Package ‘SpatialExperiment’

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Title S4 Class for Spatially Resolved -omics Data
Description Defines an S4 class for storing data from spatial -omics experiments. The class extends SingleCellExperiment to support storage and retrieval of additional information from spot-based and molecule-based platforms, including spatial coordinates, images, and image metadata. A specialized constructor function is included for data from the 10x Genomics Visium platform.

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BugReports https://github.com/drighelli/SpatialExperiment/issues

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Methods for handling image-related data

Description

The set of functions described below is designed to handle the image-related data stored inside a `SpatialExperiment`'s `imgData int_metadata` field. These include:

- `getImg`, `addImg`, `rmvImg` to retrieve/add/remove an image entry to/from the `imgData` DataFrame
- `imgSource`, `imgRaster` to retrieve the path/URL and raster object, respectively, associated with an image or set of images

Usage

```r
## S4 method for signature 'SpatialExperiment'
getImg(x, sample_id = NULL, image_id = NULL)

## S4 method for signature 'SpatialExperiment'
addImg(x, imageSource, scaleFactor, sample_id, image_id, load = TRUE)

## S4 method for signature 'SpatialExperiment'
```
### imgData-methods

```r
rmvImg(x, sample_id = NULL, image_id = NULL)
```

```
## S4 method for signature 'SpatialExperiment'
imgSource(x, sample_id = NULL, image_id = NULL, path = FALSE)
```

```
## S4 method for signature 'SpatialExperiment'
imgRaster(x, sample_id = NULL, image_id = NULL)
```

```
## S4 method for signature 'SpatialExperiment'
rotateImg(x, sample_id = NULL, image_id = NULL, degrees = 90)
```

```
## S4 method for signature 'SpatialExperiment'
mirrorImg(x, sample_id = NULL, image_id = NULL, axis = c("h", "v"))
```

#### Arguments

- `x` : a `SpatialExperiment`
- `sample_id` : character string, TRUE or NULL specifying sample/image identifier(s); here, TRUE is equivalent to all samples/images and NULL specifies the first available entry (see details)
- `image_id` : see `sample_id`
- `imageSource` : a character string specifying an image file name (.png, .jpg or .tif) or URL to source the image from
- `scaleFactor` : single numeric scale factor used to rescale spatial coordinates according to the image’s resolution
- `load` : logical; should the image(s) be loaded into memory as a raster object? if FALSE, will store the path/URL instead
- `path` : logical; for RemoteSpatialImages, TRUE returns the path to the image’s cached file, and FALSE its URL. For Stored/LoadedSpatialImages, a path/NA is returned, irrespective of path.
- `degrees` : single numeric in +/-[0.90,.....,360] specifying how many degrees to rotate. A negative/positive value corresponds to counter-clockwise rotation
- `axis` : character string specifying whether to mirror horizontally ("h") or vertically ("v")

#### Value

- `getImg()` returns a single or list of SpatialImage(s).
- `add/rmvImg()` return a `SpatialExperiment` with modified imgData; specifically, they create/remove an image entry (row) in the imgData DataFrame.
- `imgRaster/Source()` access relevant data in the SpatialImage(s) stored inside the imgData’s data field. Depending on whether or not multiple entries are accessed, a character string or vector is returned by `imgSource()`, and a single or list of raster object(s) is returned by `imgRaster()`.
- `rotate/mirrorImg()` return a LoadedSpatialImage with modified a raster matrix.
Author(s)

Helena L. Crowell

Examples

eexample(read10xVisium)

# 'SpatialImage' accession
(spi <- getImg(spe))
plot(imgRaster(spi))

# remove an image
imgData(spe)
sp <- rmvImg(spe,
  sample_id = "section1",
  image_id = "lowres")
imgData(spe)

# add an image
url <- "https://i.redd.it/3pw5uh7xo041.jpg"
spe <- addImg(spe,
  sample_id = "section1",
  image_id = "pomeranian",
  imageSource = url,
  scaleFactor = NA_real_,
  load = FALSE)

# extract image
img <- imgRaster(spe,
  sample_id = "section1",
  image_id = "pomeranian")
plot(img)

# transformations

# clockwise rotation
spe1 <- rotateImg(spe,
  degrees = 90) # first image

spe2 <- rotateImg(spe,
  sample_id = TRUE,
  image_id = TRUE,
  degrees = 90) # all images

par(mfrow = c(1, 3))
plot(imgRaster(spe))
plot(imgRaster(spe1))
plot(imgRaster(spe2))

# horizontal/vertical mirroring
**read10xVisium**

Load data from a 10x Genomics Visium experiment

---

**Description**

Creates a `SpatialExperiment` from the Space Ranger output directories for 10x Genomics Visium spatial gene expression data.

