols(x)</code> , by default <code>rowData(x)</code> propagates the rownames of x to the returned <code>DataFrame</code> object (note that for a SummarizedExperiment object or derivative, the rownames are also the names i.e. <code>rownames(x)</code> is always the same as <code>names(x)</code> ). Setting <code>use.names=FALSE</code> suppresses this propagation i.e. it returns a <code>DataFrame</code> object with no rownames. Use this when <code>rowData(x)</code> fails, which can happen when the rownames contain NAs (because the rownames of a SummarizedExperiment object or derivative can contain NAs, but the rownames of a <code>DataFrame</code> object cannot). For <code>combineRows</code> and <code>combineCols</code> : See Combining section below.
value	An object of a class specified in the S4 method signature or as outlined in 'Details'.
object	A <code>SingleCellExperiment</code> object, which includes SFE.
type	Name or numeric index to indicate which <code>reducedDim</code> to get, such as "PCA". By default the first item in <code>reducedDims</code> .

---

removeEmptySpace	<i>Remove empty space</i>
------------------	---------------------------

---

### Description

For each sample independently, all geometries and `spatialCoords` are translated so the origin is at the minimum coordinates of the bounding box of all geometries of the sample. This way coordinates of different samples will be more comparable. This removes empty space in the images if present.

### Usage

```
removeEmptySpace(sfe, sample_id = "all")
```

### Arguments

sfe	An SFE object.
sample_id	Sample to remove empty space.

**Value**

An SFE object with empty space removed.

**Note**

Unlike other functions in this package, this function operates on all samples by default.

**Examples**

```
library(SFEData)
library(SingleCellExperiment)
sfe <- McKellarMuscleData("full")
# Only keep spots on tissue
sfe <- sfe[, colData(sfe)$in_tissue]
# Move the coordinates of the tissue
sfe <- removeEmptySpace(sfe)
```

---

rotateImg

*Rotate image*

---

**Description**

As in SpatialExperiment, rotation here must be a multiple of 90 degrees.

**Usage**

```
## S4 method for signature 'SpatRasterImage'
rotateImg(x, degrees, maxcell = 1e+07, ...)

## S4 method for signature 'BioFormatsImage'
rotateImg(x, degrees, ...)

## S4 method for signature 'ExtImage'
rotateImg(x, degrees, ...)
```

**Arguments**

x	An object of class *Image as implemented in this package.
degrees	How many degrees to rotate. Positive number means clockwise and negative number means counterclockwise.
maxcell	Max number of pixels to load SpatRasterImage into memory. The default 1e7 is chosen because this is the approximate number of pixels in the medium resolution image at resolution = 4L in Xenium OME-TIFF to make different methods of this function consistent.
...	Ignored. It's there so different methods can all be passed to the same lapply in the method for SFE objects. Some methods have extra arguments.

**Value**

SpatRasterImage will be loaded into memory and converted to ExtImage. Otherwise \*Image object of the same class.

**See Also**

Other image methods: [SFE-image](#), [affineImg\(\)](#), [cropImg\(\)](#), [dim,BioFormatsImage-method](#), [dim,ExtImage-method](#), [ext\(\)](#), [imgRaster\(\)](#), [imgSource\(\)](#), [mirrorImg\(\)](#), [scaleImg\(\)](#), [translateImg\(\)](#), [transposeImg\(\)](#)

---

 rotateMinRect

*Rotate the tissue to reduce empty space in plots*


---

**Description**

This function finds the minimum bounding rotated rectangle and then rotates the sample so the minimum bounding rotated rectangle is horizontal or vertical. This will reduce empty space in plots when the sample has a long shape.

**Usage**

```
## S4 method for signature 'matrix'
rotateMinRect(x, orientation = c("horizontal", "vertical"), ...)

## S4 method for signature 'sfc'
rotateMinRect(x, orientation = c("horizontal", "vertical"), ...)

## S4 method for signature 'sf'
rotateMinRect(x, orientation = c("horizontal", "vertical"), ...)

## S4 method for signature 'SpatialFeatureExperiment'
rotateMinRect(x, orientation = c("horizontal", "vertical"), maxcell = 1e+07)
```

**Arguments**

x	Object of interest to be rotated, can be a matrix with 2 columns, sf, sfc, or SpatialFeatureExperiment.
orientation	Whether you want the rotated sample to be horizontal or vertical.
...	Ignored
maxcell	Rotating SpatRasterImage will convert it into ExtImage, loading the image into memory. This argument specifies the maximum number of pixels in the image loaded into memory. The image will be down sampled to approximately this number of pixels.

**Value**

Object of the same class as x, rotated.

---

`rowGeometries`*Row geometry getters and setters*

---

### Description

`rowGeometries` are geometries that corresponding to rows of the gene count matrix, such as sm-FISH transcript spots. The `txSpots()` function is a convenience wrapper for transcript spots, although this entirely depends on the `rowGeometry` being named `txSpots`.

### Usage

```
rowGeometry(x, type = 1L, sample_id = 1L, withDimnames = TRUE)
```

```
rowGeometry(  
  x,  
  type = 1L,  
  sample_id = 1L,  
  withDimnames = TRUE,  
  partial = FALSE,  
  translate = TRUE  
) <- value
```

```
rowGeometries(x, sample_id = "all", withDimnames = TRUE)
```

```
rowGeometries(  
  x,  
  sample_id = "all",  
  withDimnames = TRUE,  
  partial = FALSE,  
  translate = TRUE  
) <- value
```

```
rowGeometryNames(x)
```

```
rowGeometryNames(x) <- value
```

```
txSpots(x, sample_id = 1L, withDimnames = TRUE)
```

```
txSpots(  
  x,  
  sample_id = 1L,  
  withDimnames = TRUE,  
  partial = FALSE,  
  translate = TRUE  
) <- value
```

### Arguments

`x` A `SpatialFeatureExperiment` object.

type	An integer specifying the index or string specifying the name of the *Geometry to query or replace. If missing, then the first item in the *Geometries will be returned or replaced.
sample_id	Sample ID to get or set geometries.
withDimnames	Logical. If TRUE, then the dimnames (colnames or rownames) of the gene count matrix should correspond to row names of the sf data frames of interest.
partial	In setters, if a rowGeometry of the same name exists, whether to only replace the rows present in value.
translate	Logical. Only used if <code>removeEmptySpace</code> has been run of the SFE object. If that's the case, this argument indicates whether the new value to be assigned to the geometry is in the coordinates prior to removal of empty space so it should be translated to match the new coordinates after removing empty space. Default to TRUE.
value	Value to set. For dimGeometry, must be a sf data frame with the same number of rows as size in the dimension of interest, or an ordinary data frame that can be converted to such a sf data frame (see <code>df2sf</code> ). For dimGeometries, must be a list of such sf or ordinary data frames.

## Details

When there are multiple samples in the SFE object, rowGeometries for each sample has the `sample_id` appended to the name of the geometry. For example, if the name is `txSpots` and the sample ID is `sample01`, then the actual name of the rowGeometry is `txSpots_sample01`. In the getter, one can still specify `rowGeometry(sfe, "txSpots", sample_id = "sample01")`.

