# Lab 11 HW

## Lorena Cuellar

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```
expr <- read.table("rs8067378_ENSG00000172057.6.txt")
head(expr)

## sample geno exp
## 1 HG00367 A/G 28.96038
## 2 NA20768 A/G 20.24449
## 3 HG00361 A/A 31.32628
## 4 HG00135 A/A 34.11169
## 5 NA18870 G/G 18.25141
## 6 NA11993 A/A 32.89721

nrow(expr)
```

### ## [1] 462

Q13: Read this file into R and determine the sample size for each genotype and their corresponding median expression levels for each of these genotypes.

The sample size for each genotype

```
table(expr$geno)
```

```
##
## A/A A/G G/G
## 108 233 121
```

Median expression levels for each of these genotypes

```
library(doBy)
summaryBy(exp~geno, data = expr, FUN=median)
```

```
## geno exp.median
## 1 A/A 31.24847
## 2 A/G 25.06486
## 3 G/G 20.07363
```

### summary(expr)

```
##
       sample
                            geno
                                                  exp
##
    Length:462
                        Length:462
                                                    : 6.675
                                             Min.
##
    Class : character
                        Class : character
                                             1st Qu.:20.004
    Mode :character
##
                        Mode :character
                                             Median :25.116
##
                                             Mean
                                                    :25.640
##
                                             3rd Qu.:30.779
##
                                             Max.
                                                    :51.518
```

## library(ggplot2)

Q14: Generate a boxplot with a box per genotype, what could you infer from the relative expression value between A/A and G/G displayed in this plot? Does the SNP effect the expression of ORMDL3?

It can be inferred that the A/A genotype has higher expression levels than the G/G genotype from this plot. Additionally, it does seem like this SNP effects the expression of ORMDL3, it looks like the SNP increases ORMDL3 expression.

## ggplot(expr) + aes(x=geno, exp, fill=geno) + geom\_boxplot(notch=TRUE)

