

**WORKSHOP: R SHINY** 

By: Hiba Ben Aribi

### WORKSHOP

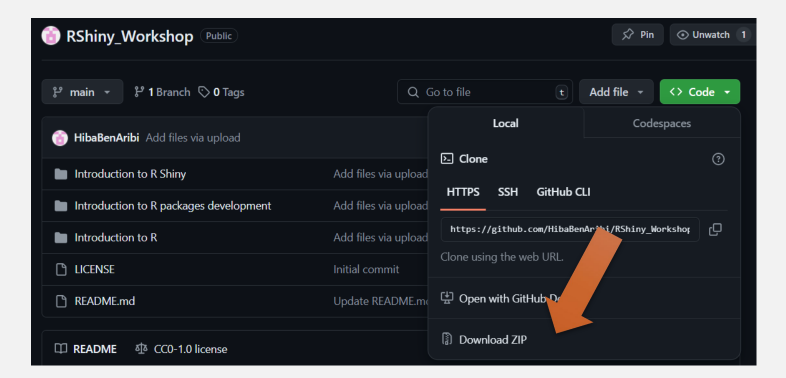


#### GitHub Repository:

https://bit.ly/RShiny\_Workshop

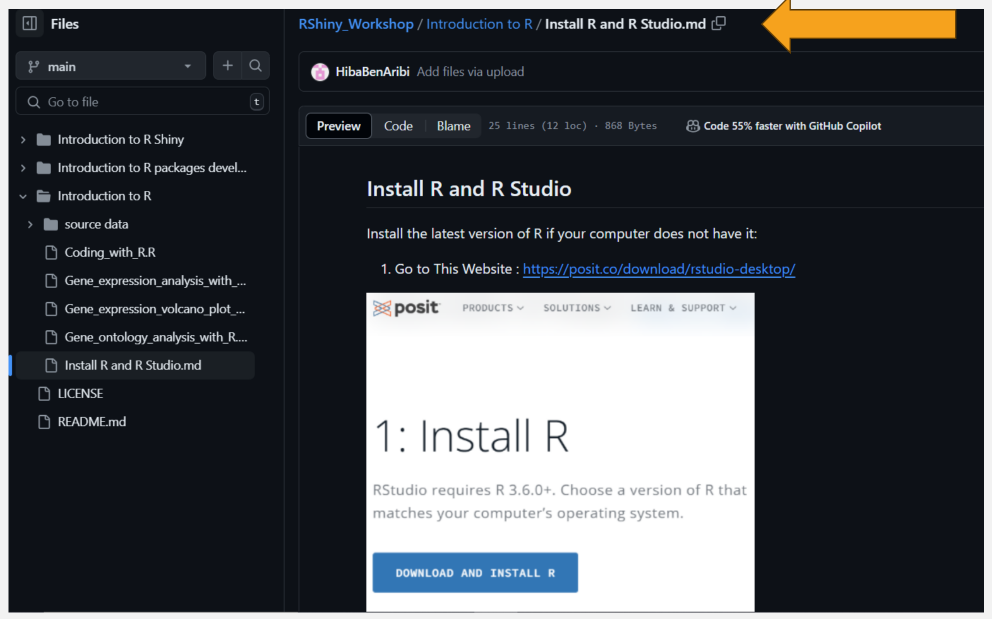
#### Or

https://github.com/HibaBenAribi/RShiny\_Workshop



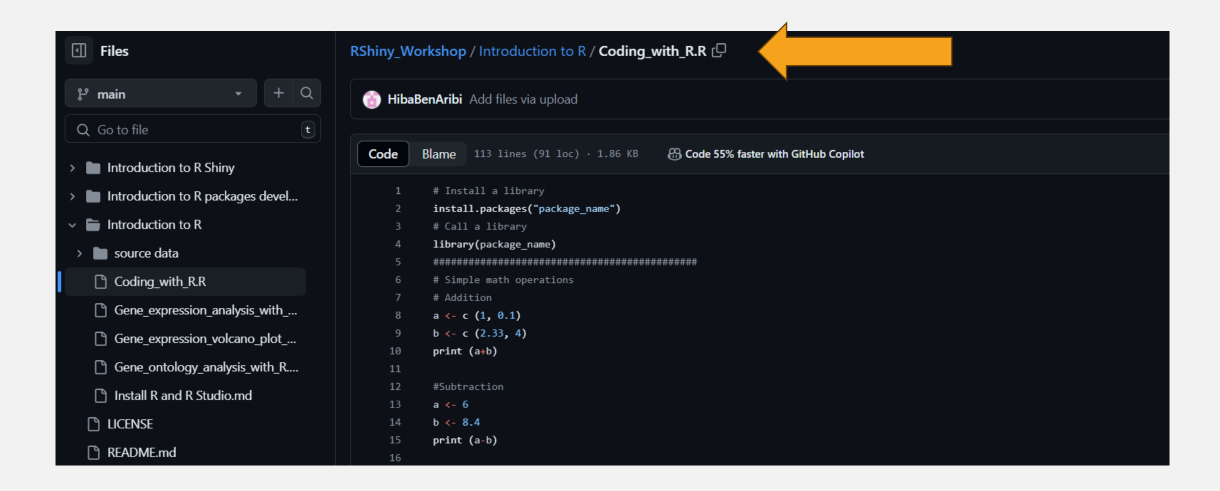


