Making your packages accessible to non-programmer collaborators using the VisRseq framework

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June 26, 2016 | BioC 2016

Abstract

I will introduce the VisRseq framework and walk the participants through the quick process of creating modules called R-apps from R packages. I expect this to be specially useful for bioinformaticians and package developers that develop R-based analysis tools and would like to make them accessible to their non-programmer collaborators or to the public without having to spend significant time on creating extensive graphical user interfaces. I will walk the participants through several examples of creating diverse types of apps, from simple plotting (e.g. plots) to intermediate (e.g. clustering) to more advanced (e.g. edgeR and DEseq) packages. I will also show how several R-apps can be linked together to create more complex workflows. Participants will require having beginner/intermediate knowledge of R and a machine with R and Java installation.

Goals for this workshop

- Learn about VisRseq framework.
- Learn how to create R-apps through several examples.

Requirements

To get the most out of this workshop you are recommended to install the latest version of VisRseqfrom visrseq.github.io/downloads. You can still follow the instructions and create R-apps using plain R, however you will require VisRseq to be able to run the apps and see the results.

VisRseq Overview

Structure of an R-App

To create an R app you need a .R code containing the main script together with a .json specifying parameters. Each R app can have any number of "parameters", grouped into "categories". The .json file can be created manually or using the helper function in visrutils.R

```
source("https://raw.githubusercontent.com/hyounesy/visr-apps/master/visrutils.R")
```

which defines the following functions among other things.

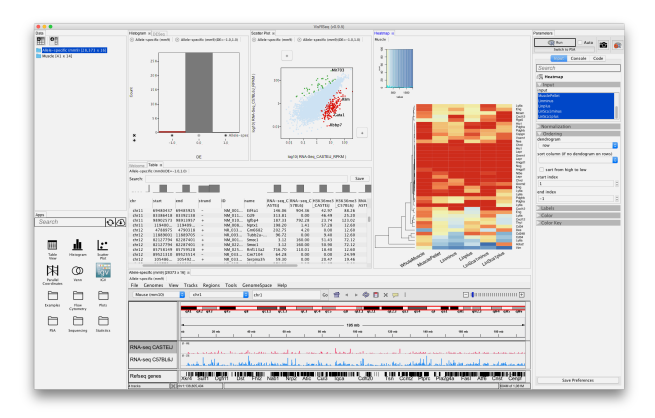


Figure 1:

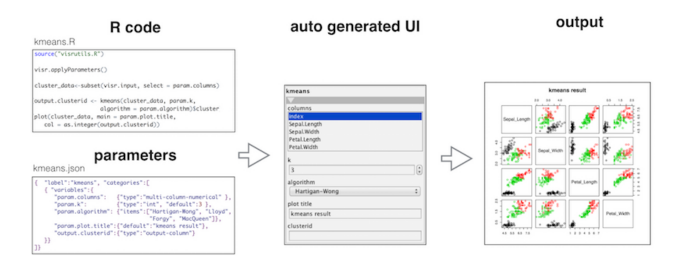


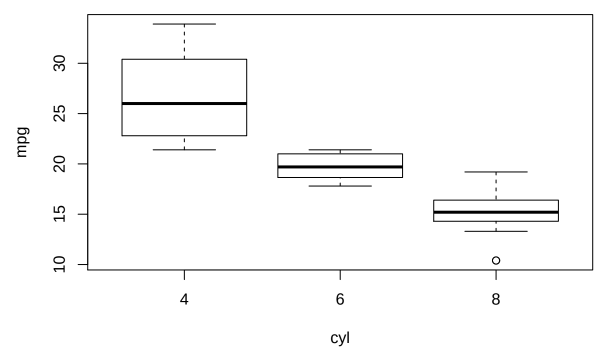
Figure 2:

Let's get started and create our first app!

