# 3. Exploration Gene

Fay

2022-10-05

### Import data

```
MICE <- read.csv("output_data/MICE.csv")
```

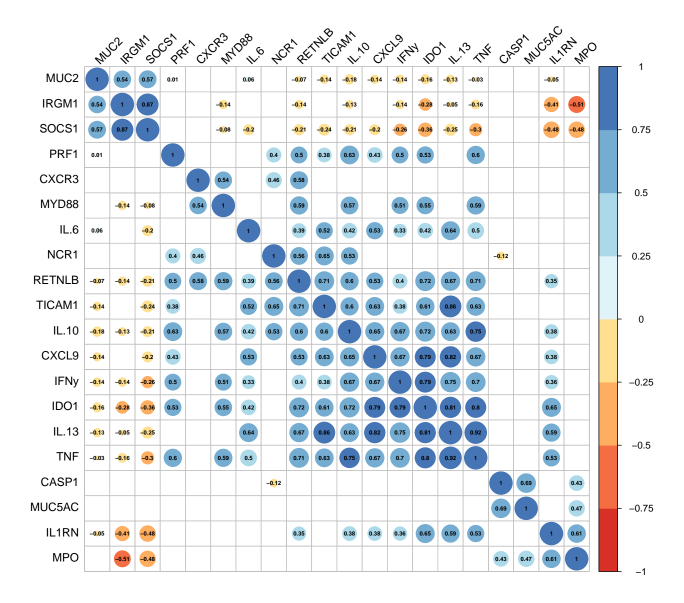
### Vectors for selecting genes

#### Lab data

Correlations between the genes

Corrplot of correlations - Laboratory gene expression data

Here is a corrplot of the correlations between the genes.

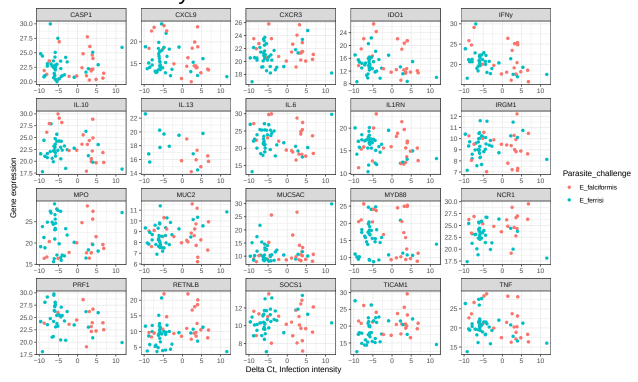


### Gene expression versus infection intensities - laboratory infections

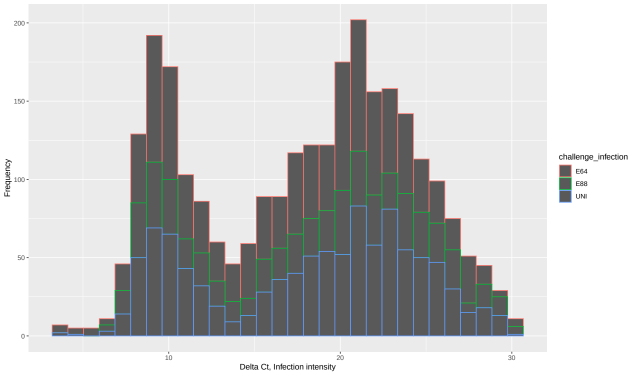
# Gene expression in response to infection group, laboratory infections



