Package 'cytoviewer'

November 28, 2024

Version 1.6.0

Title An interactive multi-channel image viewer for R

Description This R package supports interactive visualization of multi-channel images and segmentation masks generated by imaging mass cytometry and other highly multiplexed imaging techniques using shiny. The cytoviewer interface is divided into image-level (Composite and Channels) and cell-level visualization (Masks). It allows users to overlay individual images with segmentation masks, integrates well with SingleCellExperiment and SpatialExperiment objects for metadata visualization and supports image downloads.

License GPL-3

Imports shiny, shinydashboard, utils, colourpicker, shinycssloaders, svgPanZoom, viridis, archive, grDevices, RColorBrewer, svglite, EBImage, methods, cytomapper, SingleCellExperiment, S4Vectors, SummarizedExperiment

Suggests BiocStyle, knitr, rmarkdown, markdown, testthat

biocViews ImmunoOncology, Software, SingleCell, OneChannel, TwoChannel, MultiChannel, Spatial, DataImport

VignetteBuilder knitr

URL https://github.com/BodenmillerGroup/cytoviewer

BugReports https://github.com/BodenmillerGroup/cytoviewer/issues

RoxygenNote 7.2.3

Encoding UTF-8

git_url https://git.bioconductor.org/packages/cytoviewer

git_branch RELEASE_3_20

git_last_commit cbda7eb

git_last_commit_date 2024-10-29

Repository Bioconductor 3.20

Date/Publication 2024-11-28

Author Lasse Meyer [aut, cre] (<https://orcid.org/0000-0002-1660-1199>), Nils Eling [aut] (<https://orcid.org/0000-0002-4711-1176>)

Maintainer Lasse Meyer <lasse.meyer@dqbm.uzh.ch>

cytoviewer

5

Contents

| cytoviewer | | | | ••• | | • • | | • | • | | • | | | | • | | · | • | • | • | | • | • | • | • | 2 |
|------------|--|--|--|-----|--|-----|--|---|---|--|---|--|--|--|---|--|---|---|---|---|--|---|---|---|---|---|
|------------|--|--|--|-----|--|-----|--|---|---|--|---|--|--|--|---|--|---|---|---|---|--|---|---|---|---|---|

Index

cytoviewer

cytoviewer - Shiny application to interactively browse multi-channel images

Description

This shiny R application allows users to interactively visualize multi-channel images and segmentation masks generated by imaging mass cytometry and other highly multiplexed imaging techniques. The cytoviewer interface is divided into image-level (Composite and Channels) and cell-level visualization (Masks). It allows users to overlay individual images with segmentation masks, integrates well with SingleCellExperiment and SpatialExperiment objects for metadata visualization and supports image downloads.

Usage

```
cytoviewer(
  image = NULL,
  mask = NULL,
  object = NULL,
  cell_id = NULL,
  img_id = NULL
)
```

Arguments

| image | (optional) a CytoImageList object containing single or multi-channel Image objects. |
|---------|---|
| mask | (optional) a CytoImageList containing single-channel Image objects. |
| object | (optional) a SingleCellExperiment or SpatialExperiment object. |
| cell_id | character specifying the colData(object) entry, in which the integer cell IDs are stored. These IDs should match the integer pixel values in the segmentation mask object (mask). |
| img_id | character specifying the colData(object) and mcols(mask) and/or mcols(image) entry, in which the image IDs are stored. |

Value

A Shiny app object for interactive multi-channel image visualization and exploration

The input objects

The functionality of cytoviewer depends on which input objects are user-provided. Below we describe the four use cases in respect to input objects and functionality.

1. Usage of cytoviewer with images, masks and object

2

cytoviewer

The full functionality of cytoviewer can be leveraged when image, mask and object are provided. This allows image-level visualization (Composite and Channels), cell-level visualization, overlaying images with segmentation masks as well as metadata visualization.

2. Usage of cytoviewer with images only

If only image is specified, image-level visualization (Composite and Channels) is possible.

3. Usage of cytoviewer with images and masks

Image-level visualization (Composite and Channels), overlaying of images with masks and celllevel visualization is feasible when image and mask are provided.

4. Usage of cytoviewer with masks and object

If mask and object are specified, cell-level visualization as well as metadata visualization is possible.

Author(s)

Lasse Meyer (<lasse.meyer@dqbm.uzh.ch>)

See Also

plotPixels for the function underlying image-level visualization

plotCells for the function underlying cell-level visualization

cytomapperShiny for a shiny application that visualizes gated cells on images

Examples

```
# Load example datasets from cytomapper
library(cytomapper, quietly = TRUE)
data("pancreasImages")
data("pancreasMasks")
data("pancreasSCE")
# 1. Use cytoviewer with images, masks and object
app <- cytoviewer(image = pancreasImages,</pre>
                  mask = pancreasMasks,
                  object = pancreasSCE,
                  img_id = "ImageNb",
                  cell_id = "CellNb")
if (interactive()) {
shiny::runApp(app, launch.browser = TRUE)
}
## Other input variations (see "The input objects" section):
# 2. Use cytoviewer with images
app_1 <- cytoviewer(image = pancreasImages)</pre>
if (interactive()) {
shiny::runApp(app_1, launch.browser = TRUE)
}
# 3. Use cytoviewer with images and masks
app_2 <- cytoviewer(image = pancreasImages,</pre>
                  mask = pancreasMasks,
                  img_id = "ImageNb")
```

Index

cytomapperShiny, 3
cytoviewer, 2

plotCells, 3
plotPixels, 3