Appending the `sample_id` is unnecessary when there is only one sample, but `sample_id` will be appended when to SFE objects are combined with `cbind`. It is necessary to distinguish between different samples because they can have overlapping coordinate values.

## See Also

`[dimGeometries()]`, `[colGeometries()]`

## Examples

```
library(SFEData)
library(RBioFormats)
fp <- tempfile()
dir_use <- XeniumOutput("v2", file_path = fp)
# RBioFormats issue
try(sfe <- readXenium(dir_use, add_molecules = TRUE))
sfe <- readXenium(dir_use, add_molecules = TRUE)
rowGeometries(sfe)
rowGeometryNames(sfe)
tx <- rowGeometry(sfe, "txSpots")
txSpots(sfe)
unlink(dir_use, recursive = TRUE)
```

---

sampleIDs	<i>Get all unique sample IDs</i>
-----------	----------------------------------

---

**Description**

The title is self-explanatory.

**Usage**

```
sampleIDs(sfe)
```

**Arguments**

sfe                    A SpatialFeatureExperiment object.

**Value**

A character vector of all unique entries of the sample\_id column in colData(x).

**Examples**

```
library(SFEData)
sfe <- McKellarMuscleData(dataset = "small")
sampleIDs(sfe)
```

---

saveRDS,SpatialFeatureExperiment-method	<i>Save SpatialFeatureExperiment as RDS file</i>
---	--

---

**Description**

Saving SFE objects as RDS files is complicated by the SpatRaster class of the images. If present, the images need to be wrapped with the [wrap](#) function in terra before serializing the SFE object. Otherwise the images will be invalid pointers when the RDS is reloaded. If the image does not fit in memory and its file source is unknown, then it will be written to a temporary file, which is reloaded when the RDS file is loaded. When an SFE object with images is read from an RDS file, the images will not be unwrapped until necessary.

**Usage**

```
## S4 method for signature 'SpatialFeatureExperiment'
saveRDS(
  object,
  file = "",
  ascii = FALSE,
  version = NULL,
  compress = TRUE,
  refhook = NULL
)
```

**Arguments**

object	A SpatialFeatureExperiment object.
file	a <a href="#">connection</a> or the path name of the file where the R object is saved to or read from.
ascii	a logical. If TRUE or NA, an ASCII representation is written; otherwise (default), a binary one is used. See the comments in the help for <a href="#">save</a> .
version	the workspace format version to use. NULL specifies the current default version (3). The only other supported value is 2, the default from R 1.4.0 to R 3.5.0.
compress	a logical specifying whether saving to a named file is to use "gzip" compression, or one of "gzip", "bzip2", "xz" or "zstd" to indicate the type of compression to be used. Ignored if file is a connection.
refhook	a hook function for handling reference objects. See <a href="#">unserialize</a> .

**Value**

Invisibly NULL.

**Examples**

```
outdir <- system.file("extdata", package = "SpatialFeatureExperiment")
samples <- file.path(outdir, paste0("sample0", 1:2))
sfe <- read10xVisiumSFE(samples, type = "sparse", data = "filtered")
saveRDS(sfe, "foo.rds")
# Clean up
file.remove("foo.rds")
```

---

scaleImg

*Scale image*

---

**Description**

This function scales the image about its center. After scaling, the center of the image is not shifted.

**Usage**

```
## S4 method for signature 'AlignedSpatialImage'
scaleImg(x, factor, ...)

## S4 method for signature 'BioFormatsImage'
scaleImg(x, factor, ...)
```

**Arguments**

x	An object of class *Image as implemented in this package.
factor	Numeric, scaling factor.
...	Ignored. It's there so different methods can all be passed to the same lapply in the method for SFE objects. Some methods have extra arguments.

**Value**

A \*Image object of the same class that has been scaled. Behind the scene, it's only the extent that has been changed and the images are not changed. The center of the image is unchanged.

**See Also**

Other image methods: [SFE-image](#), [affineImg\(\)](#), [cropImg\(\)](#), [dim,BioFormatsImage-method,dim,ExtImage-method,ext\(\)](#), [imgRaster\(\)](#), [imgSource\(\)](#), [mirrorImg\(\)](#), [rotateImg\(\)](#), [translateImg\(\)](#), [transposeImg\(\)](#)

---

SFE-image

*Methods for handling image-related data*


---

**Description**

Generics of these functions are defined in SpatialExperiment, except for transposeImg. These SFE methods cater to the new image-related classes in SFE. The SPE method for getImg, rmvImg, and imgRaster don't need to be modified for SFE and are hence not implemented here, but are simply re-exported.

**Usage**

```
## S4 method for signature 'SpatialFeatureExperiment'
addImg(x, imageSource, sample_id = 1L, image_id, extent = NULL, scale_fct = 1)
```

```
## S4 method for signature 'SpatialFeatureExperiment'
transposeImg(
  x,
  sample_id = 1L,
  image_id = NULL,
  maxcell = 1e+07,
  filename = ""
)
```

```
## S4 method for signature 'SpatialFeatureExperiment'
mirrorImg(
  x,
  sample_id = 1L,
  image_id = NULL,
  direction = "vertical",
  maxcell = 1e+07,
  filename = ""
)
```

```
## S4 method for signature 'SpatialFeatureExperiment'
rotateImg(x, sample_id = 1L, image_id = NULL, degrees, maxcell = 1e+07)
```

```
## S4 method for signature 'SpatialFeatureExperiment'
translateImg(x, sample_id = 1L, image_id = NULL, v)
```

```
## S4 method for signature 'SpatialFeatureExperiment'
```

```

scaleImg(x, sample_id = 1L, image_id = NULL, factor)

## S4 method for signature 'SpatialFeatureExperiment'
affineImg(x, sample_id = 1L, image_id = NULL, M, v)

```

### Arguments

x	A SFE object.
imageSource	a character string specifying an image file name (.png, .jpg or .tif) or URL to source the image from
sample_id	Which sample the image is associated with. Use <a href="#">sampleIDs</a> to get sample IDs present in the SFE object.
image_id	Image ID, such as "lowres" and "hires" for Visium data and "DAPI" and "PolyT" for Vizgen MERFISH data.
extent	A numeric vector of length 4 with names of the set xmin, ymin, xmax, and ymax, specifying the extent of the image.
scale_fct	Scale factor – multiply pixel coordinates in full resolution image by this scale factor should yield pixel coordinates in a different resolution. extent takes precedence over scale_fct.
maxcell	Max number of pixels to load SpatRasterImage into memory. The default 1e7 is chosen because this is the approximate number of pixels in the medium resolution image at resolution = 4L in Xenium OME-TIFF to make different methods of this function consistent.
filename	character. Output filename
direction	character. Should (partially) match "vertical" to flip by rows, or "horizontal" to flip by columns
degrees	How many degrees to rotate. Positive number means clockwise and negative number means counterclockwise.
v	A numeric vector of length 2 specifying the vector in the xy plane to translate the SFE object.
factor	Numeric, scaling factor.
M	A 2x2 numeric matrix for the linear transformation in the xy plane.

### Details

Method of [transposeImg](#), [mirrorImg](#), and [rotateImg](#) perform the method on all images within the SFE object that are specified with `sample_id` and `image_id`. For images that are not loaded into memory, `rotateImg` will load `SpatRasterImage` into memory and all image operations except translate will load `BioFormatsImage` into memory.

