

Programming: practical 2a solutions

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IN THIS question, we are going to use a `for` statement to loop over a large data set and construct some scatter plots. To generate the data, run the following piece of R code

```
library("nclRprogramming")
data(dummy_data)
dd = dummy_data
```

The data frame `dd` represents an experiment, where we have ten treatments: A, B, \dots, J and measurements at some time points. We want to create a scatter plot of measurement against time, for each treatment type.

1. First we create a scatter plot of one treatment:

```
plot(dd[dd$treatments=="A",]$time,
     dd[dd$treatments=="A",]$measurement)
```

Since the colnames are a bit long, let's shorten them:

```
colnames(dd) = c("m", "t", "trts")
```

2. To generate a scatter-plot for each treatment, we need to iterate over the different treatment types:

```
for(treat in unique(dd$trts)) {
  plot(dd[dd$trts==treat,]$t,
       dd[dd$trts==treat,]$m)
  readline("Hit return for next plot\n")
}
```

A few questions:

- What does `unique(dd$trts)` give?

```
## It gives all treatments.
```

- In the `for` loop, what variable is changing? What are its possible values?

```
# The treat variable is changing. It goes through
# the different treatments.
```

- What does the `readline` function do?

3. The default axis labels aren't great. So we can change the x -axis label using `xlab`:

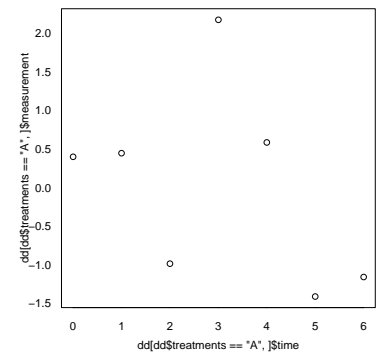


Figure 1: Measurements against time.

```
plot(dd[dd$trts==treat,]$t,
     dd[dd$trts==treat,]$m,
     xlab="Time", ylab="Measurement")
```

Use ylab to alter the y-axis label.

4. To add a title to a plot we use the main argument, viz:

```
plot(dd[dd$trts=="A",]$t,
     dd[dd$trts=="A",]$m,
     main="Treatment",
     xlab="Time", ylab="Measurement")
```

We can combine strings/characters using the paste function,

```
paste("Treatment", treat)

## [1] "Treatment J"
```

Rather than have a static title, make the title of each plot display the treatment type.

```
plot(dd[dd$trts==treat,]$t,
     dd[dd$trts==treat,]$m,
     main=paste("Treatment", treat),
     xlab="Time", ylab="Measurement")
```

5. The y-axis range should really be the same in all graphics. Add a ylim argument to fix the range.¹

```
range(dd$m)

## [1] -1.639  8.113

plot(dd[dd$trts==treat,]$t,
     dd[dd$trts==treat,]$m,
     main=paste("Treatment", treat),
     xlab="Time", ylab="Measurement",
     ylim=c(-2, 10))
```

6. At each iteration, use the message function to print the average measurement level across all time points.

```
## Within the for loop have the line
message(mean(dd[dd$trts == treat, ]$m))
```

7. On each graph, highlight any observations with a blue point if they are larger than the mean + standard deviations or less than the mean - standard deviations. Use the points function to highlight a point.² For example, to highlight the points (1,2) and (3, 4) we use the command:

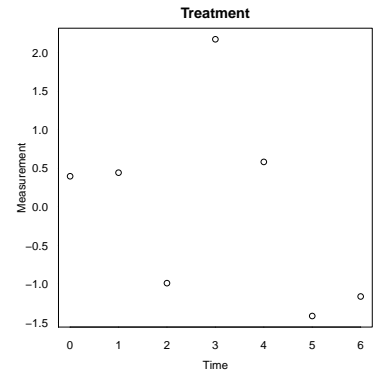


Figure 2: Measurements against time with a title.

¹ Hint: Work out the range before the for loop.

² Hint: You don't need if statements here. Just subset your data frame and pass this new data frame to the points function.

```
points(c(1, 3), c(2, 4), col = 2)
```

```
plot(dd[sel,]$t, dd[sel,]$m,
      ylab=treat, xlab="Time",
      main=paste("Treatment", treat))
##Select a particular treatment
sel = (dd$trts == treat)

##Calculate the limits
values = dd[sel,]$m
message(mean(values))
upper_lim = mean(values) + sd(values)
lower_lim = mean(values) - sd(values)

##Extract the points
up_row = dd[sel & dd$m > upper_lim,]
low_row = dd[sel & dd$m < lower_lim,]

##pch=19 gives a solid dot
##See ?points
points(up_row$t, up_row$m, col=4, pch=19)
points(low_row$t, low_row$m, col=4, pch=19)
```

8. Suppose we wanted to save individual graphs. Add in the pdf function to save the resulting graph. To get unique file names, use the paste command:

```
filename = paste("file", treat, ".pdf", sep = "")
```

9. Put the your code, i.e. the for loop and plotting commands, in a function which takes the data frame as an argument.
10. Alter your function to take another argument where you can save the graph in a different directory.

Final piece of code

```
viewgraphs = function(dd, colour=TRUE, save=FALSE) {
  for(treat in unique(dd$trts)) {
    if(save) {
      filename = paste("file", treat, ".pdf", sep="")
      pdf(filename)
    }

    ##Use a different shape in the points
    if(colour) pch = 19
    else pch = 22
  }
}
```

```

##Do selection one
sel = (dd$trts == treat)
plot(dd[sel,]$t, dd[sel,]$m,
      ylab=treat, xlab="Time",
      main=paste("Treatment", treat))

##Calculate the limits
values = dd[sel,]$m
message(mean(values))
upper_lim = mean(values) + sd(values)
lower_lim = mean(values) - sd(values)

##Extract the points
up_row = dd[sel & dd$m > upper_lim,]
low_row = dd[sel & dd$m < lower_lim,]

##pch=19 gives a solid dot
##See ?points
points(up_row$t, up_row$m, col=4, pch=19)
points(low_row$t, low_row$m, col=4, pch=19)
if(save){
  dev.off()
} else {
  readline("Hit return for next plot\n")
}
}
}

```

Solutions

Solutions are contained within this package:

```

library("nclRprogramming")
vignette("solutions2a", package = "nclRprogramming")

```