# Heatmaps - the gene expression edition

Jeff Oliver

12 September, 2017

### Learning objectives

- 1. one
- 2. two
- 3. three

[DESCRIPTION O announcement]	R MOTIVATION;	2-4 sentences th	at would be u	ised for an
Getting started				
[TOPIC ONE]				

# [TOPIC TWO]

Install two packages (in console)

```
install.packages("ggplot2")
install.packages("tidyr")
```

Create a script that will save all our work; also, load additional packages

```
# Expression heatmap
# Jeff Oliver
# jcoliver@email.arizona.edu
# 2017-09-14

library("tidyr")
library("ggplot2")
```

#### Read in the data

```
# Expression heatmap
# Jeff Oliver
# jcoliver@email.arizona.edu
# 2017-09-14
```

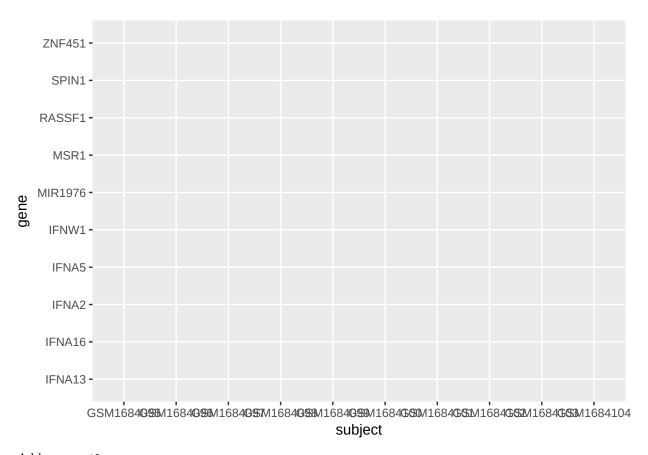
```
library("tidyr")
library("ggplot2")
exp.data <- read.csv(file = "data/GSE68849-expression.csv", stringsAsFactors = FALSE)
Investigate data with summary
summary(exp.data)
                      treatment
                                          IFNA5
##
     subject
  Length:10
                     Length:10
                                                78.12
                                      Min. :
  Class :character
                     Class : character
                                                 86.80
##
                                      1st Qu.:
## Mode :character Mode :character
                                      Median: 3578.65
##
                                      Mean : 5019.12
##
                                      3rd Qu.: 9093.19
##
                                      Max. :15018.41
##
       IFNA13
                         IFNA2
                                         SPIN1
                                                        ZNF451
## Min. : 98.34 Min.
                          : 100.7
                                    Min.
                                            :104.2 Min.
                                                           :232.2
  1st Qu.: 110.58 1st Qu.: 124.9
                                    1st Qu.:109.9 1st Qu.:289.3
## Median : 7765.56 Median : 9665.8
                                     Median: 119.4 Median: 302.4
## Mean : 8617.42 Mean
                          :10647.0 Mean :118.2 Mean
                                                          :333.9
## 3rd Qu.:16655.33 3rd Qu.:20046.6
                                     3rd Qu.:123.3
                                                    3rd Qu.:325.3
                                                          :569.3
## Max. :18974.16 Max. :24029.1 Max. :134.8 Max.
##
       IFNA16
                       RASSF1
                                      IFNW1
                                                        MSR1
## Min. : 101.2 Min.
                         :244.8 Min.
                                         : 89.26 Min.
                                                          : 89.58
## 1st Qu.: 158.9 1st Qu.:291.6 1st Qu.: 95.79
                                                  1st Qu.: 95.48
## Median: 8968.3 Median: 330.5 Median: 2883.71
                                                   Median :100.83
## Mean :10378.3
                   Mean :323.5 Mean :3834.03
                                                   Mean :103.43
## 3rd Qu.:20543.8
                    3rd Qu.:353.3 3rd Qu.:7297.51
                                                   3rd Qu.:108.38
## Max.
         :23060.5
                    Max. :386.8 Max. :9198.55
                                                   Max. :125.94
      MIR1976
##
         : 82.85
## Min.
## 1st Qu.: 87.79
## Median :103.58
## Mean :100.16
## 3rd Qu.:108.23
## Max. :120.63
Transform data to long format
# Expression heatmap
# Jeff Oliver
\# jcoliver@email.arizona.edu
# 2017-09-14
library("tidyr")
library("ggplot2")
exp.data <- read.csv(file = "data/GSE68849-expression.csv", stringsAsFactors = FALSE)
exp.long <- gather(data = exp.data,
```