### Note

If the image is already a GeoTIFF file that already has an extent, then the extent associated with the file will be honored and the `extent` and `scale_fct` arguments are ignored. Transposing the image is just like transposing a matrix. It's flipped about the line going from the top left to the bottom right.

**See Also**

Other image methods: [affineImg\(\)](#), [cropImg\(\)](#), [dim,BioFormatsImage-method](#), [dim,ExtImage-method](#), [ext\(\)](#), [imgRaster\(\)](#), [imgSource\(\)](#), [mirrorImg\(\)](#), [rotateImg\(\)](#), [scaleImg\(\)](#), [translateImg\(\)](#), [transposeImg\(\)](#)

**Examples**

```
library(SFEData)
sfe <- McKellarMuscleData("small")
img_path <- system.file(file.path("extdata", "sample01", "outs", "spatial",
"tissue_lowres_image.png"), package = "SpatialFeatureExperiment")
sfe <- addImg(sfe, img_path, sample_id = "Vis5A", image_id = "lowres", scale_fct =
0.023)
img <- getImg(sfe)
# SpatRasterImage method
img_t <- transposeImg(img)
# SFE method
sfe <- transposeImg(sfe, sample_id = "Vis5A", image_id = "lowres")
```

---

SFE-transform

*Affine transformtaion of SFE object in histological space*


---

**Description**

These functions perform affine transformations on SFE objects, including all geometries and images. The transformation is performed on each sample as a whole. This differs from functions such as [mirrorImg](#) in that [mirrorImg](#) and [rotateImg](#) transform the image with the center at the center of the image itself. In contrast, the center of transformation here is the center of the bounding box of the entire sample, including images.

**Usage**

```
transpose(sfe, sample_id = "all", maxcell = NULL, filename = "")
```

```
mirror(
  sfe,
  sample_id = "all",
  direction = c("vertical", "horizontal"),
  maxcell = NULL,
  filename = ""
)
```

```
rotate(sfe, sample_id = "all", degrees, maxcell = 1e+07)
```

```
translate(sfe, sample_id = "all", v)
```

```
scale(sfe, sample_id = "all", factor)
```

```
affine(sfe, sample_id = "all", M, v, maxcell = 1e+07)
```

**Arguments**

sfe	An SFE object.
sample_id	Sample(s) to transform.
maxcell	Rotating SpatRasterImage will convert it into ExtImage, loading the image into memory. This argument specifies the maximum number of pixels in the image loaded into memory. The image will be down sampled to approximately this number of pixels.
filename	character. Output filename
direction	character. Should (partially) match "vertical" to flip by rows, or "horizontal" to flip by columns
degrees	How many degrees to rotate. Positive number means clockwise and negative number means counterclockwise.
v	Vector to spatially translate the SFE object.
factor	Numeric, scaling factor.
M	A 2x2 numeric matrix for the linear transformation in the xy plane.

**Details**

For images that are not loaded into memory, rotateImg will load SpatRasterImage into memory and all image operations except translate will load BioFormatsImage into memory.

**Value**

An SFE object with the sample(s) transformed.

**Examples**

```
library(SFEData)
sfe <- McKellarMuscleData("small")
sfe2 <- SpatialFeatureExperiment::transpose(sfe)
sfe3 <- SpatialFeatureExperiment::mirror(sfe)
```

---

show, SpatialFeatureExperiment-method

*Print method for SpatialFeatureExperiment*

---

**Description**

Printing summaries of colGeometries, rowGeometries, and annotGeometries in addition to what's shown for SpatialExperiment. Geometry names and types are printed.

**Usage**

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

**Arguments**

object	A SpatialFeatureExperiment object.
--------	------------------------------------

**Value**

None (invisible NULL).

**Examples**

```
library(SFEData)
sfe <- McKellarMuscleData(dataset = "small")
sfe # The show method is implicitly called
```

---

SpatialFeatureExperiment

*Constructor of SpatialFeatureExperiment object*

---

**Description**

Create a SpatialFeatureExperiment object.

**Usage**

```
SpatialFeatureExperiment(
  assays,
  colData = DataFrame(),
  rowData = NULL,
  sample_id = "sample01",
  spatialCoordsNames = c("x", "y"),
  spatialCoords = NULL,
  colGeometries = NULL,
  rowGeometries = NULL,
  annotGeometries = NULL,
  spotDiameter = NA_real_,
  annotGeometryType = "POLYGON",
  spatialGraphs = NULL,
  unit = c("full_res_image_pixel", "micron"),
  ...
)
```

**Arguments**

- |         |   |
|---------|---|
| assays  | A list or SimpleList of matrix-like elements, or a matrix-like object (e.g. an ordinary matrix, a data frame, a <a href="#">DataFrame</a> object from the <b>S4Vectors</b> package, a <a href="#">SparseMatrix</a> derivative from the <b>SparseArray</b> package, a <a href="#">sparseMatrix</a> derivative from the <b>Matrix</b> package, a <a href="#">DelayedMatrix</a> object from the <b>DelayedArray</b> package, etc...). All elements of the list must have the same dimensions, and dimension names (if present) must be consistent across elements and with the row names of rowRanges and colData. |
| colData | An optional <a href="#">DataFrame</a> describing the samples. Row names on colData, if present, become the column names of the returned object.   |
| rowData | NULL (the default) or a <a href="#">DataFrame</a> object describing the rows. Row names, if present, become the row names of the constructed SummarizedExperiment object. The number of rows of the <a href="#">DataFrame</a> must equal the number of rows of the matrices in assays.  |

sample_id	A character sample identifier, which matches the sample_id in <code>imgData</code> . The sample_id will also be stored in a new column in <code>colData</code> , if not already present. Default = sample01.
spatialCoordsNames	A character vector of column names if <code>*Geometries</code> arguments have ordinary data frames, to identify the columns in the ordinary data frames that specify the spatial coordinates. If <code>colGeometries</code> is not specified, then this argument will behave as in <code>SpatialExperiment</code> , but <code>colGeometries</code> will be given precedence if provided.
spatialCoords	A numeric matrix containing columns of spatial coordinates, as in <code>SpatialExperiment</code> . The coordinates are centroids of the entities represented by the columns of the gene count matrix. If <code>colGeometries</code> is also specified, then it will be given priority and a warning is issued. Otherwise, the sf representation of the centroids will be stored in the colGeometry called centroids.
colGeometries	<p>Geometry of the entities that correspond to the columns of the gene count matrix, such as cells and Visium spots. It must be a named list of one of the following:</p> <p><b>An sf data frame</b> The geometry column specifies the geometry of the entities.</p> <p><b>An ordinary data frame specifying centroids</b> Column names for the coordinates are specified in the <code>spatialCoordsNames</code> argument. For Visium and ST, in addition to the centroid coordinate data frame, the spot diameter in the same unit as the coordinates can be specified in the <code>spotDiameter</code> argument.</p> <p><b>An ordinary data frame specifying polygons</b> Also use <code>spatialCoordsNames</code>. There should an additional column "ID" to specify which vertices belong to which polygon. The coordinates should not be in list columns. Rather, the data frame should look like it is passed to <code>ggplot2::geom_polygon</code>. If there are holes, then there must also be a column "subID" that differentiates between the outer polygon and the holes.</p> <p>In all cases, the data frame should specify the same number of geometries as the number of columns in the gene count matrix. If the column "barcode" is present, then it will be matched to column names of the gene count matrix. Otherwise, the geometries are assumed to be in the same order as columns in the gene count matrix. If the geometries are specified in an ordinary data frame, then it will be converted into sf internally. Named list of data frames because each entity can have multiple geometries, such as whole cell and nuclei segmentations. The geometries are assumed to be POINTs for centroids and POLYGONs for segmentations. If polygons are specified in an ordinary data frame, then anything with fewer than 3 vertices will be removed. For anything other than POINTs, attributes of the geometry will be ignored.</p>
rowGeometries	Geometry associated with genes or features, which correspond to rows of the gene count matrix.
annotGeometries	Geometry of entities that do not correspond to columns or rows of the gene count matrix, such as tissue boundary and pathologist annotations of histological regions, and nuclei segmentation in a Visium dataset. Also a named list as in <code>colGeometries</code> . The ordinary data frame may specify POINTs, POLYGONs, or LINESTRINGs, or their MULTI versions. Each data frame can only specify one type of geometry. For MULTI versions, there must be a column "group" to identify each MULTI geometry.
spotDiameter	Spot diameter for technologies with arrays of spots of fixed diameter per slide, such as Visium, ST, DBiT-seq, and slide-seq. The diameter must be in the same