**Usage**

```r
read10xVisium(
  samples = "",
  sample_id = paste0("sample", sprintf("%02d", seq_along(samples))),
  type = c("HDF5", "sparse"),
  data = c("filtered", "raw"),
  images = "lowres",
  load = TRUE
)
```

**Arguments**

- **samples**: a character vector specifying one or more directories, each corresponding to a 10x Genomics Visium sample (see Details); if provided, names will be used as sample identifiers.
- **sample_id**: character string specifying unique sample identifiers, one for each directory specified via `samples`; ignored if `!is.null(names(samples))`.
- **type**: character string specifying the type of format to read count data from (see `read10xCounts`).
- **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".
- **load**: logical; should the image(s) be loaded into memory as a `grob`? If `FALSE`, will store the path/URL instead.
Details

The constructor assumes data from each sample are located in a single output directory as returned by Space Ranger, thus having the following file organization (where "raw/filtered" refers to either "raw" or "filtered" to match the 'data' argument.) The base directory "outs/" from Space Ranger can either be included manually in the paths provided in 'samples', or can be ignored; if ignored, it will be added automatically. The `.h5` files are used if 'type = "HDF5"'. (Note that 'tissue_positions.csv' was renamed in Space Ranger v2.0.0.)

```r
# sample
  · | — outs
  · · | — raw/filtered_feature_bc_matrix.h5
  · · | — raw/filtered_feature_bc_matrix
  · · · | — barcodes.tsv.gz
  · · · | — features.tsv.gz
  · · · | — matrix.mtx.gz
  · · | — spatial
  · · · | — scalefactors_json.json
  · · · | — tissue_lowres_image.png
  · · · | — tissue_positions.csv
```

Value

A `SpatialExperiment` object

Author(s)

Helena L. Crowell

Examples

```r
dir <- system.file(
  file.path("extdata", "10xVisium"),
  package = "SpatialExperiment")

sample_ids <- c("section1", "section2")
samples <- file.path(dir, sample_ids, "outs")

list.files(samples[1])
list.files(file.path(samples[1], "spatial"))
file.path(samples[1], "raw_feature_bc_matrix")

(spe <- read10xVisium(samples, sample_ids,
  type = "sparse", data = "raw",
  images = "lowres", load = FALSE))

# base directory 'outs/' from Space Ranger can also be omitted
samples2 <- file.path(dir, sample_ids)
(spe2 <- read10xVisium(samples2, sample_ids,
  type = "sparse", data = "raw",
  images = "lowres", load = FALSE))
```
# tabulate number of spots mapped to tissue
cd <- colData(spe)
table(
    in_tissue = cd$in_tissue,
    sample_id = cd$sample_id)

# view available images
imgData(spe)

---

**Description**

Function to read in images and scale factors for 10x Genomics Visium data, and return as a valid `imgData` DataFrame.

**Usage**

```r
readImgData(
  path = ".", 
  sample_id = names(path), 
  imageSources = file.path(path, "tissue_lowres_image.png"), 
  scaleFactors = file.path(path, "scalefactors_json.json"), 
  load = TRUE
)
```

**Arguments**

- `path` : a path where to find one or more images
- `sample_id` : the `sample_id` for the `SpatialExperiment` object
- `imageSources` : the images source path(s)
- `scaleFactors` : the .json file where to find the scale factors
- `load` : logical; should the image(s) be loaded into memory as a `grob`? If FALSE, will store the path/URL instead.

**Value**

- a `DataFrame`

**Author(s)**

Helena L. Crowell
Examples

dir <- system.file(  
  file.path("extdata", "10xVisium", "section1", "outs", "spatial"),  
  package = "SpatialExperiment")

# base directory contains
# - scale factors (scalefactors_json.json)
# - one image (tissue_lowres_image.png)
list.files(dir)

# read in images & scale factors
# as valid 'imgData' 'DFrame'
readImgData(dir, sample_id = "foo")

SpatialExperiment-assays

Methods for named assays

Description

The SpatialExperiment class provides methods for getting or setting named assays. For example, molecules(spe) will get or set an assay named molecules from object spe, equivalent to assay(spe, i = "molecules"). This provides a convenient interface for users and encourages standardization of assay names across packages.

Available methods

In the following code, spe is a SpatialExperiment object, value is a BumpyMatrix-like object with the same dimensions as spe, and ... are further arguments passed to assay (for the getter) or assay<- (for the setter).

molecules(x, ...), molecules(x, ...) <- value: Get or set an assay named molecules, which is usually assumed to be a BumpyMatrix-formatted object containing spatial coordinates (and any other information) of the individual molecules per gene per cell.