## **REQUIREMENTS**





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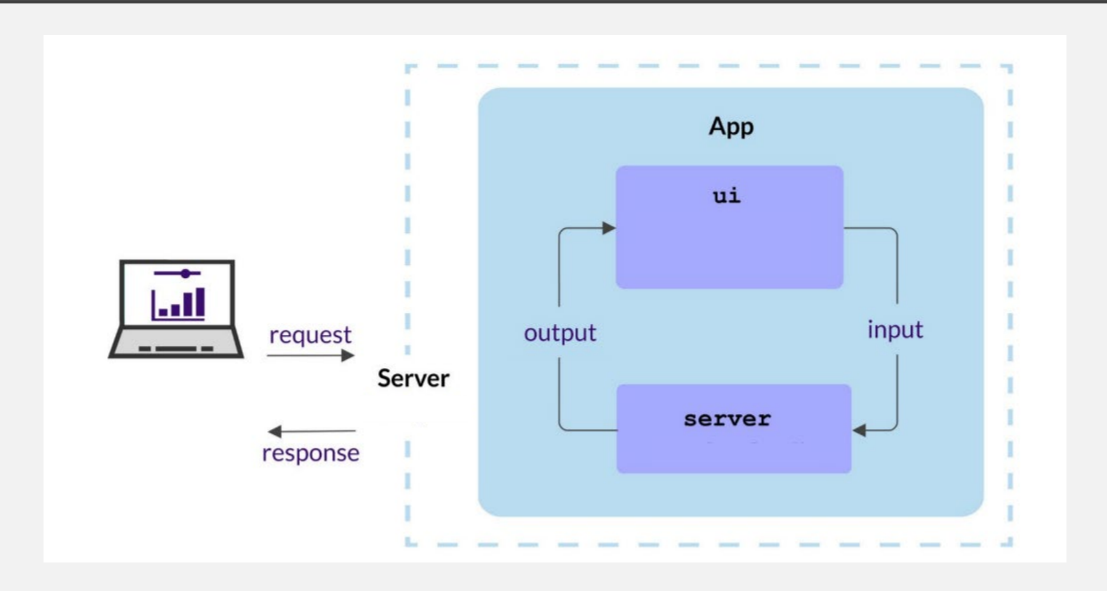
## SHINY



## Shiny is:

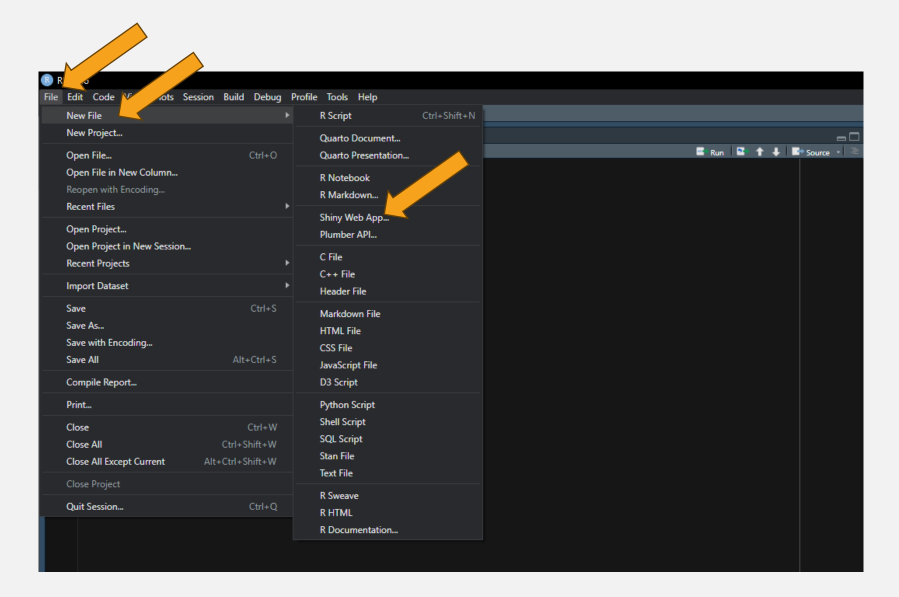
- ✓ An R package
- ✓ Allows the creation of web applications
- ✓ With interactive graphical user interfaces
- ✓ Using R instead of JS