	unit as the coordinates in the *Geometry arguments. Ignored for geometries that are not POINT or MULTIPOINT.
annotGeometryType	Character vector specifying geometry type of each element of the list if annotGeometry is specified. Each element of the vector must be one of POINT, LINESTRING, POLYGON, MULTIPOINT, MULTILINESTRING, and MULTIPOLYGON. Must be either length 1 (same for all elements of the list) or the same length as the list. Ignored if the corresponding element is an sf object.
spatialGraphs	A named list of listw objects (see spdep) for spatial neighborhood graphs.
unit	Unit the coordinates are in, either microns or pixels in full resolution image.
...	Additional arguments passed to the <a href="#">SpatialExperiment</a> and <a href="#">SingleCellExperiment</a> constructors.

### Value

A SFE object. If neither colGeometries nor spotDiameter is specified, then a colGeometry called "centroids" will be made, which is essentially the spatial coordinates as sf POINTs. If spotDiameter is specified, but not colGeometries, then the spatial coordinates will be buffered by half the diameter to get spots with the desired diameter, and the resulting colGeometry will be called "spotPoly", for which there's a convenience getter and setter, [spotPoly](#).

### Examples

```
library(Matrix)
data("visium_row_col")
coords1 <- visium_row_col[visium_row_col$col < 6 & visium_row_col$row < 6, ]
coords1$row <- coords1$row * sqrt(3)
cg <- df2sf(coords1[, c("col", "row")], c("col", "row"), spotDiameter = 0.7)

set.seed(29)
col_inds <- sample(seq_len(13), 13)
row_inds <- sample(seq_len(5), 13, replace = TRUE)
values <- sample(seq_len(5), 13, replace = TRUE)
mat <- sparseMatrix(i = row_inds, j = col_inds, x = values)
colnames(mat) <- coords1$barcode
rownames(mat) <- sample(LETTERS, 5)
rownames(cg) <- colnames(mat)

sfe <- SpatialFeatureExperiment(list(counts = mat),
  colData = coords1,
  spatialCoordsNames = c("col", "row"),
  spotDiameter = 0.7
)
sfe2 <- SpatialFeatureExperiment(list(counts = mat),
  colGeometries = list(foo = cg)
)
```

---

SpatialFeatureExperiment-class

*The SpatialFeatureExperiment class*

---

**Description**

This class inherits from the [SpatialExperiment](#) (SPE) class, which in turn inherits from [SingleCellExperiment](#) (SCE). `SpatialFeatureExperiment` stores geometries of spots or cells in `sf` objects which form columns of a `DataFrame` which is in turn a column of the `int_colData` `DataFrame` of the underlying SCE object, just like `reducedDim` in SCE. Geometries of the tissue outline, pathologist annotations, and objects (e.g. nuclei segmentation in a Visium dataset) are stored in `sf` objects in a named list called `annotGeometries` in `int_metadata`.

---

SpatialFeatureExperiment-coercion

*SpatialFeatureExperiment coercion methods*

---

**Description**

The `SpatialFeatureExperiment` class inherits from `SpatialExperiment`, which in turn inherits from `SingleCellExperiment`. A `SpatialExperiment` object with geometries in `colGeometries` in the `int_colData`, `rowGeometries` in the `int_elementMetadata`, or `annotGeometries` in the `int_metadata` can be directly converted to `SpatialFeatureExperiment` with `as(spe, "SpatialFeatureExperiment")`. A `SpatialExperiment` object without the geometries can also be converted; the coordinates in the `spatialCoords` field will be used to make POINT geometries named "centroids" to add to `colGeometries`. The geometries can also be supplied separately when using `toSpatialFeatureExperiment`. Images are converted to `SpatRaster`.

**Usage**

```
## S4 method for signature 'SpatialExperiment'
toSpatialFeatureExperiment(
  x,
  colGeometries = NULL,
  rowGeometries = NULL,
  annotGeometries = NULL,
  spatialCoordsNames = c("x", "y"),
  annotGeometryType = "POLYGON",
  spatialGraphs = NULL,
  spotDiameter = NA,
  unit = NULL
)

## S4 method for signature 'SingleCellExperiment'
toSpatialFeatureExperiment(
  x,
  sample_id = "sample01",
  spatialCoordsNames = c("x", "y"),
  spatialCoords = NULL,
  colGeometries = NULL,
  rowGeometries = NULL,
  annotGeometries = NULL,
  annotGeometryType = "POLYGON",
  spatialGraphs = NULL,
  spotDiameter = NA,
```

```

scaleFactors = 1,
imageSources = NULL,
image_id = NULL,
loadImage = TRUE,
imgData = NULL,
unit = NULL
)

## S4 method for signature 'Seurat'
toSpatialFeatureExperiment(
  x,
  add_molecules = TRUE,
  flip = c("geometry", "image", "none"),
  image_scalefactors = c("lowres", "hires"),
  unit = NULL,
  BPPARAM = SerialParam()
)