Author(s)

Dario Righelli

See Also

assay and assay<-  

Examples

example(SpatialExperiment)
molecules(spe_mol)
The SpatialExperiment class

Description

The SpatialExperiment class is designed to represent spatially resolved transcriptomics (ST) data. It inherits from the SingleCellExperiment class and is used in the same manner. In addition, the class supports storage of spatial information via spatialCoords and storage of images via imgData.

Arguments

... Arguments passed to the SingleCellExperiment constructor to fill the slots of the base class.

sample_id A character sample identifier, which matches the sample_id in imgData. The sample_id will also be stored in a new column in colData, if not already present. Default = sample01.

spatialCoordsNames A character vector of column names from colData containing spatial coordinates, which will be accessible with spatialCoords. Alternatively, the spatialCoords argument may be provided. If both are provided, spatialCoordsNames is given precedence, and a warning is returned. Default = c("x", "y").

spatialCoords A numeric matrix containing columns of spatial coordinates, which will be accessible with spatialCoords. Alternatively, spatialCoordsNames may be provided. If both are provided, spatialCoordsNames is given precedence, and a warning is returned.

scaleFactors 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 imageSources. Default = 1.

imgData Optional DataFrame containing the image data. Alternatively, this can be built from the arguments imageSources and image_id (see Details).

imageSources Optional file path(s) or URL(s) for one or more image sources.

image_id Optional character vector (same length as imageSources) containing unique image identifiers.

loadImage Logical indicating whether to load image into memory. Default = FALSE.

spatialDataNames (Deprecated) A character vector of column names from colData to include in spatialData. Alternatively, the spatialData argument may be provided. If both are provided, spatialDataNames is given precedence, and a warning is returned. (Note: spatialData and spatialDataNames have been deprecated; colData and spatialCoords should be used for all columns. The arguments
have been retained for backward compatibility but may be removed in the future.)

spatialData  (Deprecated) A DataFrame containing columns to store in spatialData, which
must contain at least the columns of spatial coordinates. Alternatively, spatialDataNames
may be provided. If both are provided, spatialDataNames is given precedence,
and a warning is returned. (Note: spatialData and spatialDataNames have
been deprecated; colData and spatialCoords should be used for all columns.
The arguments have been retained for backward compatibility but may be re-
moved in the future.)

Details

In this class, rows represent genes, and columns represent spots (for spot-based ST platforms) or
cells (for molecule-based ST platforms). As for SingleCellExperiment, counts and logcounts
can be stored in the assays slot, and row and column metadata in rowData and colData. For
molecule-based ST data, the additional measurements per molecule per cell can be stored in a
BumpyMatrix-formatted assay named molecules.

The additional arguments in the constructor documented above (e.g. spatialCoords, imgData,
and others) represent the extensions to the SingleCellExperiment class to store associated spatial
and imaging information for ST data.

The constructor expects colData to contain a column named sample_id. If this is not present,
it will assign the value from the sample_id argument. If the imgData argument is provided, the
constructor expects the imgData DataFrame to already be built. Otherwise, it will build it from
the imageSources and (optional) image_id arguments. If image_id is not provided, this will be
assumed from sample_id and imageSources instead. To combine multiple samples within a single
object, see combine.

For 10x Genomics Visium datasets, the function read10xVisium can be used to load data and create
a SpatialExperiment object directly from Space Ranger output files.

Value

A SpatialExperiment object.

Author(s)

Dario Righelli and Helena L. Crowell

See Also

?"SpatialExperiment-methods", which includes: spatialCoords, spatialCoordsNames, imgData,
scaleFactors

?"SpatialExperiment-assays", which includes: molecules

?"SpatialExperiment-colData"

?"SpatialExperiment-combine"

?"SpatialExperiment-subset"

?"SpatialExperiment-misc"
`readImgData`

?"imgData-methods"

`SpatialImage`

`read10xVisium`

### Examples

#### Example 1: Spot-based ST (10x Visium) using constructor

```r
dir <- system.file(
  file.path("extdata", "10xVisium", "section1", "outs"),
  package = "SpatialExperiment")

# read in counts
fnm <- file.path(dir, "raw_feature_bc_matrix")
sce <- DropletUtils::read10xCounts(fnm)

# read in image data
img <- readImgData(
  path = file.path(dir, "spatial"),
  sample_id="foo")

# read in spatial coordinates
fnm <- file.path(dir, "spatial", "tissue_positions_list.csv")
xyz <- read.csv(fnm, header = FALSE,
  col.names = c(
    "barcode", "in_tissue", "array_row", "array_col",
    "pxl_row_in_fullres", "pxl_col_in_fullres"))

# construct observation & feature metadata
rd <- S4Vectors::DataFrame(
  symbol = rowData(sce)$Symbol)

# construct `SpatialExperiment`
(spe <- SpatialExperiment(
  assays = list(counts = assay(sce)),
  rowData = rd,
  colData = DataFrame(xyz),
  spatialCoordsNames = c("pxl_col_in_fullres", "pxl_row_in_fullres"),
  imgData = img,
  sample_id = "foo"))
```