## SHINY

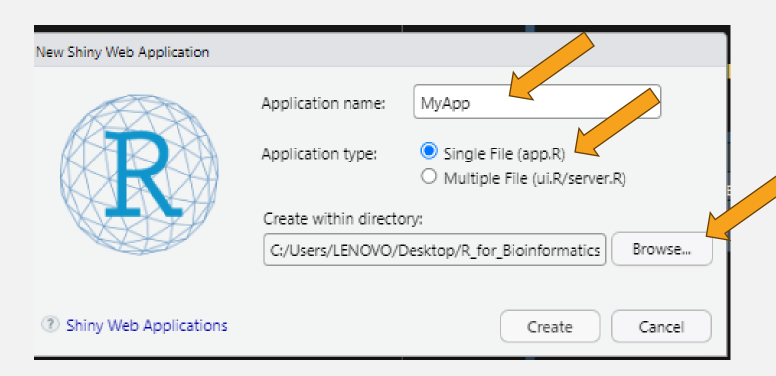


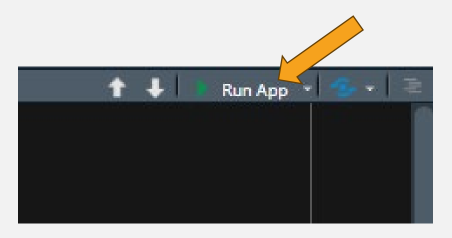
# LET'S CREATE A SHINY APP

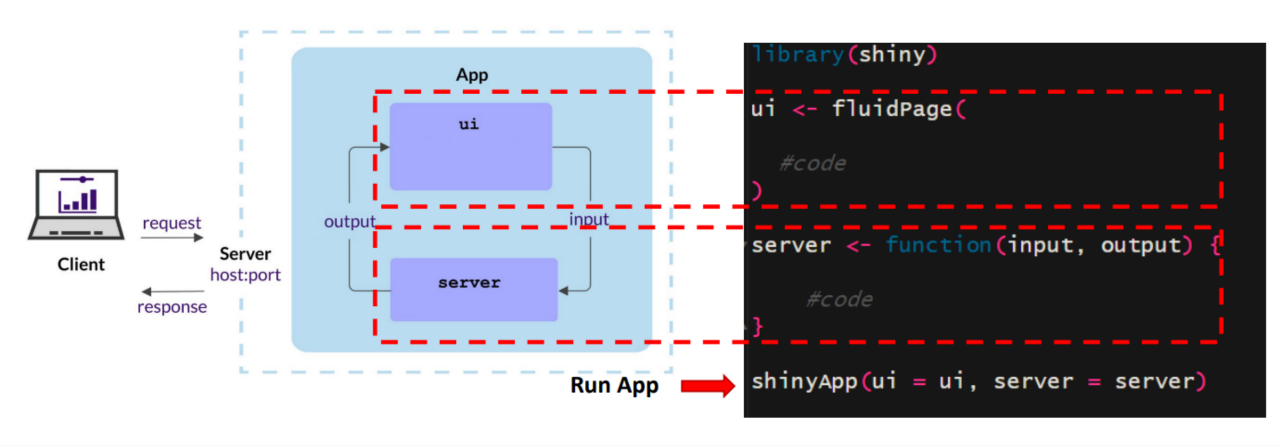
- I. Open R Studio
- 2. New File
- 3. Shiny web app