# Gene expression in response to infection intensity – laboratory infections



# Gene expression in response to infecting parasite – laboratory infections



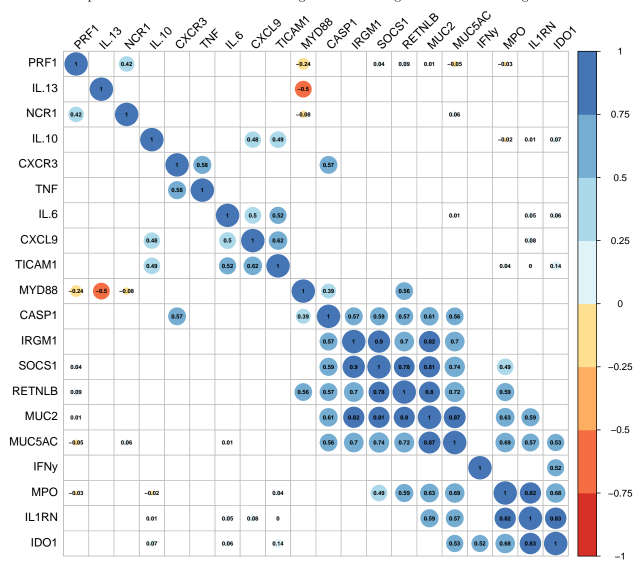
## Repeating for field infections

#### Field data

#### Correlations between the genes

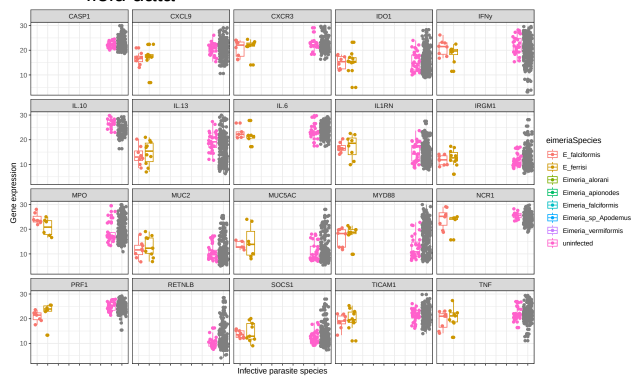
### Corrplot of correlations - Field gene expression data

Here is a corrplot of the correlations between the genes. I am using the non-normalized genes

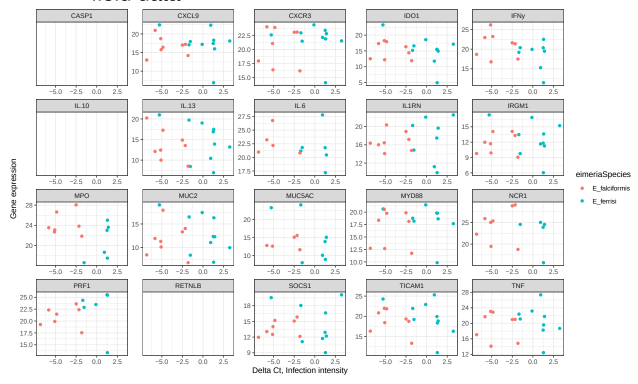


### Gene expression versus infection intensities - Field infections

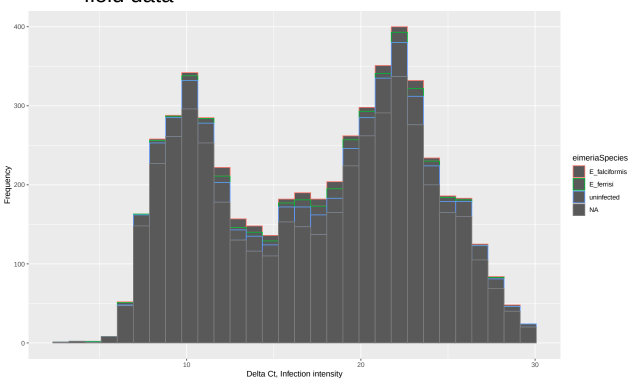
# Gene expression in response to parasite species – field data



# Gene expression in response to infection intensity – field data



# Gene expression in response to infecting parasite – field data



### Gene expression correlations in lab and field data

#### Corrplot of correlations - Laboratory and field gene expression data

Here is a corrplot of the correlations between the genes.