#### Example 2: Spot-based ST (10x Visium) using `read10xVisium`

```
# see ?read10xVisium for details
example(read10xVisium)
```
# Example 3: Molecule-based ST

# create simulated data
n <- 1000; ng <- 50; nc <- 20
# sample xy-coordinates in [0,1]
x <- runif(n)
y <- runif(n)
# assign each molecule to some gene-cell pair
gs <- paste0("gene", seq(ng))
cs <- paste0("cell", seq(nc))
gene <- sample(gs, n, TRUE)
cell <- sample(cs, n, TRUE)
# construct data.frame of molecule coordinates
df <- data.frame(gene, cell, x, y)

# (assure gene & cell are factor so that
# missing observations aren’t dropped)
df$gene <- factor(df$gene, gs)
df$cell <- factor(df$cell, cs)

# construct BumpyMatrix
mol <- BumpyMatrix::splitAsBumpyMatrix(
  df[, c("x", "y")],
  row = df$gene, column = df$cell)

# get count matrix
y <- with(df, table(gene, cell))
y <- as.matrix(unclass(y))

# construct SpatialExperiment
(spe_mol <- SpatialExperiment(
  assays = list(
    counts = y,
    molecules = mol)))

---

SpatialExperiment-coercion

SpatialExperiment coercion methods

Description

The SpatialExperiment class inherits from the SingleCellExperiment class making it necessary to coerce between these classes. To do so, we designed two different methods: the traditional as method and the toSpatialExperiment function (recommended). The as method checks if the SingleCellExperiment object has already populated int_colData with three elements: spatialData, spatialCoords, and imgData. It also checks if colData already contains a sample_id. In case these checks pass the new SpatialExperiment will have the same values as the SingleCellExperiment passed object. Otherwise a SpatialExperiment with default values for these slots will be created.
The `toSpatialExperiment` method expects a `SingleCellExperiment` object and additional arguments as explained in the related section of this documentation. In case the `SingleCellExperiment` object has already populated `int_colData` with `spatialData` and/or `spatialCoords` and/or `imgData`, these will be respectively overwritten in case the arguments `spatialData/spatialDataNames` and/or `spatialCoords/spatialCoordsNames` and/or `imgData` are not NULL.

**Arguments**

- **sce** A `SingleCellExperiment` object.
- **sample_id** A character sample identifier, which matches the `sample_id` in `imgData`. The `sample_id` will also be stored in a new column in `colData`, if not already present. Default = `sample01`.
- **spatialCoordsNames** A character vector of column names from `colData` containing spatial coordinates, which will be accessible with `spatialCoords`. Alternatively, the `spatialCoords` argument may be provided. If both are provided, `spatialCoordsNames` is given precedence, and a warning is returned. Default = `c("x", "y")`.
- **spatialCoords** A numeric matrix containing columns of spatial coordinates, which will be accessible with `spatialCoords`. Alternatively, `spatialCoordsNames` may be provided. If both are provided, `spatialCoordsNames` is given precedence, and a warning is returned.
- **scaleFactors** 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 `imageSources`. Default = `1`.
- **imgData** Optional DataFrame containing the image data. Alternatively, this can be built from the arguments `imageSources` and `image_id` (see Details).
- **imageSources** Optional file path(s) or URL(s) for one or more image sources.
- **image_id** Optional character vector (same length as `imageSources`) containing unique image identifiers.
- **loadImage** Logical indicating whether to load image into memory. Default = `FALSE`.
- **spatialDataNames** (Deprecated) A character vector of column names from `colData` to include in `spatialData`. Alternatively, the `spatialData` argument may be provided. If both are provided, `spatialDataNames` is given precedence, and a warning is returned. (Note: `spatialData` and `spatialDataNames` have been deprecated; `colData` and `spatialCoords` should be used for all columns. The arguments have been retained for backward compatibility but may be removed in the future.)
- **spatialData** (Deprecated) A DataFrame containing columns to store in `spatialData`, which must contain at least the columns of spatial coordinates. Alternatively, `spatialDataNames` may be provided. If both are provided, `spatialDataNames` is given precedence, and a warning is returned. (Note: `spatialData` and `spatialDataNames` have been deprecated; `colData` and `spatialCoords` should be used for all columns. The arguments have been retained for backward compatibility but may be removed in the future.)
Examples

dir <- system.file(
    file.path("extdata", "10xVisium", "section1", "outs"),
    package = "SpatialExperiment")

