## Exploratory Data Analysis

```
# Load packages and read Files
library(tidyverse)
library(data.table)
df.ZIKV <- fread("GSE207347_A1B1_vs_A2B2_ZIKV_ribodiff_name.txt.gz")</pre>
df.DENV <- fread("GSE207347_A1B1_vs_A3B3_DENV_ribodiff_name.txt.gz")</pre>
df.ZIKV_DE <- fread("GSE207347_ZIKV_DESeq2_result_name.txt.gz")</pre>
df.DENV_DE <- fread("GSE207347_DENV_DESeq2_result_name.txt.gz")</pre>
# Figure 1 B
fig1b <- ggplot(subset(df.ZIKV_DE, !is.na(padj)),</pre>
                aes(log2FoldChange, -log10(padj)))
fig1b +
  geom_point(aes(colour = -log10(padj) > -log10(0.05)), show.legend = FALSE) +
  labs(title = "ZIKV") + ylab("p adj(-log10)") +
  xlab("RNA log2Fold change \n (Infected vs uninfected)") +
  geom_vline(xintercept = 0) + geom_hline(yintercept = -log10(0.05)) +
  scale_color_manual(values = c('black', 'red')) +
  annotate("text", x = -1.5, y = 20,
           label = "Down regulated n < 0.05 \ (n=335)",
           col = "black") +
  annotate("text", x = 1.5, y = 20,
           label = "Up regulated n < 0.05 \ n (n=445)",
           col = "black") +
  theme(panel.grid.major = element_blank(),
        panel.grid.minor = element_blank(),
        panel.background = element_blank(),
        plot.title = element_text(hjust = 0.5),
        axis.line = element_line(colour = "black"),
        panel.border = element_rect(colour = "black",
                                     fill = NA, linewidth = 1.5))
```

## ZIKV

```
Down regulated
                                                                 Up regulated
   20
                                        q<0.05
                                                                     q<0.05
                                        (n=335)
                                                                     (n=445)
   15
p adj(-log10)
    5
                                 -2.5
         -5.0
                                                         0.0
                                                                                 2.5
                                          RNA log2Fold change
                                          (Infected vs uninfected)
```

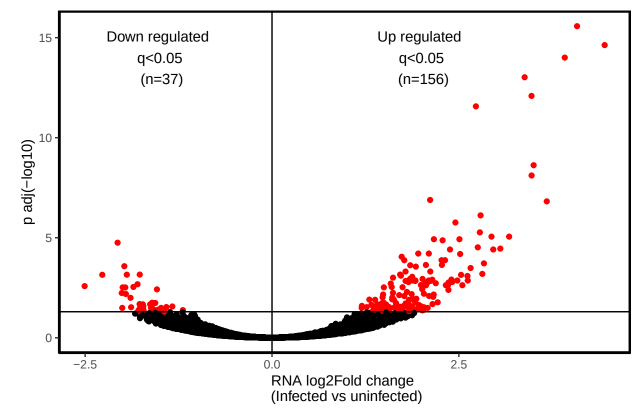
```
df.ZIKV_DE.sub <- subset(df.ZIKV_DE, !is.na(padj))
# Count of Up regulated with q < 0.05
(df.ZIKV_DE.sub$log2FoldChange > 0 & df.ZIKV_DE.sub$padj < 0.05) %>% sum()
```

## [1] 445

```
# Count of Down regulated with q < 0.05 (df.ZIKV_DE.sub$log2FoldChange < 0 & df.ZIKV_DE.sub$padj < 0.05) %>% sum()
```

## [1] 335

## **DENV**



```
df.DENV_DE.sub <- subset(df.DENV_DE, !is.na(padj))
# Count of Up regulated with q < 0.05
(df.DENV_DE.sub$log2FoldChange > 0 & df.DENV_DE.sub$padj < 0.05) %>% sum()
```

## [1] 156

```
# Count of Down regulated with q < 0.05 (df.DENV_DE.sub$log2FoldChange < 0 & df.DENV_DE.sub$padj < 0.05) %>% sum()
```

## [1] 37