Example 1: Boxplot

Let's start simple by creating a basic app that draws a barplot. Let's first write a basic boxplot script in a file simple_boxplot.R:

```
boxplot(mtcars[["mpg"]]~mtcars[["cyl"]], xlab = "cyl", ylab = "mpg")
```



Assume we would like to allow user to use their own data set and be able to specify x and y columns from their data.

We start the app definition by calling visr.app.start and specifying the app name. If we want be able to debug this app in R, we pass a test data set to debugdata. This parameter will be ignored when running the app in VisRseq. The data assigned to an app will be store in visr.input variable.

```
visr.app.start("Simple Boxplot", debugdata=mtcars)
```

We then add two parameters: y the numeric vector of data values that will be split into groups according to the parameter group. The parameter y has to be a numerical column in the input data, so we specify the type="column-numerical". The parameter group can be any column, so we specify type="column". Inorder to be able the debug the app inside the R, we also specify the values using debugvalue. This is optional but recommended.

```
visr.param("y", type="column-numerical", debugvalue="mpg")
visr.param("group", type="column", debugvalue="cyl")
```

This will create two parameters visr.param.y and visr.param.group. When running the script in R, they will be initialized with their corresponding debugvalue column names. When running the app in VisRseq, they will contain the string name of the column selected in the GUI.

We end the definition by calling visr.app.end. If argument printjson == TRUE, the json data for the app parameters will be printed to console. If argument writefile == TRUE, the output json will be written to a file specified by filename argument. If filename is not specified, the json file will be created at the same path of the R script, with the same name, but .json extension (i.e. simple_boxplot.json).

```
visr.app.end(printjson=TRUE, writefile = FALSE, filename = NULL)
```

```
## {
     "label": "Simple Boxplot",
##
     "info": "",
##
##
     "categories":[ {
##
       "label": "",
       "info": "",
##
       "variables": {
##
         "visr.param.y": {
##
            "type": "column-numerical"
##
##
         },
##
         "visr.param.group": {
##
            "type": "column"
##
##
##
     }]
## }
```

We now have the json file. So we just have to modify our original R script to use the correct parameter names.

```
visr.applyParameters()
boxplot(visr.input[[visr.param.y]]~visr.input[[visr.param.group]], xlab = visr.param.group, ylab = visr
```

Done! now if we place the simple_boxplot.R and simple_boxplot.json inside VisRseq/visr/srcR and hit the Refresh icon above the apps pane, we will have our first custom app in VisRseq. For demonstration purpose here we have used the app to draw a boxplot for the iris dataset.

here is the complete source code for simple_boxplot.R app:

```
source("visrutils.R")
visr.app.start("Simple Boxplot", debugdata=mtcars)
visr.param("y", type="column-numerical", debugvalue="mpg")
visr.param("group", type="column", debugvalue="cyl")
visr.app.end(printjson=TRUE, writefile = FALSE, filename = NULL)
visr.applyParameters()
boxplot(visr.input[[visr.param.y]]~visr.input[[visr.param.group]], xlab = visr.param.group, ylab = visr
```