# read in counts
fnm <- file.path(dir, "raw_feature_bc_matrix")
sce <- DropletUtils::read10xCounts(fnm)

# read in spatial coordinates
fnm <- file.path(dir, "spatial", "tissue_positions_list.csv")
xyz <- read.csv(fnm, header = FALSE,
    col.names = c("barcode", "in_tissue", "array_row", "array_col",
    "pxl_row_in_fullres", "pxl_col_in_fullres"))

# read in image data
img <- readImgData(
    path = file.path(dir, "spatial"),
    sample_id = "sample01")

## as method
(spe <- as(sce, "SpatialExperiment"))

colData(sce) <- DataFrame(xyz[,c(1:4)])
int_colData(sce)$spatialCoords <- as.matrix(xyz[,c(5,6)])

## Coercing an sce without imgData
(spe <- as(sce, "SpatialExperiment"))

## Coercing an sce with imgData
int_colData(sce)$imgData <- img
(spe <- as(sce, "SpatialExperiment"))

## toSpatialExperiment method
colData(sce) <- DataFrame(xyz)
(spe <- toSpatialExperiment(sce,
    imgData = img,
    spatialCoordsNames = c("pxl_col_in_fullres", "pxl_row_in_fullres"),
    sample_id = "sample01"))

---

SpatialExperiment-colData

SpatialExperiment colData

---

Description

The SpatialExperiment class provides a modified colData setter, which ensures that the SpatialExperiment object remains valid.
Usage

```r
## S4 replacement method for signature 'SpatialExperiment,DataFrame'
colData(x) <- value

## S4 replacement method for signature 'SpatialExperiment,`NULL`'
colData(x) <- value
```

Arguments

- **x**
  - a `SpatialExperiment`
- **value**
  - a `DataFrame`

Details

The `colData` setter performs several checks to ensure validity. If the replacement `colData` does not contain a `sample_id` column, the existing `sample_ids` will be retained. If the replacement `colData` contains `sample_ids`, a check is performed to ensure the number of unique `sample_ids` is the same, i.e. a one-to-one mapping is possible. If the replacement is `NULL`, the `sample_ids` are retained. In addition, checks are performed against the `sample_ids` in `imgData`.

Value

- a `SpatialExperiment` object with updated `colData`

Examples

```r
eexample(read10xVisium)

# empty replacement retains sample identifiers
colData(spe) <- NULL
names(colData(spe))

# replacement of sample identifiers
# requires one-to-one mapping

## invalid replacement
tryCatch(
  spe$sample_id <- seq(ncol(spe)),
  error = function(e) message(e))

## valid replacement
old <- c("section1", "section2")
new <- c("sample_A", "sample_B")
idx <- match(spe$sample_id, old)
tmp <- spe

tmp$sample_id <- new[idx]
table(spe$sample_id, tmp$sample_id)
```
Combining SpatialExperiment objects

Description

The SpatialExperiment class provides modified methods to combine multiple SpatialExperiment objects by column, for example from multiple samples. These methods ensure that all data fields remain synchronized when samples are added or removed.

Usage

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

Arguments

- `...`: a list of SpatialExperiment objects
- `deparse.level`: refer to ?rbind

Value

A combined SpatialExperiment object

Combining

The `...` argument is assumed to contain one or more SpatialExperiment objects.

cbind(..., deparse.level=1): Returns a SpatialExperiment where all objects in `...` are combined column-wise, i.e., columns in successive objects are appended to the first object.

Each SpatialExperiment object in `...` must have the same colData (with the same spatialCoords). If multiple objects use the same sample_id, the method will proceed by assigning unique sample_ids.

Additionally, the method combines imgData by row using rbind.

Refer to ?"cbind,SingleCellExperiment-method" for details on how metadata and other inherited attributes are combined in the output object.

Refer to ?cbind for the interpretation of deparse.level.

Author(s)

Dario Righelli
Examples

```r
example(read10xVisium, echo = FALSE)

# merging with duplicated 'sample_id's
# will automatically assign unique identifiers
spe1 <- spe2 <- spe
spe3 <- cbind(spe1, spe2)
unique(spe3$sample_id)

# assign unique sample identifiers
spe1 <- spe2 <- spe
spe1$sample_id <- paste(spe1$sample_id, "sample1", sep = ".")
spe2$sample_id <- paste(spe2$sample_id, "sample2", sep = ".")

# combine into single object
spe <- cbind(spe1, spe2)

# view joint 'imgData'
imgData(spe)

# tabulate number of spots mapped to tissue
cd <- colData(spe)
table(
  in_tissue = cd$in_tissue,
  sample_id = cd$sample_id)
```

Description

The `SpatialExperiment` class provides a family of methods to get and set spatial data attributes in `SpatialExperiment` objects. Spatial attributes include `spatialCoords`, `imgData`, and `scaleFactors`, as well as methods to rotate and mirror `SpatialExperiment` objects and their spatial coordinates.