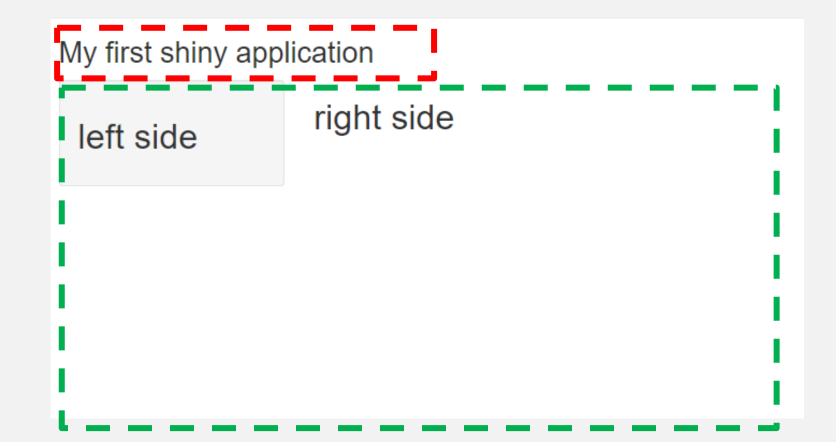


- I. Name it
- 2. Choose single file
- 3. Define directory
- 4. Run App







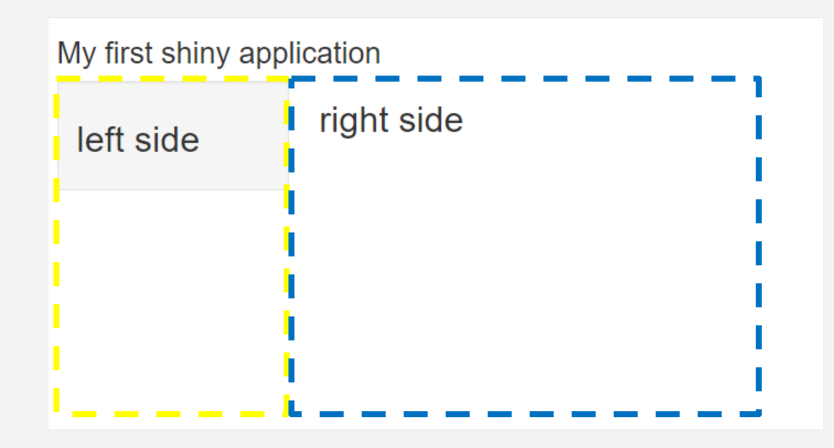


```
library(shiny)

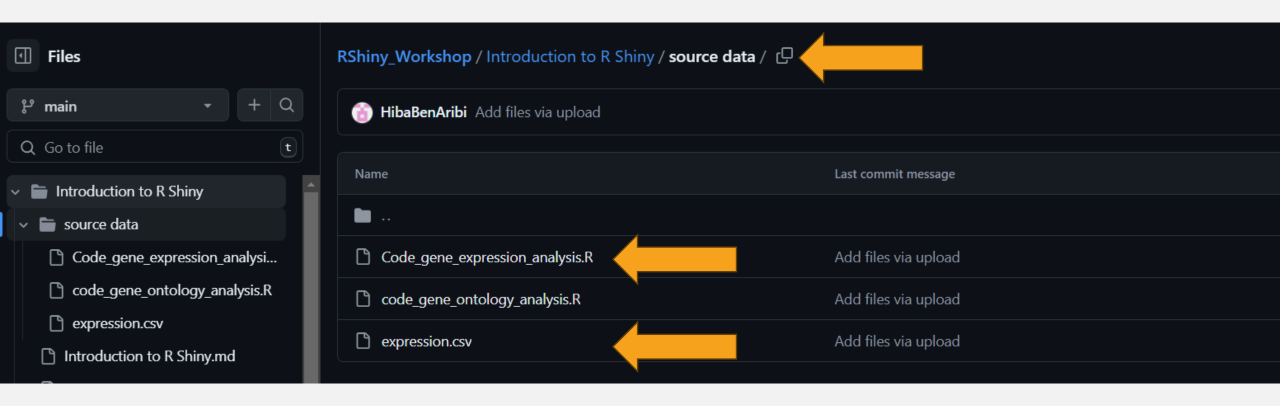
# Define UI

ui <- fluidPage(
    # Application title
    titlePanel("My first shiny application"),
    # Layout
    sidebarLayout(
    #left side
    sidebarPanel(
        h1("left side")
    ),

mainPanel(
    h1("right side")
)
</pre>
```

