

Package ‘interactiveDisplayBase’

December 11, 2024

Type Package

Title Base package for enabling powerful shiny web displays of
Bioconductor objects

Version 1.45.0

Date 2021-07-30

Imports shiny, DT

Depends R (>= 2.10), methods, BiocGenerics

Suggests knitr, markdown

Enhances rstudioapi

Description The interactiveDisplayBase package contains the the basic
methods needed to generate interactive Shiny based display
methods for Bioconductor objects.

License Artistic-2.0

Collate interactiveDisplayBase.R dataframe.R dot_runApp.R zzz.R

VignetteBuilder knitr

biocViews GO, GeneExpression, Microarray, Sequencing, Classification,
Network, QualityControl, Visualization, Visualization,
Genetics, DataRepresentation, GUI, AnnotationData, ShinyApps

RoxygenNote 7.1.1

git_url <https://git.bioconductor.org/packages/interactiveDisplayBase>

git_branch devel

git_last_commit 5218b82

git_last_commit_date 2024-10-29

Repository Bioconductor 3.21

Date/Publication 2024-12-10

Author Bioconductor Package Maintainer [cre],
Shawn Balcome [aut],
Marc Carlson [ctb],
Marcel Ramos [ctb]

Maintainer Bioconductor Package Maintainer <maintainer@bioconductor.org>

Contents

.runApp	2
display	3
Index	5

.runApp	<i>Run a shiny app, capturing results to the R session</i>
---------	--

Description

This utility function launches a shiny visualization application, either in the RStudio viewer pane (if run under RStudio) or in the browser.

Usage

```
.runApp(app, ...)
```

Arguments

app	The shiny application definition, see ?shiny::runApp.
...	additional arguments passed to shiny::runApp().

Value

The return value of shiny::runApp.

Author(s)

Martin Morgan

Examples

```
if (interactive()) {
  require(shiny)

  app <- list(
    ui = fluidPage(
      title="Who Am I?",
      sidebarLayout(
        position="left",
        sidebarPanel(
          h1("Your name"),
          textInput("your_name", "Your name?", "Anonymous"),
          actionButton("done", "Done")),
        mainPanel(
          "Hi", textOutput("your_name", inline=TRUE))
      )),
  )
```

```
server = function(input, output) {
  output$your_name <- renderText(input$your_name)
  observe({
    if (input$done > 0)
      isolate(stopApp(returnValue = input$your_name))
  })
})

.runApp(app)
}
```

display

display: Open a Shiny application for a Bioconductor object

Description

This opens a shiny visualization application in the browser based on the submitted object.

Usage

```
display(object, ...)
```

Arguments

object	data object to display
...	additional arguments passed to methods; currently unused.

Value

Usually some variation of the initial input object, but it may be altered by the display widget (subset for example).

Author(s)

Shawn Balcome and Marc Carlson

See Also

<http://bioconductor.org/packages/2.13/bioc/html/interactiveDisplayBase.html>

Examples

```
if(interactive()) {
  ## draw a data.frame
  display(mtcars)

  ## subset a data.frame:
```

4

display

```
mtcars2 <- display(mtcars)
```

```
}
```

Index

* **manip, internal**

.runApp, [2](#)

* **methods**

display, [3](#)

.runApp, [2](#)

display, [3](#)

display, ANY-method (display), [3](#)

display, data.frame-method (display), [3](#)

display, missing-method (display), [3](#)