Example 2: Kmeans

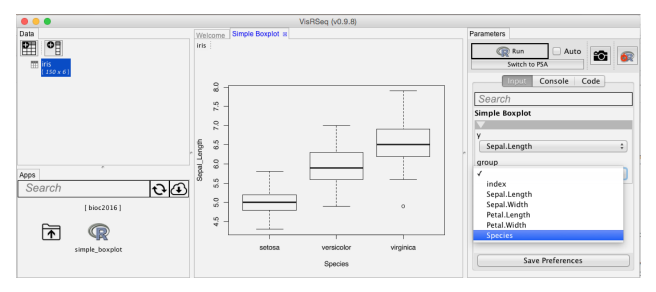


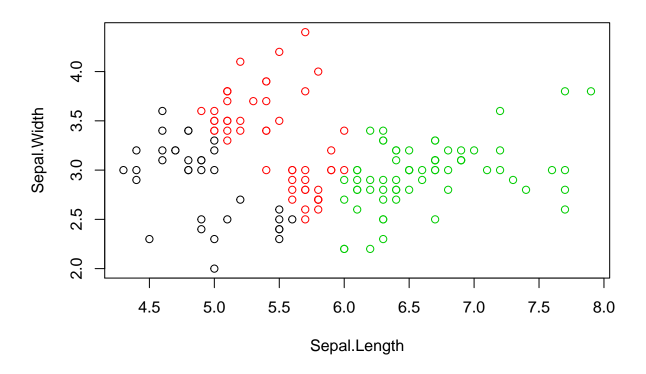
Figure 3:

```
# start parameter definition
visr.app.start("kmeans", debugdata = iris)
visr.category("clustering parameters")
visr.param("columns", type = "multi-column-numerical", debugvalue = c("Sepal.Length", "Sepal.Width"))
visr.param("k", default = 3)
visr.param("algorithm", items = c("Hartigan-Wong", "Lloyd", "Forgy", "MacQueen"))
visr.category("output")
visr.param("plot.type", items = c("scatter plot", "histogram"))
visr.param("plot.title", default = "kmeans results")
visr.param("output.clusterid", type = "output-column")
visr.app.end(printjson=TRUE)
## {
##
     "label": "kmeans",
     "info": "",
##
##
     "categories":[ {
##
       "label": "clustering parameters",
       "info": "",
##
       "variables": {
##
##
         "visr.param.columns": {
##
           "type": "multi-column-numerical"
##
         },
         "visr.param.k": {
##
##
           "type": "int",
           "default": 3
##
##
         },
##
         "visr.param.algorithm": {
##
           "type": "string",
           "items": [ "Hartigan-Wong", "Lloyd", "Forgy", "MacQueen" ]
##
##
         }
##
       }
##
     },
     {
```

##

```
##
       "label": "output",
       "info": "",
##
##
       "variables": {
##
         "visr.param.plot.type": {
           "type": "string",
##
##
           "items": [ "scatter plot", "histogram" ]
##
         },
##
         "visr.param.plot.title": {
           "type": "string",
##
##
           "default": "kmeans results"
##
         },
         "visr.param.output.clusterid": {
##
           "type": "output-column"
##
##
##
##
     }]
## }
visr.applyParameters()
cluster_data<-subset(visr.input, select = visr.param.columns)</pre>
visr.param.output.clusterid <- kmeans(cluster_data, visr.param.k, algorithm = visr.param.algorithm)$clu
# plotting options
if (visr.param.plot.type == "scatter plot") {
    plot(cluster_data, main = visr.param.plot.title, col = as.integer(visr.param.output.clusterid))
} else {
  clustersTable <- table(visr.param.output.clusterid)</pre>
  lbls<-as.character(clustersTable)</pre>
  bplt<-barplot(clustersTable , xlab="cluster ID", ylab = "cluster size", main=visr.param.plot.title)</pre>
  text(y = 0, x = bplt, labels=lbls, xpd=TRUE, adj=c(0.5, -1))
}
```