```

### Arguments

- x** A SpatialExperiment or Seurat object to be coerced to a SpatialFeatureExperiment object.
- colGeometries** Geometry of the entities that correspond to the columns of the gene count matrix, such as cells and Visium spots. It must be a named list of one of the following:
- An sf data frame** The geometry column specifies the geometry of the entities.
  - An ordinary data frame specifying centroids** Column names for the coordinates are specified in the spatialCoordsNames argument. For Visium and ST, in addition to the centroid coordinate data frame, the spot diameter in the same unit as the coordinates can be specified in the spotDiameter argument.
  - An ordinary data frame specifying polygons** Also use spatialCoordsNames. There should be an additional column "ID" to specify which vertices belong to which polygon. The coordinates should not be in list columns. Rather, the data frame should look like it is passed to `ggplot2::geom_polygon`. If there are holes, then there must also be a column "subID" that differentiates between the outer polygon and the holes.
- In all cases, the data frame should specify the same number of geometries as the number of columns in the gene count matrix. If the column "barcode" is present, then it will be matched to column names of the gene count matrix. Otherwise, the geometries are assumed to be in the same order as columns in the gene count matrix. If the geometries are specified in an ordinary data frame, then it will be converted into sf internally. Named list of data frames because each entity can have multiple geometries, such as whole cell and nuclei segmentations. The geometries are assumed to be POINTs for centroids and POLYGONs for segmentations. If polygons are specified in an ordinary data frame, then anything with fewer than 3 vertices will be removed. For anything other than POINTs, attributes of the geometry will be ignored.
- rowGeometries** Geometry associated with genes or features, which correspond to rows of the gene count matrix.
- annotGeometries** Geometry of entities that do not correspond to columns or rows of the gene count matrix, such as tissue boundary and pathologist annotations of histological

regions, and nuclei segmentation in a Visium dataset. Also a named list as in `colGeometries`. The ordinary data frame may specify POINTs, POLYGONs, or LINESTRINGs, or their MULTI versions. Each data frame can only specify one type of geometry. For MULTI versions, there must be a column "group" to identify each MULTI geometry.

<code>spatialCoordsNames</code>	A character vector of column names if <code>*Geometries</code> arguments have ordinary data frames, to identify the columns in the ordinary data frames that specify the spatial coordinates. If <code>colGeometries</code> is not specified, then this argument will behave as in <a href="#">SpatialExperiment</a> , but <code>colGeometries</code> will be given precedence if provided.
<code>annotGeometryType</code>	Character vector specifying geometry type of each element of the list if <code>annotGeometry</code> is specified. Each element of the vector must be one of POINT, LINESTRING, POLYGON, MULTIPOINT, MULTILINESTRING, and MULTIPOLYGON. Must be either length 1 (same for all elements of the list) or the same length as the list. Ignored if the corresponding element is an <code>sf</code> object.
<code>spatialGraphs</code>	A named list of <code>listw</code> objects (see <code>spdep</code> ) for spatial neighborhood graphs.
<code>spotDiameter</code>	Spot diameter for technologies with arrays of spots of fixed diameter per slide, such as Visium, ST, DBiT-seq, and slide-seq. The diameter must be in the same unit as the coordinates in the <code>*Geometry</code> arguments. Ignored for geometries that are not POINT or MULTIPOINT.
<code>unit</code>	# Default unit is "micron". However for Visium one can choose between "micron" or "full_res_image_pixel".
<code>sample_id</code>	A character sample identifier, which matches the <code>sample_id</code> in <code>imgData</code> . The <code>sample_id</code> will also be stored in a new column in <code>colData</code> , if not already present. Default = <code>sample01</code> .
<code>spatialCoords</code>	A numeric matrix containing columns of spatial coordinates, as in <a href="#">SpatialExperiment</a> . The coordinates are centroids of the entities represented by the columns of the gene count matrix. If <code>colGeometries</code> is also specified, then it will be given priority and a warning is issued. Otherwise, the <code>sf</code> representation of the centroids will be stored in the <code>colGeometry</code> called <code>centroids</code> .
<code>scaleFactors</code>	Optional scale factors associated with the image(s). This can be provided as a numeric value, numeric vector, list, or file path to a JSON file for the 10x Genomics Visium platform. For 10x Genomics Visium, the correct scale factor will automatically be selected depending on the resolution of the image from <code>imageSources</code> . Default = 1.
<code>imageSources</code>	Optional file path(s) or URL(s) for one or more image sources.
<code>image_id</code>	Optional character vector (same length as <code>imageSources</code> ) containing unique image identifiers.
<code>loadImage</code>	Logical indicating whether to load image into memory. Default = FALSE.
<code>imgData</code>	Optional <code>DataFrame</code> containing the image data. Alternatively, this can be built from the arguments <code>imageSources</code> and <code>image_id</code> (see Details).
<code>add_molecules</code>	Logical, whether to add transcripts coordinates to an object.
<code>flip</code>	To flip the image, geometry coordinates, or none. Because the image has the origin at the top left while the geometry has origin at the bottom left, one of them needs to be flipped for them to match. If one of them is already flipped, then use "none". The image will not be flipped if it's GeoTIFF.

image\_scalefactors  
 # A character, choose between "lowres" or " hires". Only for 10X Visium, image scaling factors are from 'scalefactors\_json.json' file.

BPPARAM  
 Deprecated when coercing from SpatialExperiment, but is used when coercing from Seurat object.

### Value

A SpatialFeatureExperiment object

### Examples

```
library(VisiumIO)
# From examples of TENxVisium()
sample_dir <- system.file(
  file.path("extdata", "10xVisium", "section1"),
  package = "VisiumIO"
)
## using spacerangerOut folder
tv <- TENxVisium(
  spacerangerOut = file.path(sample_dir, "outs"), processing = "raw", images = "lowres"
)
spe <- import(tv)
# There can't be duplicate barcodes
colnames(spe) <- make.unique(colnames(spe), sep = "-")
rownames(spatialCoords(spe)) <- colnames(spe)
sfe <- toSpatialFeatureExperiment(spe)
# For coercing Seurat to SFE see this -> ./vignettes/seurat_sfe_coerce.Rmd
```

---

SpatialFeatureExperiment-subset

*Subsetting SpatialFeatureExperiment objects*

---

### Description

The SFE method has special treatment for the spatial graphs. In `listw`, the neighbors are indicated by indices, which will change after subsetting. The `SFE_graph_subset` option determines whether the graphs are subsetting or reconstructed. In the default (`options(SFE_graph_subset = TRUE)`), the graphs are subsetting, in which case singletons may be produced. For `options(SFE_graph_subset = FALSE)`, which is the behavior of versions earlier than Bioc 3.20, the graphs are reconstructed with the parameters recorded in an attribute of the graphs. This option can result into different graphs. For example, suppose we start with a `k` nearest neighbor graph. After subsetting, cells at the boundary of the region used to subset the SFE object may lose some of their neighbors. In contrast, when the graph is reconstructed, these same edge cells will gain other cells that remain after subsetting as neighbors in the new KNN graph.

### Usage

```
## S4 method for signature 'SpatialFeatureExperiment,ANY,ANY,ANY'
x[i, j, ..., drop = FALSE]
```

**Arguments**

x	A SpatialFeatureExperiment object.
i	Row indices for subsetting.
j	column indices for subsetting.
...	Passed to the SingleCellExperiment method of [].
drop	Only used if graphs are reconstructed (options(SFE_graph_subset = FALSE)). If TRUE then colGraphs are dropped but annotGraphs are kept.

**Details**

The option SFE\_graph\_subset was introduced because subsetting is usually faster than reconstructing and in some cases such as distance-based neighbors and Visium spot adjacency give the same results. It was introduced also because of the development of alabaster.sfe for a language-agnostic on-disk serialization of SFE objects and some parameters used to construct graphs have special classes whose alabaster methods have not been implemented, such as BPPARAM and BNPARAM, so when reconstructing, the defaults for those arguments will be used.