Look at the top of the long data:

key = gene,

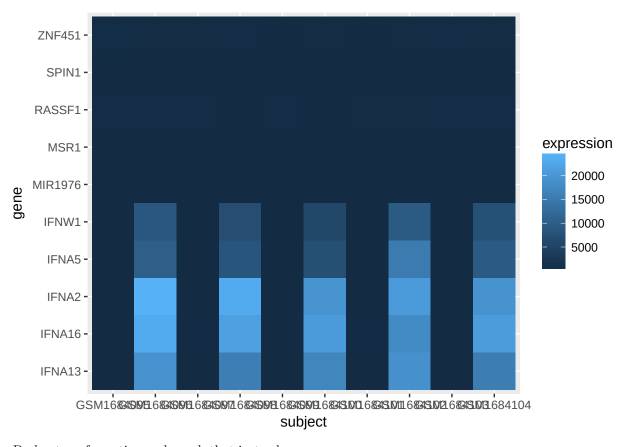
value = expression,
-subject, -treatment)

```
head(exp.long)
##
        subject treatment gene expression
## 1 GSM1684095 control IFNA5
                                   83.12937
## 2 GSM1684096 influenza IFNA5 10096.47000
## 3 GSM1684097 control IFNA5 97.80374
## 4 GSM1684098 influenza IFNA5 8180.98900
## 5 GSM1684099 control IFNA5 81.70878
## 6 GSM1684100 influenza IFNA5 7054.90200
Plot!
# Expression heatmap
# Jeff Oliver
# jcoliver@email.arizona.edu
# 2017-09-14
library("tidyr")
library("ggplot2")
exp.data <- read.csv(file = "data/GSE68849-expression.csv", stringsAsFactors = FALSE)</pre>
exp.long <- gather(data = exp.data,</pre>
                   key = gene,
                   value = expression,
                   -subject, -treatment)
exp.heatmap <- ggplot(data = exp.long, mapping = aes(x = subject,</pre>
                                                      y = gene,
                                                      fill = expression))
exp.heatmap
```



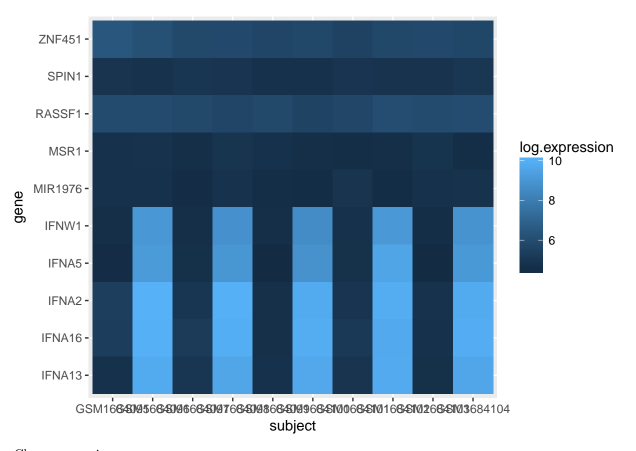
### $\operatorname{Add}$ geom\_tile

```
# Expression heatmap
# Jeff Oliver
# jcoliver@email.arizona.edu
# 2017-09-14
library("tidyr")
library("ggplot2")
exp.data <- read.csv(file = "data/GSE68849-expression.csv", stringsAsFactors = FALSE)</pre>
exp.long <- gather(data = exp.data,</pre>
                   key = gene,
                    value = expression,
                    -subject, -treatment)
exp.heatmap <- ggplot(data = exp.long, mapping = aes(x = subject,</pre>
                                                        y = gene,
                                                        fill = expression)) +
  geom_tile()
exp.heatmap
```