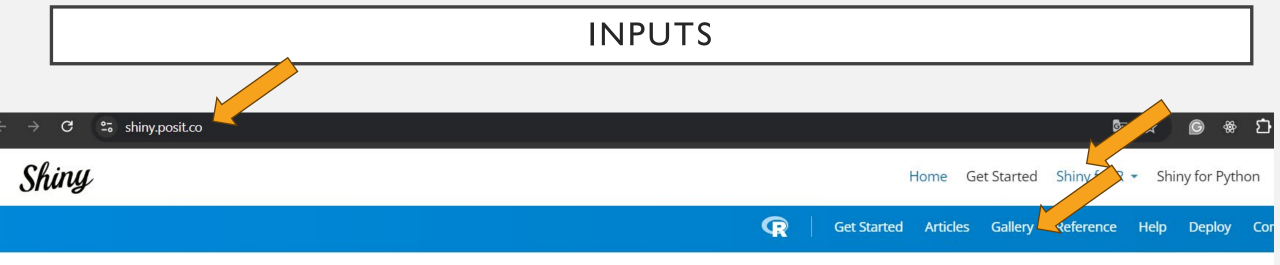


# INTEGRATE OUR CODE



```
2 library(readr)
   library (dplyr)
                                 Call Libraries
   library(ggplot2)
   expression <- read.csv("source data/expression.csv")</pre>
                                                                       Inputs
                                                                                                            df manipulation
   deg_pval <- 0.05 # max p value to define DEGs</pre>
   dea_logFC <- 2 # min logFC value</pre>
13 # gene expression analysis
   expression$diffexpressed <- "Not Significant"</pre>
   expression$diffexpressed[expression$log2FoldChange > deg_logFC & expression$pvalue < deg_pval ] <- "Upregulated genes"
   expression$diffexpressed[expression$log2FoldChange < -deg_logFC & expression$pvalue < deg_pval] <- "Downregulated genes"
     Visualize the results as a volcano plot
20 ggplot(data=expression, aes(x=log2FoldChange, y=-log10(pvalue), col=diffexpressed)) +
     geom_point() + theme_minimal() + theme(text = element_text(size = 15))
```

**Create plot** 

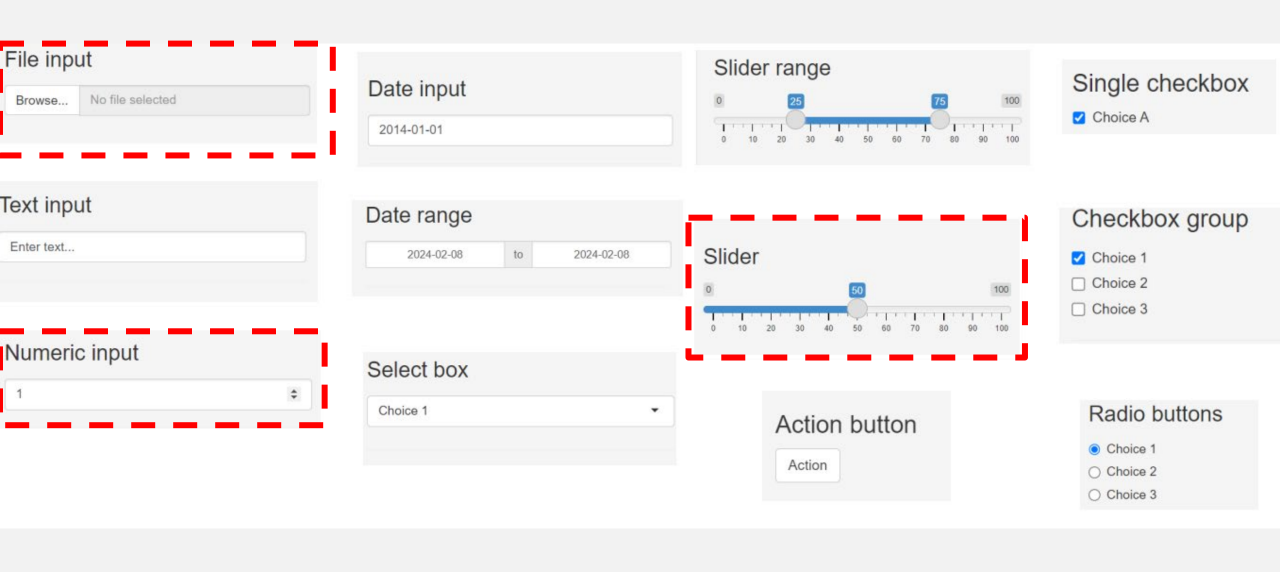


# Easy web apps for data science without the compromises

No web development skills required

Get started in R

Get started in Python



#### FILE INPUT

```
library(DT) # package for table management
sidebarPanel(
```

```
sidebarPanel(
   fileInput('ID', 'Text...'),
),
```



Request File Path in UI

```
server <- function(input, output) {
    #import file

data<- reactive({
    inFile <- input$data_file
    df <- read.csv(inFile$datapath, header = TRUE)
    return(df)
})</pre>
```