Usage

```r
## S4 method for signature 'SpatialExperiment'
spatialData(x)

## S4 replacement method for signature 'SpatialExperiment,DFrame'
spatialData(x) <- value

## S4 replacement method for signature 'SpatialExperiment,"NULL"'
spatialData(x) <- value

## S4 method for signature 'SpatialExperiment'
```
spatialDataNames(x)

## S4 replacement method for signature 'SpatialExperiment,character'
spatialDataNames(x) <- value

## S4 replacement method for signature 'SpatialExperiment,"NULL"'
spatialDataNames(x) <- value

## S4 method for signature 'SpatialExperiment'
spatialCoords(x)

## S4 replacement method for signature 'SpatialExperiment,matrix'
spatialCoords(x) <- value

## S4 replacement method for signature 'SpatialExperiment,"NULL"'
spatialCoords(x) <- value

## S4 method for signature 'SpatialExperiment'
spatialCoordsNames(x)

## S4 replacement method for signature 'SpatialExperiment,character'
spatialCoordsNames(x) <- value

## S4 replacement method for signature 'SpatialExperiment,"NULL"'
spatialCoordsNames(x) <- value

## S4 method for signature 'SpatialExperiment'
scaleFactors(x, sample_id = TRUE, image_id = TRUE)

## S4 method for signature 'SpatialExperiment'
x$name

## S4 method for signature 'SpatialExperiment'
imgData(x)

## S4 replacement method for signature 'SpatialExperiment,DataFrame'
imgData(x) <- value

## S4 replacement method for signature 'SpatialExperiment,"NULL"'
imgData(x) <- value

### Arguments

**x**  
A *SpatialExperiment* object.

**value**  
Replacement value for replacement methods.

**sample_id**  
Logical value or character vector specifying sample identifier(s) for `scaleFactors`.  
Default = TRUE (all samples).
### image_id
Logical value or character vector specifying image identifier(s) for scaleFactors. Default = TRUE (all images).

### name
The name of the colData column to extract.

### Details
Additional details for each type of data attribute are provided below.

Note: spatialData and spatialDataNames (previously used to store a subset of columns from colData) have been deprecated. All columns should be stored in either spatialCoords (numeric matrix containing spatial coordinates) or colData (all other columns). The spatialData/spatialDataNames functionality has been retained for backward compatibility but may be removed in the future.

See rotateCoords, mirrorCoords, rotateObject, or mirrorObject for details on methods to rotate and mirror SpatialExperiment objects and their spatialCoords.

### Value
Return value varies depending on method, as described below.

#### spatialData and spatialCoords methods

**spatialData(x) <- value:** The spatialData setter expects a DataFrame. If the input does not contain an in_tissue column, this will be included with a default value of 1.

**spatialCoords(x):** Getter for numeric matrix of spatial coordinates.

**spatialCoords(x) <- value:** Setter for numeric matrix of spatial coordinates.

#### spatialDataNames and spatialCoordsNames methods

**spatialDataNames(x):** Returns the names of the colData associated with the spatial information, which are stored in the int_metadata.

**spatialDataNames(x) <- value:** Setter to replace column names in the spatialData DataFrame.

**spatialCoordsNames(x):** Returns the defined names of the spatial coordinates (e.g. c("x", "y")).

**spatialCoordsNames(x) <- value:** Setter to define the names of the spatial coordinate columns.

#### imgData methods

**imgData(x):** Getter to return the imgData DataFrame.

**imgData(x) <- value:** Setter to provide a DataFrame object as imgData of the SpatialExperiment object.

#### Other methods

**scaleFactors(x, sample_id, image_id):** Getter to return the scale factors associated with the sample_id(s) and image_id(s) provided. This is related to the stored image(s) in the SpatialExperiment imgData structure. See argument descriptions for further details.
Examples

```r
example(read10xVisium)

# spatialCoords returns a numeric matrix
head(spatialCoords(spe))

# change spatial coordinate names
spatialCoordsNames(spe)
spatialCoordsNames(spe) <- c("x", "y")
head(spatialCoords(spe))

# imgData and scale factors
imgData(spe)
scaleFactors(spe)

# tabulate number of spots mapped to tissue
cd <- colData(spe)
table(
  in_tissue = cd$in_tissue,
  sample_id = cd$sample_id)
```

Description

Miscellaneous methods for the `SpatialExperiment` class and its descendants that do not fit into any other documentation category such as, for example, show methods.

