## Lab 11

Edwin Ruiz

2024-05-19

## Section 1

```
mxl <- read.csv("373531-SampleGenotypes-Homo_sapiens_Variation_Sample_rs8067378 (2).csv")
head(mxl)
     Sample.. Male. Female. Unknown. Genotype.. forward. strand. Population.s. Father
##
## 1
                       NA19648 (F)
                                                          A|A ALL, AMR, MXL
## 2
                       NA19649 (M)
                                                          G|G ALL, AMR, MXL
## 3
                       NA19651 (F)
                                                          A|A ALL, AMR, MXL
## 4
                       NA19652 (M)
                                                          G|G ALL, AMR, MXL
## 5
                       NA19654 (F)
                                                          G|G ALL, AMR, MXL
                                                          A|G ALL, AMR, MXL
## 6
                       NA19655 (M)
##
     Mother
## 1
## 2
## 3
## 4
## 5
## 6
table(mxl$Genotype..forward.strand.)
##
## A|A A|G G|A G|G
  22 21 12
table(mxl$Genotype..forward.strand.) / nrow(mxl) * 100
##
##
               A|G
                        G|A
                                G|G
       A \mid A
## 34.3750 32.8125 18.7500 14.0625
```

## Extra Credit

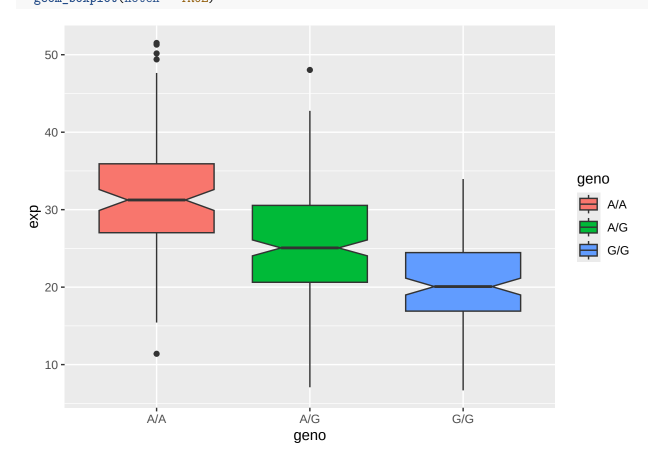
So, you processed about  $\sim 230$  samples and did the normalization on a genome level. Now, you want to find whether there is any association of the 4 asthma-associated SNPs (rs8067378...) on ORMDL3 expression.

How many samples do we have?

```
## 3 HG00361 A/A 31.32628
## 4 HG00135 A/A 34.11169
## 5 NA18870 G/G 18.25141
## 6 NA11993 A/A 32.89721
```

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

```
nrow(expr)
## [1] 462
table(expr$geno)
##
## A/A A/G G/G
## 108 233 121
library(ggplot2)
ggplot(expr) + aes(geno, exp, fill = geno) +
    geom_boxplot(notch = TRUE)
```



```
ggplot_data<- ggplot_build(boxplot_data)$data[[1]]

median_values <- ggplot_data %>%
    group_by(x) %>%
    summarize(median = middle)

median_values
```