The edge weights will be recomputed from the binary neighborhood indicator with the same normalization style as the original graph, such as "W" for row normalization. When distance-based edge weights are used instead of the binary indicator, the edge weights will be re-normalized, which is mostly some rescaling. This should give the same results as recomputing the distance based edge weights for styles "raw", "W", and "B" since the distances themselves don't change, but the effects of other more complicated styles of re-normalization on spatial statistics should be further investigated.

By default, upon subsetting, the images are cropped to the bounding box of the remaining cells. However, when the image is large and the bounding box contains most of the original image, cropping is slow. Cropping can be disabled by options(SFE\_subset\_crop = FALSE). Also, when the remaining part of the image is larger than a threshold, the image will not be cropped; the threshold can be set with the SFE\_subset\_crop\_max option, such as options(SFE\_subset\_crop\_max = "100MB").

**Value**

A subsetted SpatialFeatureExperiment object.

**Examples**

```
# Just like subsetting matrices and SingleCellExperiment
library(SFEData)
sfe <- McKellarMuscleData(dataset = "small")
sfe_subset <- sfe[seq_len(10), seq_len(10), drop = TRUE]
# Gives warning as graph reconstruction fails

sfe_subset <- sfe[seq_len(10), seq_len(10)]
```

spatialGraphs

*Spatial graph methods***Description**

Spatial neighborhood graphs as `spdep`'s `listw` objects are stored in the `int_metadata` of the SFE object. The `listw` class is used because `spdep` has many useful methods that rely on the neighborhood graph as `listw`. See the [spdep documentation website](#) for functions to edit the spatial neighborhood graph, or the `nb` object within the `listw`.

**Usage**

```
## S4 method for signature 'SpatialFeatureExperiment'
spatialGraphs(x, MARGIN = NULL, sample_id = "all", name = "all")

colGraphs(x, sample_id = "all", name = "all")

rowGraphs(x, sample_id = "all", name = "all")

annotGraphs(x, sample_id = "all", name = "all")

## S4 replacement method for signature 'SpatialFeatureExperiment'
spatialGraphs(x, MARGIN = NULL, sample_id = "all", name = "all") <- value

colGraphs(x, sample_id = "all", name = "all") <- value

rowGraphs(x, sample_id = "all", name = "all") <- value

annotGraphs(x, sample_id = "all", name = "all") <- value

## S4 method for signature 'SpatialFeatureExperiment,numeric'
spatialGraphNames(x, MARGIN, sample_id = 1L)

## S4 replacement method for signature 'SpatialFeatureExperiment,numeric,ANY,character'
spatialGraphNames(x, MARGIN, sample_id = 1L) <- value

colGraphNames(x, sample_id = 1L)

rowGraphNames(x, sample_id = 1L)

annotGraphNames(x, sample_id = 1L)

colGraphNames(x, sample_id = 1L) <- value

rowGraphNames(x, sample_id = 1L) <- value

annotGraphNames(x, sample_id = 1L) <- value

## S4 method for signature 'SpatialFeatureExperiment'
spatialGraph(x, type = 1L, MARGIN, sample_id = 1L)
```

```

colGraph(x, type = 1L, sample_id = 1L)

rowGraph(x, type = 1L, sample_id = 1L)

annotGraph(x, type = 1L, sample_id = 1L)

## S4 replacement method for signature 'SpatialFeatureExperiment'
spatialGraph(x, type = 1L, MARGIN, sample_id = NULL) <- value

colGraph(x, type = 1L, sample_id = 1L) <- value

rowGraph(x, type = 1L, sample_id = 1L) <- value

annotGraph(x, type = 1L, sample_id = 1L) <- value

```

### Arguments

x	A <code>SpatialFeatureExperiment</code> object.
MARGIN	As in <code>apply</code> . 1 stands for rows and 2 stands for columns. In addition, 3 stands for spatial neighborhood graphs that correspond to <code>annotGeometries</code> .
sample_id	Name of the sample the graph is associated with. This is useful when multiple pieces of tissues are in the same SFE object (say for a joint dimension reduction and clustering) and the spatial neighborhood is only meaningful within the same piece of tissue. See the <code>sample_id</code> argument in <code>SpatialExperiment</code> .
name	Name of the graphs to add to each <code>sample_id</code> ; used in the <code>spatialGraphs</code> replacement method as it must be character while <code>type</code> can be either an integer index or a name.
value	A <code>listw</code> object ( <code>*Graph</code> ), or a named list of list of <code>listw</code> objects ( <code>*Graphs</code> ) where the names of the top level list are <code>sample_ids</code> when adding graphs for all samples in the margin of interest, or a list of <code>listw</code> objects when adding graphs for one sample in one margin.
type	An integer specifying the index or string specifying the name of the <code>*Graph</code> to query or replace. If missing, then the first item in the <code>*Graph</code> will be returned or replaced.

### Value

Getters for multiple graphs return a named list. Getters for names return a character vector of the names. Getters for single graphs return a `listw` object. Setters return an SFE object.

### Examples

```

library(SFEData)
sfe <- McKellarMuscleData(dataset = "small")
g1 <- findVisiumGraph(sfe)
g2 <- findSpatialNeighbors(sfe)

# Set all graphs of a margin by a named list
spatialGraphs(sfe, MARGIN = 2L, sample_id = "Vis5A") <-
  list(tri2nb = g2, visium = g1)
# Or equivalently
colGraphs(sfe, sample_id = "Vis5A") <- list(tri2nb = g2, visium = g1)

```

```

# Get all graphs of a margin, returning a named list
gs <- spatialGraphs(sfe, MARGIN = 2L)
# Or equivalently
gs <- colGraphs(sfe)

# Set graph of the same name and same margin for multiple samples
# Each sample has a separate graph
sfe2 <- McKellarMuscleData("small2")
sfe_combined <- cbind(sfe, sfe2)
colGraphs(sfe_combined, name = "visium", sample_id = "all") <-
  findVisiumGraph(sfe_combined, sample_id = "all")

# Get graph names
spatialGraphNames(sfe, MARGIN = 2L, sample_id = "Vis5A")
# Or equivalently (sample_id optional as only one sample is present)
colGraphNames(sfe)

# Set graph names
spatialGraphNames(sfe, MARGIN = 2L) <- c("foo", "bar")
colGraphNames(sfe) <- c("tri2nb", "visium")

# MARGIN = 1 means rowGraphs; MARGIN = 3 means annotation graphs (annotGraphs)
# for both getters and setters

# Set single graph by
# Spatial graph for myofibers
g_myofiber <- findSpatialNeighbors(sfe,
  type = "myofiber_simplified",
  MARGIN = 3L
)
spatialGraph(sfe, type = "myofiber", MARGIN = 3L) <- g_myofiber
# Or equivalently
annotGraph(sfe, "myofiber") <- g_myofiber

# Get a specific graph by name
g <- spatialGraph(sfe, "myofiber", MARGIN = 3L)
g2 <- spatialGraph(sfe, "visium", MARGIN = 2L)
# Or equivalently
g <- annotGraph(sfe, "myofiber")
g2 <- colGraph(sfe, "visium")

```

---

SpatRasterImage

*SpatRaster representation of images in SFE objects*


---

## Description

SpatialFeatureExperiment and the Voyager package work with images differently from SpatialExperiment. In SFE and Voyager's, plotting functions for SFE objects, the images can be read with `rast` and represented as `SpatRaster`, so the image is not entirely loaded into memory unless necessary. Plotting will not load a large image into memory; rather the image will be downsampled and the downsampled version is plotted. A `SpatRasterImage` object (as of Bioc 3.19 or SFE version 1.6 and above) is a `SpatRaster` object but also inheriting from `VirtualSpatialImage` as required by `SpatialExperiment`.