Read the file in server

#### NUMERIC INPUTS

Request File Path in UI

```
Numeric input
sidebarPanel(
  # Numeric input
 numericInput("ID", label = h3("Text"), value = Default_Value),
  sliderInput("ID", label = h3("Text"), min = number, max = number, value = Default_Value)
                                                                              Slider
```

value <- input\$ID</pre>

Call value in server

## TABLES / PLOTS

## In UI

```
DTOutput('table_name')
```

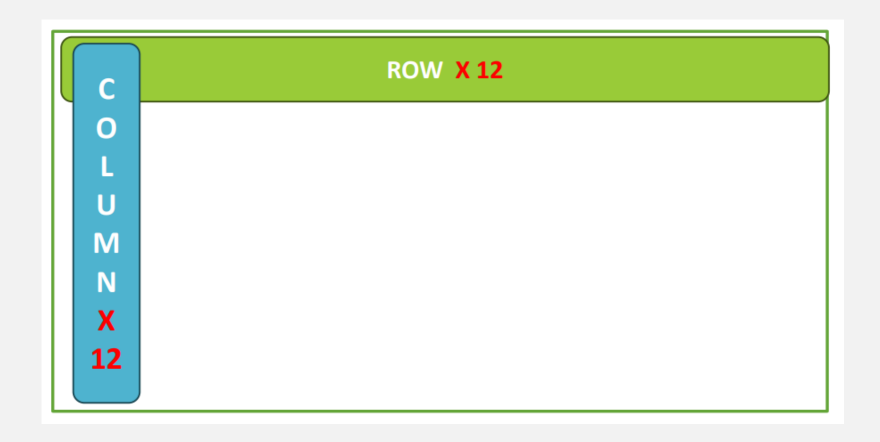
```
plotOutput("plot_name")
```

## **In Server**

```
output$table_name <- renderDT({
    #code
})</pre>
```

```
output$plot_name <- renderPlot({
    #code
})</pre>
```

## CUSTOMIZE UI



#### **CUSTOMIZE UI**

RShiny\_Workshop / Introduction to R Shiny / Introduction to R Shiny.md

#### Resources to Customize the UI

https://shiny.posit.co/r/getstarted/build-an-app/customizing-ui/theming.html

https://shiny.posit.co/r/articles/build/dashboards/

https://rstudio.github.io/shinydashboard/get\_started.html

https://daattali.com/shiny/shinycssloaders-demo/

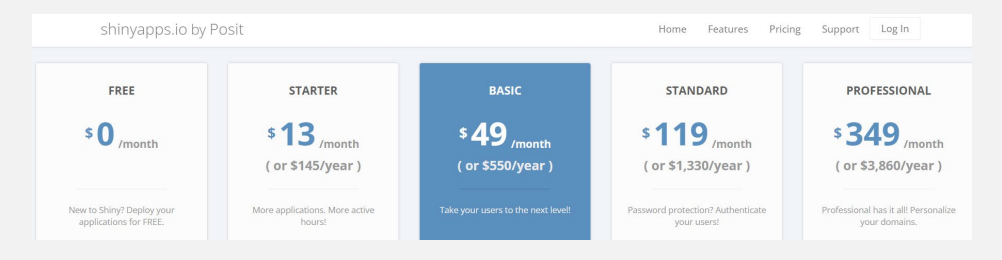
https://github.com/daattali/shinycssloaders

# **DEPLOYMENT**

#### SHINY APP DEPLOYMENT

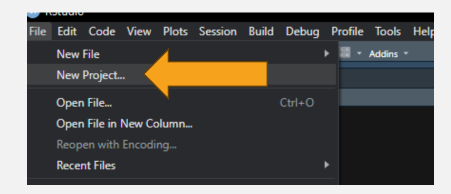
## Shiny could be shared as:

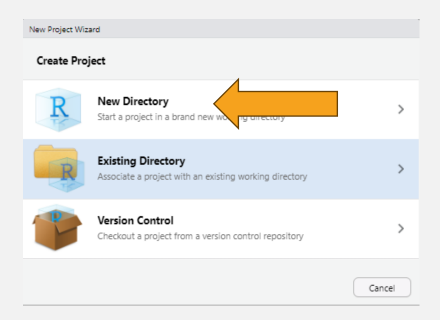
- √R code
- ✓ Host it on <a href="https://www.shinyapps.io/">https://www.shinyapps.io/</a>

