lab12part2extracredits

Kai Zhao (PID:A17599942)

##Section 4: Population Scale Analysis [HOMEWORK] One sample is obviously not enough to know what is happening in a population. You are interested in assessing genetic differences on a population scale. So, you processed about ~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.

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. How many samples do we have?

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

sample geno exp
HG00367 A/G 28.96038
NA20768 A/G 20.24449
HG00135 A/A 31.32628
HG00135 A/A 34.11169
NA18870 G/G 18.25141
NA11993 A/A 32.89721

nrow(expr)

[1] 462

summary(expr)
```

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

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

A/A A/G G/G 108 233 121

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?

Yes

```
library(ggplot2)
```

let make a boxplot

```
ggplot(expr)+aes(geno,exp,fill=geno)+geom_boxplot(notch=TRUE)
```