**Usage**

```
SpatRasterImage(img)

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

**Arguments**

```
img          A SpatRaster or PackedSpatRaster object.
object       A SpatRasterImage object.
```

**Value**

A SpatRasterImage object.

**Examples**

```
# Example code
```

---

<code>splitByCol</code>	<i>Split SFE object with categorical vector or geometry</i>
-------------------------	---

---

**Description**

The split methods for SFE split an SFE object into multiple SFE objects by geometries (all cells/spots intersecting with each geometry will become a separate SFE object). The `splitSamples` function splits the SFE object by `sample_id` so each sample will become a separate SFE object. The `splitContiguity` function splits the SFE object by contiguity of an `annotGeometry`, which by default is "tissueBoundary". The `splitComponent` function splits the SFE object by graph component, so if there are disconnected components in the graph, then each component will become a new SFE object.

**Usage**

```
## S4 method for signature 'SpatialFeatureExperiment,sf'
splitByCol(x, f, sample_id = "all", colGeometryName = 1L, cover = FALSE)

## S4 method for signature 'SpatialFeatureExperiment,sfc'
splitByCol(x, f, sample_id = 1L, colGeometryName = 1L, cover = FALSE)

## S4 method for signature 'SpatialFeatureExperiment,list'
splitByCol(x, f, sample_id = "all", colGeometryName = 1L, cover = FALSE)

splitSamples(x)

splitContiguity(
  x,
  colGeometryName = 1L,
  annotGeometryName = "tissueBoundary",
  min_area = 0,
  cover = FALSE
)
```

**Arguments**

x	An SFE object
f	It can be a sf data frame or sfc to split by geometry. Each row of the sf data frame or each element in the sfc will correspond to a new SFE object. The sf data frame must have a column sample_id when splitting multiple samples. Can also be a list of sfc whose names correspond to sample_ids to split.
sample_id	Which samples to split.
colGeometryName	Which colGeometry to use to determine which cells or spots should belong to which new SFE object when splitting by sf or sfc. Default to the first one.
cover	Logical, whether the geometries in x must be entirely covered by y if op = st_intersection or whether x must be entirely outside y if op = st_difference. Only relevant when keep_whole != "none".
annotGeometryName	Name of annotGeometry to use to split by contiguity.
min_area	Minimum area in the same unit as the geometry coordinates (squared) for each piece to be considered a separate piece when splitting by contiguity. Only pieces that are large enough are considered.

**Value**

A list of SFE objects.

**Examples**

```
# example code
```

---

splitComponent	<i>Split by graph component</i>
----------------	---------------------------------

---

**Description**

Split either a SFE object or sf or sfc by graph component of a spatial neighborhood graph.

**Usage**

```
## S4 method for signature 'SpatialFeatureExperiment'
splitComponent(x, colGraphName = 1L, min_cells = 100)
```

```
## S4 method for signature 'sf'
splitComponent(
  x,
  min_cells = 100,
  distance_cutoff = 50,
  BNPARAM = NULL,
  BPPARAM = SerialParam()
)
```

```
## S4 method for signature 'sfc'
```

```

splitComponent(
  x,
  min_cells = 100,
  distance_cutoff = 50,
  BNPARAM = NULL,
  BPPARAM = SerialParam()
)

```

### Arguments

x	A SFE object, or sf or sfc.
colGraphName	Name of graph to use for splitComponent. We recommend distance based neighbors ('dnearest' in <a href="#">findSpatialNeighbors</a> ), and recommend NOT using k nearest neighbors ('knearest') or triangulation ('tri2nb').
min_cells	Minimum number of cells per graph component; components with fewer than this number of cells are considered debris and removed.
distance_cutoff	For the sf and sfc methods, the distance cutoff used to construct a distance-based spatial neighborhood graph, in the same unit as the spatial coordinates. Anything within this distance is considered a neighbor.
BNPARAM	A <a href="#">BiocNeighborParam</a> object specifying the algorithm to find k nearest neighbors and distance based neighbors with nn_method = "bioc". For distance based neighbors, only <a href="#">KmknnParam</a> and <a href="#">VptreeParam</a> are applicable.
BPPARAM	A <a href="#">BiocParallelParam</a> object for multithreading. Only used for k nearest neighbor and distance based neighbor with nn_method = "bioc".

### Value

A list of the same object type as the input, each element for one component.

---

st_any_pred	<i>Simple geometry predicates</i>
-------------	-----------------------------------

---

### Description

Unlike functions in sf like `st_intersects`, this function simply returns a logical vector indicating whether each geometry in x intersects (or returns TRUE from other predicates) anything in y, preferably when y only contains a small number of geometries or is one single MULTI geometry. This is useful when cropping or subsetting an SFE object with a geometry, such as tissue boundary or histological region polygons or a bounding box.

### Usage

```
st_any_pred(x, y, pred, yx = FALSE, sparse = FALSE, ...)
```

```
st_any_intersects(x, y, yx = FALSE, sparse = FALSE)
```

```
st_n_pred(x, y, pred, ...)
```

```
st_n_intersects(x, y)
```

**Arguments**

x	An object of class sf, sfc, or sfg.
y	Another object of class sf, sfc, or sfg.
pred	A geometric binary predicate function, such as <code>st_intersects</code> . It should return an object of class sgbp, for sparse predicates.
yx	Whether to do <code>pred(y, x)</code> instead of <code>pred(x, y)</code> . For symmetric predicates, the results should be the same. When x has a large number of geometries and y has few, <code>pred(y, x)</code> is much faster than <code>pred(x, y)</code> for <code>st_intersects</code> , <code>st_disjoint</code> , and <code>st_is_within_distance</code> .
sparse	If TRUE, returns numeric indices rather than logical vector. Defaults to FALSE for backward compatibility, though the default in <code>st_intersects</code> is TRUE.
...	Arguments passed to pred.

**Value**

For `st_any_*`, a logical vector indicating whether each geometry in x intersects (or other predicates such as is covered by) anything in y or a numeric vector of indices of TRUE when `sparse = TRUE`. Simplified from the sgbp results which indicate which item in y each item in x intersects, which might not always be relevant. For `st_n_*`, an integer vector indicating the number of geometries in y returns TRUE for each geometry in x.

**Examples**