Usage

```r
## S4 method for signature 'SpatialExperiment'
show(object)
```

Arguments

- `object`: a `SpatialExperiment` object

Value

Returns NULL

Author(s)

Dario Righelli and Helena L. Crowell
Examples

```
example(read10xVisium)
spe
```

---

**SpatialExperiment-rotate-mirror**

*Methods for spatial attributes*

**Description**

The `SpatialExperiment` class provides methods to rotate and mirror `SpatialExperiment` objects and their `spatialCoords`.

**Usage**

```r
## S4 method for signature 'SpatialExperiment'
rotateCoords(x, sample_id = NULL, degrees = 90, warn = TRUE)

## S4 method for signature 'SpatialExperiment'
mirrorCoords(x, sample_id = NULL, axis = c("h", "v"), warn = TRUE)

## S4 method for signature 'SpatialExperiment'
rotateObject(x, sample_id = NULL, image_id = NULL, degrees = 90)

## S4 method for signature 'SpatialExperiment'
mirrorObject(x, sample_id = NULL, image_id = NULL, axis = c("h", "v"))
```

**Arguments**

- `x` A `SpatialExperiment` object.
- `sample_id` Logical value or character vector specifying sample identifier(s) for scaleFactors. Default = TRUE (all samples).
- `degrees` single numeric in +/-[0,90,...,360] specifying how many degrees to rotate. A negative/positive value corresponds to counter-clockwise rotation. Applicable for `rotateCoords` and `rotateObject` methods.
- `warn` Logical value indicating whether to print a warning about mismatches between coordinates and images, possible with the `spatialCoords` transformation methods `rotateCoords` and `mirrorCoords`.
- `axis` character string specifying whether to mirror horizontally ("h") or vertically ("v"). Applicable for `mirrorCoords` and `mirrorObject` methods.
- `image_id` Logical value or character vector specifying image identifier(s) for scaleFactors. Default = TRUE (all images).

**Details**

Additional details for each type of data attribute are provided below.
Value

Return value varies depending on method, as described below.

**spatialCoords transformation methods**

rotateCoords(x, sample_id, degrees, warn): Apply a rotation to the spatialCoords of x, potentially subsetted to sample sample_id (or without subsetting if sample_id is NULL), by the specified number of degrees clockwise. Warn about mismatches with images if warn.

mirrorCoords(x, sample_id, axis, warn): Reflect the spatialCoords of x across either the horizontal or vertical axis, specified by supplying "h" or "v" to the axis argument, respectively. Subset x to just the sample sample_id, if not NULL. Warn about mismatches with images if warn.

**SpatialExperiment transformation wrapper methods**

rotateObject(x, sample_id, image_id, degrees): Apply a rotation to the spatialCoords and imgData of x, potentially subsetted to sample sample_id (or without subsetting if sample_id is NULL), by the specified number of degrees clockwise. Wrapper around rotateCoords and rotateImg.

mirrorObject(x, sample_id, image_id, axis): Reflect the spatialCoords and imgData of x across either the horizontal or vertical axis, specified by supplying "h" or "v" to the axis argument, respectively. Subset x to just the sample sample_id, if not NULL. Wrapper around mirrorCoords and mirrorImg.

**Author(s)**

Nicholas J. Eagles

**Examples**

```r
example(read10xVisium)

# rotateCoords(), mirrorCoords(), rotateObject(), and mirrorObject() return a
# SpatialExperiment, potentially subsetted by sample.

# Subset to just "section1"; rotate coordinates 90 degrees clockwise followed
# by a reflection across the vertical axis
spe_coords <- rotateCoords(spe, sample_id = "section1", degrees = 90)
spe_coords <- mirrorCoords(spe_coords, axis = "v")

# Subset to just "section2"; transform both the imgData() and spatialCoords()
# by a 180-degree rotation then reflection across the vertical axis
spe_wrapper <- rotateObject(spe, sample_id = "section2", degrees = 180)
spe_wrapper <- mirrorObject(spe_wrapper, axis = "v")
```
Subsetting SpatialExperiment objects

Description

The subsetting method for SpatialExperiment objects ensures that spatial data attributes (spatialCoords and imgData) are subbed correctly to match rows and columns with the remainder of the object.