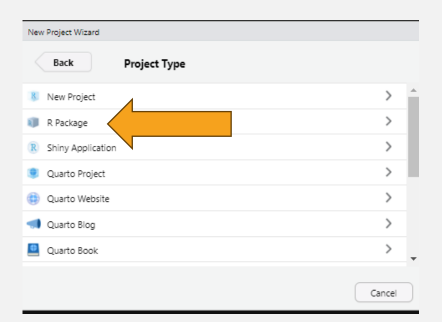


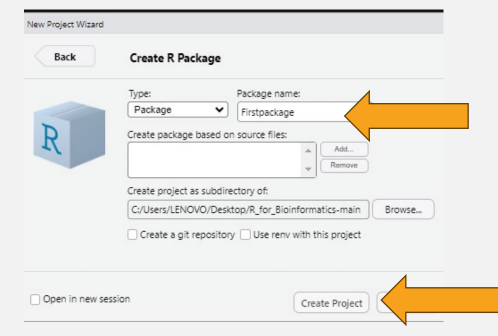
- ✓ Host on private server
- ✓ Implement as a function in an R package

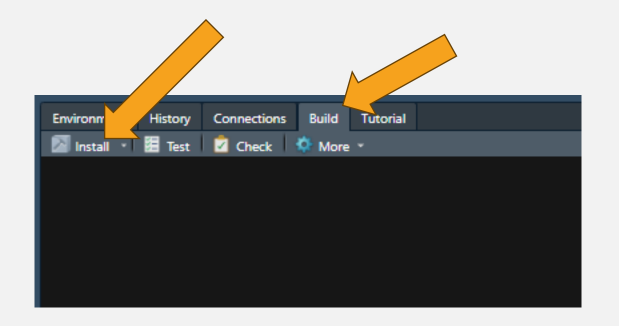
# R PACKAGE DEVELOPMENT











```
> library(Firstpackage)
> hello()
[1] "Hello, world!"
```

### **BUILD A FUNCTION**

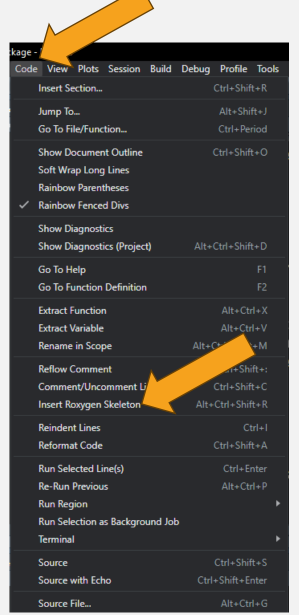
## To build a function

- ✓ Crate a R file
- ✓ Name it with the function name
- ✓ Write code inside function

```
function_name <- function() {
    #code
}</pre>
```