```
library(sf)
pts <- st_sfc(
  st_point(c(.5, .5)), st_point(c(1.5, 1.5)),
  st_point(c(2.5, 2.5))
)
pol <- st_polygon(list(rbind(c(0, 0), c(2, 0), c(2, 2), c(0, 2), c(0, 0))))
st_any_pred(pts, pol, pred = st_disjoint)
st_any_intersects(pts, pol)
st_n_pred(pts, pol, pred = st_disjoint)
st_n_intersects(pts, pol)
```

---

toExtImage

---

*Convert images to ExtImage*


---

**Description**

The ExtImage class is a thin wrapper around the Image class in ExtImage so it inherits from VirtualSpatialImage as required by SpatialExperiment and has extent as used in Voyager's plotting functions. This function converts SpatRasterImage (thin wrapper around the class in terra) and BioFormatsImage into ExtImage for image operations as implemented in the ExtImage package.

**Usage**

```
## S4 method for signature 'BioFormatsImage'
toExtImage(x, resolution = 4L, channel = NULL)

## S4 method for signature 'SpatRasterImage'
toExtImage(x, maxcell = 1e+07, channel = NULL)
```

**Arguments**

x	Either a BioFormatsImage or SpatRasterImage object.
resolution	Integer, which resolution in the BioFormatsImage to read and convert. Defaults to 4, which is a lower resolution. Ignored if only 1 resolution is present.
channel	Integer vector to indicate channel(s) to read. If NULL, then all channels will be read.
maxcell	Maximum number of pixels when SpatRasterImage is read into memory.

**Value**

A ExtImage object. The image is loaded into memory.

**See Also**

toSpatRasterImage

---

toSpatRasterImage	<i>Convert images to SpatRasterImage</i>
-------------------	--

---

**Description**

The resolution specified from the OME-TIFF file will be read into memory and written to disk as a GeoTIFF file that has the extent. The output file will have the same file name as the input file except without the ome in the extension.

**Usage**

```
## S4 method for signature 'ExtImage'
toSpatRasterImage(
  x,
  save_geotiff = TRUE,
  file_out = "img.tiff",
  overwrite = FALSE
)

## S4 method for signature 'BioFormatsImage'
toSpatRasterImage(
  x,
  save_geotiff = TRUE,
  resolution = 4L,
  channel = NULL,
  overwrite = FALSE
)
```

**Arguments**

x	Either a BioFormatsImage or EBImage object.
save_geotiff	Logical, whether to save the image to GeoTIFF file.
file_out	File to save the non-OME TIFF file for SpatRaster.

overwrite	Logical, whether to overwrite existing file of the same name.
resolution	Integer, which resolution in the BioFormatsImage to read and convert. Defaults to 4, which is a lower resolution. Ignored if only 1 resolution is present.
channel	Integer vector to indicate channel(s) to read. If NULL, then all channels will be read.

**Value**

A SpatRasterImage object

**See Also**

toExtImage

---

translateImg	<i>Translate/shift image in space</i>
--------------	---------------------------------------

---

**Description**

This function shifts the spatial extent of the image in the x-y plane.

**Usage**

```
## S4 method for signature 'SpatRasterImage'
translateImg(x, v, ...)
```

```
## S4 method for signature 'BioFormatsImage'
translateImg(x, v, ...)
```

```
## S4 method for signature 'ExtImage'
translateImg(x, v, ...)
```

**Arguments**

x	An object of class *Image as implemented in this package.
v	Numeric vector of length 2 to shift the image in the x-y plane.
...	Ignored. It's there so different methods can all be passed to the same lapply in the method for SFE objects. Some methods have extra arguments.

**Value**

A \*Image object of the same class that has been shifted in space.

**See Also**

Other image methods: [SFE-image](#), [affineImg\(\)](#), [cropImg\(\)](#), [dim,BioFormatsImage-method](#), [dim,ExtImage-method](#), [ext\(\)](#), [imgRaster\(\)](#), [imgSource\(\)](#), [mirrorImg\(\)](#), [rotateImg\(\)](#), [scaleImg\(\)](#), [transposeImg\(\)](#)

---

transposeImg	<i>Transpose images</i>
--------------	-------------------------

---

### Description

Swap rows and columns of images. In effect, this will flip the image around the diagonal running from top left to bottom right.

### Usage

```
## S4 method for signature 'SpatRasterImage'  
transposeImg(x, filename = "", maxcell = NULL, ...)  
  
## S4 method for signature 'BioFormatsImage'  
transposeImg(x, ...)  
  
## S4 method for signature 'ExtImage'  
transposeImg(x, ...)
```

### Arguments

x	An object of class *Image as implemented in this package.
filename	Output file name for transformed SpatRaster.
maxcell	Max number of pixels to load SpatRasterImage into memory. The default 1e7 is chosen because this is the approximate number of pixels in the medium resolution image at resolution = 4L in Xenium OME-TIFF to make different methods of this function consistent.
...	Ignored. It's there so different methods can all be passed to the same lapply in the method for SFE objects. Some methods have extra arguments.

### Value

For SpatRasterImage and ExtImage, object of the same class. For BioFormatsImage, the image of the specified resolution is read into memory and then the ExtImage method is called, returning ExtImage. For the extent: xmin and xmax are switched with ymin and ymax.

### See Also

Other image methods: [SFE-image](#), [affineImg\(\)](#), [cropImg\(\)](#), [dim,BioFormatsImage-method](#), [dim,ExtImage-method](#), [ext\(\)](#), [imgRaster\(\)](#), [imgSource\(\)](#), [mirrorImg\(\)](#), [rotateImg\(\)](#), [scaleImg\(\)](#), [translateImg\(\)](#)

---

```
unit, SpatialFeatureExperiment-method
Get unit of a SpatialFeatureExperiment
```

---

### Description

Length units can be microns or pixels in full resolution image in SFE objects.

### Usage

```
## S4 method for signature 'SpatialFeatureExperiment'
unit(x)
```

### Arguments

x                    A `SpatialFeatureExperiment` object.

### Value

A string for the name of the unit. At present it's merely a string and `udunits` is not used.

### Examples

```
library(SFEData)
sfe <- McKellarMuscleData(dataset = "small")
SpatialFeatureExperiment::unit(sfe)
```

---

```
updateObject                    Update a SpatialFeatureExperiment object
```

---

### Description

Update a [SpatialFeatureExperiment](#) to the latest version of object structure. This is usually called by internal functions.

### Usage

```
## S4 method for signature 'SpatialFeatureExperiment'
updateObject(object, ..., verbose = FALSE)

SFEVersion(object)
```

### Arguments

object                    An old [SpatialFeatureExperiment](#) object.

...                        Additional arguments that are ignored.

verbose                   Logical scalar indicating whether a message should be emitted as the object is updated.

**Details**

Version 1.1.4 adds package version to the SFE object. We are considering an overhaul of the `spatialGraphs` slot in a future version using the `sfdep` package to decouple the adjacency graph from the edge weights.

**Value**

An updated version of object.

**See Also**

[objectVersion](#), which is used to determine if the object is up-to-date.

**Examples**

```
library(SFEData)
sfe <- McKellarMuscleData("small")
# First version of SFE object doesn't log SFE package version, so should be NULL
SFEVersion(sfe)
sfe <- updateObject(sfe)
# See current version
SFEVersion(sfe)
```

---

`visium_row_col`*Row and columns of Visium barcodes on the slide*

---

**Description**

From Space Ranger 1.3.1.

**Usage**

```
visium_row_col
```

**Format**

A data frame with 4992 rows with columns `barcode`, `col`, and `row`.

**Source**

Space Ranger 1.3.1

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