Arguments

- **x**
  - a SpatialExperiment object
- **i**
  - row indices for subsetting
- **j**
  - column indices for subsetting

Value

- a SpatialExperiment object

subset

[>: subsetting method

Examples

```r
example(read10xVisium)

dim(spe)

set.seed(123)
idx <- sample(ncol(spe), 10)
sub <- spe[, idx]
dim(sub)
colData(sub)
spatialCoords(sub)
```
Constructor

SpatialImage(x, is.url) will return a SpatialImage object. The class of the object depends on the type of x:

- If x is a raster object, a LoadedSpatialImage is returned. This represents an image that is fully realized into memory, where the raster representation is stored inside the output object.
- If x is a string and is.url=TRUE or it starts with "http://", "http://" or "ftp://", a RemoteSpatialImage is returned. This represents an image that is remotely hosted and retrieved only on request.
- If x is a string and is.url=TRUE or it does not start with a URL-like prefix, a StoredSpatialImage is returned. This represents an image that is stored in a local file and is loaded into memory only on request.

Getting the raster image

For a SpatialImage object x, imgRaster(x, ...) will return a raster object (see ?as.raster). This is effectively a matrix of RGB colors for each pixel in the image.

For a StoredSpatialImage object x, additional arguments in ... are passed to image_read. This controls how the image is read into memory.

For a RemoteSpatialImage object x, the image file is first downloaded before the raster is returned. Here, ... may contain an extra cache argument, which should be a BiocFileCache object (from the BiocFileCache package) specifying the file cache location. The default location is determined by options("SpatialExperiment.remote.cache.path"). otherwise it defaults to a subdirectory in the R temporary directory. Any further named arguments in ... are passed to image_read.

as.raster(x, ...) is the same as imgRaster(x, ...).

In-memory caching

For StoredSpatialImage and RemoteSpatialImage objects, loading the image with imgRaster will automatically store the loaded raster object in an in-memory cache. Any subsequent imgRaster call will retrieve the raster from the cache, avoiding costly retrieval from the file system.

The cache policy is to evict the least recently used images when a new image would be added that exceeds the maximum cache size. If the new image by itself exceeds the maximum cache size, all images are evicted from the cache to trigger garbage collection and free up memory.

By default, the maximum size of the cache is 4 GB. This can be modified by setting options("SpatialExperiment.cache.size") to some number of bytes, e.g., 2^32.

Transformations

Two basic image transformations are currently supported for any SpatialImage x, namely, rotateImg(x, degrees) for clockwise (degrees > 0) and counter-clockwise (degrees < 0) rotation, and mirrorImg(x, axis) for horizontal (axis = "h") and vertical (axis = "v") mirroring.

Note that, both rotateImg() and mirrorImg() operate on the raster matrix of the input SpatialImage. Thus, any SpatialImage will automatically be coerced into a LoadedSpatialImage upon rotation/mirroring.
**SpatialImage-class**

**Other methods**

dim(x) will return an integer vector of length 2, containing the width and height of the image in pixels. Note that this calls imgRaster under the hood and thus may interact with the file and memory caches as described above.

For any SpatialImage x, `as(x, "LoadedSpatialImage")` will create a LoadedSpatialImage containing an in-memory raster object.

For a RemoteSpatialImage x, `as(x, "StoredSpatialImage")` will create a StoredSpatialImage pointing to the file cache location.

**Author(s)**

Aaron Lun

**Examples**

```r
path <- system.file(
  "extdata", "10xVisium", "section1", "outs", "spatial",
  "tissue_lowres_image.png", package="SpatialExperiment")

spi <- SpatialImage(path)
plot(imgRaster(spi))

# the following operations all use the cache
# so there is no need to reload the image
nrow(spi)
ncol(spi)
plot(as.raster(spi))

# coercing to an explicitly in-memory raster
spi <- as(spi, "LoadedSpatialImage")
plot(as.raster(spi))

############################
# transformations #
############################

# (counter-)clockwise rotation
spi1 <- rotateImg(spi, degrees = +90)
spi2 <- rotateImg(spi, degrees = -90)

par(mfrow = c(1, 3))
plot(as.raster(spi))
plot(as.raster(spi1))
plot(as.raster(spi2))

# horizontal/vertical mirroring
spi1 <- mirrorImg(spi, axis = "h")
spi2 <- mirrorImg(spi, axis = "v")

par(mfrow = c(1, 3))
plot(as.raster(spi))
```
plot(as.raster(spi1))
plot(as.raster(spi2))

---

SpatialImage-misc  Miscellaneous SpatialImage methods

Description

Miscellaneous methods for the SpatialImage class that do not fit into any other documentation category such as, for example, show methods.

Usage

## S4 method for signature 'VirtualSpatialImage'
show(object)

Arguments

object  a SpatialImage object

Value

one

Author(s)

Helena L. Crowell
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