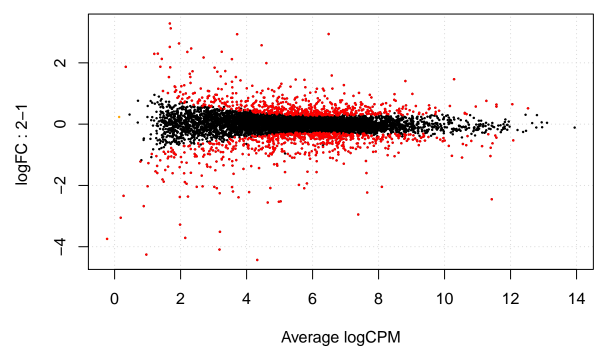
kmeans results



Example 3: Differntial expression analysis using edgeR

```
library("edgeR", quietly=T)
## Warning: package 'edgeR' was built under R version 3.2.2
## Warning: package 'limma' was built under R version 3.2.1
countdata = read.table("https://raw.githubusercontent.com/hyounesy/bioc2016.visrseq/master/data/counts.
x <- countdata
head(x)
               CT.PA.1 CT.PA.2 KD.PA.3 KD.PA.4 CT.SI.5 KD.SI.6 CT.SI.7
##
## FBgn0000008
                    76
                             71
                                     87
                                              68
                                                     137
                                                             115
                                                                       82
## FBgn0000017
                           3087
                                   3029
                                            3264
                                                    7014
                                                            4322
                                                                     3926
                  3498
## FBgn000018
                   240
                            306
                                    288
                                             307
                                                     613
                                                             528
                                                                      485
## FBgn0000032
                    611
                            672
                                    694
                                             757
                                                    1479
                                                            1361
                                                                     1351
## FBgn0000042
                 40048
                          49144
                                  70574
                                           72850
                                                   97565
                                                           95760
                                                                    99372
## FBgn0000043
                 15910
                          18194
                                  31086
                                           34085
                                                           42389
                                                                    29671
                                                   34171
group1 <- c("CT.PA.1", "CT.PA.2")</pre>
group2 <- c("KD.PA.3", "KD.PA.4")
groups <- factor(c(rep(1, length(group1)), rep(2, length(group2)))) # c(1, 1, 2, 2)
# create edgeR's container for RNA-seq count data
y <- DGEList(counts=x[, c(group1, group2)], group = groups)
```

```
# estimate normalization factors
y <- calcNormFactors(y)
# estimate tagwise dispersion (simple design)
y <- estimateCommonDisp(y)
y <- estimateTagwiseDisp(y)
# test for differential expression using classic edgeR approach
et <- exactTest(y)
# total number of DE genes in each direction
is.de <- decideTestsDGE(et, adjust.method = "BH", p.value = 0.05, lfc = 0)
# The log-fold change for each gene is plotted against the average abundance
plotSmear(y, de.tags=rownames(y)[is.de!=0])</pre>
```



```
## edgeR parameters
#countdata = read.table("https://raw.githubusercontent.com/hyounesy/bioc2016.visrseq/master/data/counts
visr.app.start("edgeR", debugdata = countdata)
visr.param("group1", type = "multi-column-numerical", debugvalue = c("CT.PA.1", "CT.PA.2"))
visr.param("group2", type = "multi-column-numerical", debugvalue = c("KD.PA.3", "KD.PA.4"))
visr.param("output.de", label = "DE clusters", type = "output-column")
visr.app.end(printjson=TRUE, writefile=T)

## {
## "label": "edgeR",
```

```
"label": "edgeR",
     "info": "",
##
##
     "categories":[ {
       "label": "",
##
       "info": "",
##
##
       "variables": {
##
         "visr.param.group1": {
##
           "type": "multi-column-numerical"
##
         "visr.param.group2": {
##
```

```
##
           "type": "multi-column-numerical"
##
         },
##
         "visr.param.output.de": {
           "label": "DE clusters",
##
           "type": "output-column"
##
##
         }
##
##
     }]
## }
```

visr.applyParameters()

```
## edgeR code
library("edgeR")
x <- visr.input
groups <- factor(c(rep(1, length(visr.param.group1)), rep(2, length(visr.param.group2)))) # c(1, 1, 2,
# create edgeR's container for RNA-seg count data
y <- DGEList(counts=x[, c(visr.param.group1, visr.param.group2)], group = groups)
# estimate normalization factors
y <- calcNormFactors(y)</pre>
# estimate tagwise dispersion (simple design)
y <- estimateCommonDisp(y)</pre>
y <- estimateTagwiseDisp(y)</pre>
# test for differential expression using classic edgeR approach
et <- exactTest(y)
# total number of DE genes in each direction
is.de <- decideTestsDGE(et, adjust.method = "BH", p.value = 0.05, lfc = 0)
# export the results to VisRseq
visr.param.output.de <- as.factor(is.de)</pre>
# The log-fold change for each gene is plotted against the average abundance
plotSmear(y, de.tags = rownames(y)[is.de != 0])
```