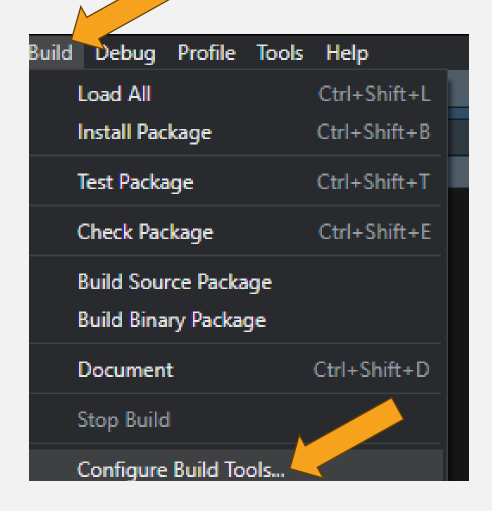
√ Build package again

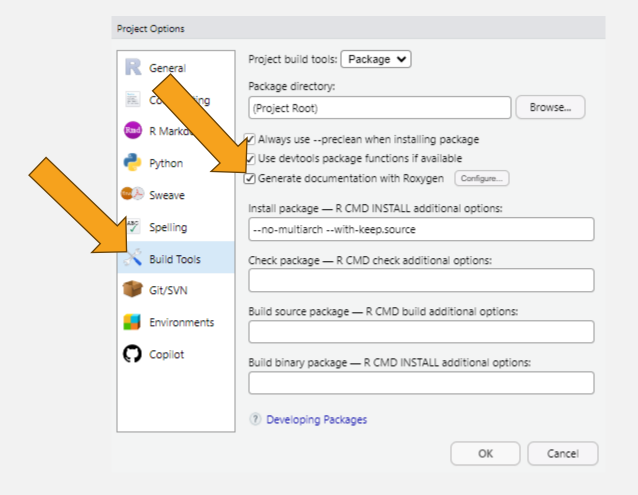
### DOCUMENT FUNCTION

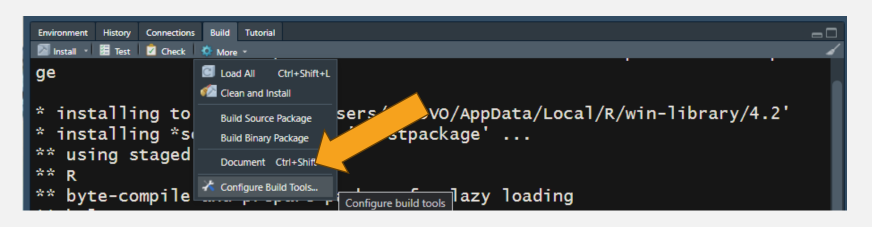


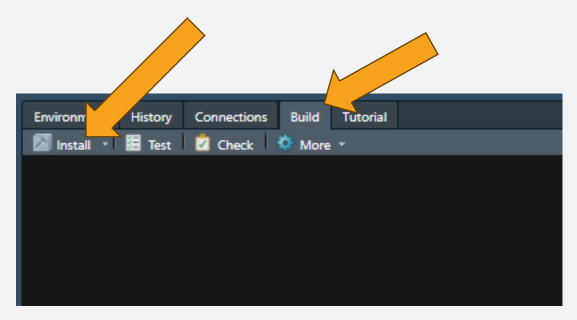
Insert Roxygen
Skeleton

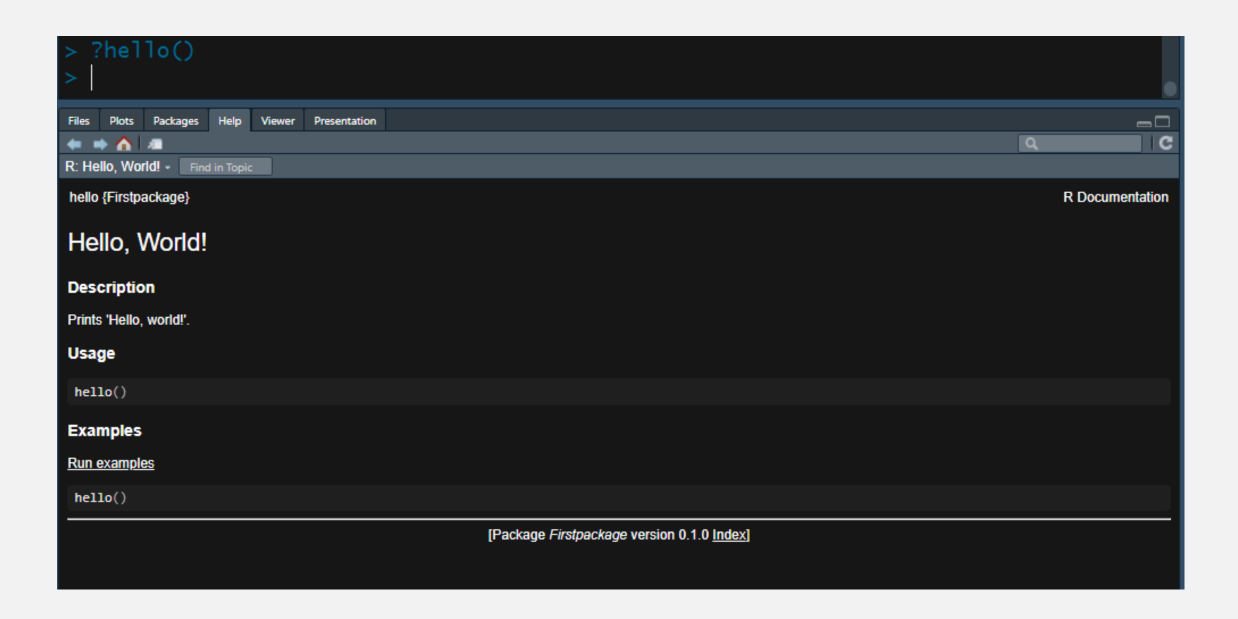
```
#' Title
#'
#' @param path
#'
#' @return
#' @export
#'
#' @examples
```











### INSTALLATION

- ✓ Import the package folder to GitHub
- √ You should be able to install it using a similar command

```
install.packages("devtools")
library(devtools)
devtools::install_github("omicscodeathon/repository/path")
library(package_name)
```

# THANK YOU FOR YOU ATTENTION!

Contact me via:

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