Do log transformation and graph that instead

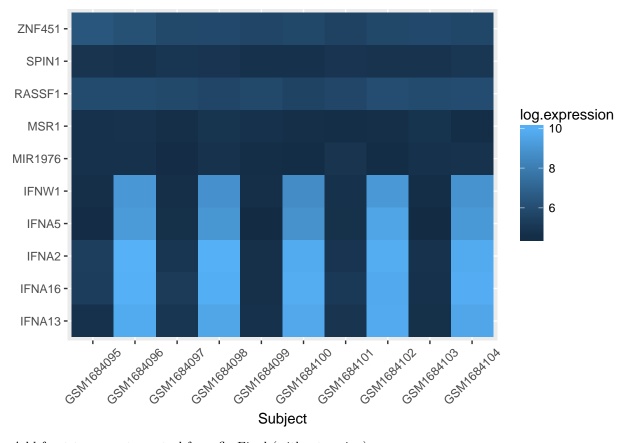
```
# Expression heatmap
# Jeff Oliver
# jcoliver@email.arizona.edu
# 2017-09-14
library("tidyr")
library("ggplot2")
exp.data <- read.csv(file = "data/GSE68849-expression.csv", stringsAsFactors = FALSE)</pre>
exp.long <- gather(data = exp.data,</pre>
                    key = gene,
                    value = expression,
                    -subject, -treatment)
exp.long$log.expression <- log(exp.long$expression)</pre>
exp.heatmap <- ggplot(data = exp.long, mapping = aes(x = subject,</pre>
                                                        y = gene,
                                                        fill = log.expression)) +
  geom_tile()
exp.heatmap
```



## Clean up x-axis

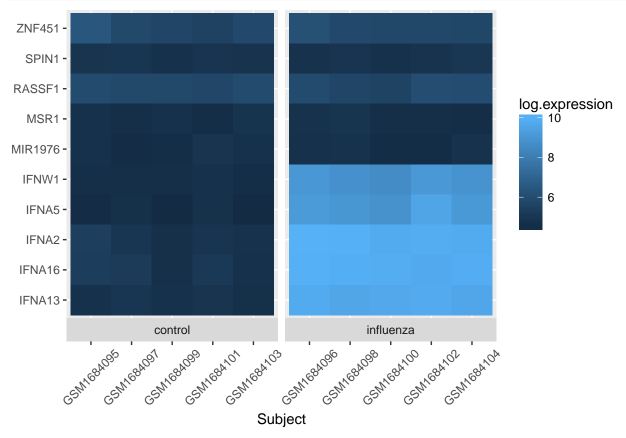
```
# Expression heatmap
# Jeff Oliver
# jcoliver@email.arizona.edu
# 2017-09-14
library("tidyr")
library("ggplot2")
exp.data <- read.csv(file = "data/GSE68849-expression.csv", stringsAsFactors = FALSE)</pre>
exp.long <- gather(data = exp.data,</pre>
                    key = gene,
                    value = expression,
                    -subject, -treatment)
exp.long$log.expression <- log(exp.long$expression)</pre>
exp.heatmap <- ggplot(data = exp.long, mapping = aes(x = subject,</pre>
                                                        y = gene,
                                                        fill = log.expression)) +
  geom_tile() +
  xlab(label = "Subject") +
  theme(axis.title.y = element_blank(),
        axis.text.x = element_text(angle = 45, vjust = 0.5))
```





Add facet to separate control from flu Final (without saving)

```
# Expression heatmap
# Jeff Oliver
\# jcoliver@email.arizona.edu
# 2017-09-14
library("tidyr")
library("ggplot2")
exp.data <- read.csv(file = "data/GSE68849-expression.csv", stringsAsFactors = FALSE)
exp.long <- gather(data = exp.data,</pre>
                   key = gene,
                    value = expression,
                    -subject, -treatment)
exp.long$log.expression <- log(exp.long$expression)</pre>
exp.heatmap <- ggplot(data = exp.long, mapping = aes(x = subject,</pre>
                                                       y = gene,
                                                       fill = log.expression)) +
  geom_tile() +
  xlab(label = "Subject") +
  facet_grid(~ treatment, switch = "x", scales = "free_x", space = "free_x") +
```



#### Final script:

#### Additional resources

- resource one
- resource two
- A PDF version of this lesson

Back to learn-r main page

Questions? e-mail me at jcoliver@email.arizona.edu.