## Taxonomic Analysis

## Before you begin:

These scripts were tailored for the analyses performed in:

Seibert et al, 2021, Mild and severe SARS-CoV-2 infection induces respiratory and intestinal microbiome changes in the K18-hACE2 transgenic mouse model

## Purpose:

The purpose of this script is to analyze taxonomic abundances of the different groups within the ceca and the lungs. This will correspond to Figures 3A-D, 4, 7A-D, and 8

## Load packages

```
library(phyloseq)
library(dplyr)
library(ggplot2)
library(RColorBrewer)
library(ggpubr)
library(forcats)
library(DESeq2)
library(microbiome)
library(Hmisc)
library(plyr)
```

# Import the PhyloSeq object needed for this analysis from the alpha diversity analysis

```
# Load the file RarifiedASVs.rds
Phylo.samples.final <- readRDS(file = "Phylo.samples.final.rds") # file found on github</pre>
```

#### Agglomerate at the phylum level

```
Phylum <- Phylo.samples.final %>%

tax_glom(taxrank = "Phylum") %>%

transform_sample_counts(function(x) {x/sum(x)}) %>%

# agglomerate at the Phylum level

# Transform to relative abundance

psmelt() %>%

# Melt to long format (this is for

ggplot), create dataframe

arrange(Phylum)

# Sort data frame alphabetically by class
```

#### Agglomerate at the family level

```
Family <- Phylo.samples.final %>%
   tax_glom(taxrank = "Family") %>%
   tax_glom(taxrank = "Family") %>%
        transform_sample_counts(function(x) {x/sum(x)}) %>%
        # agglomerate at the Family level
        transform to relative abundance
        psmelt() %>%
        # Melt to long format (this is for
        ggplot), create dataframe
        arrange(Family)
        # Sort data frame alphabetically by class
```

## Figure 3A: Taxonomic boxplots of the cecum at the Phylum level

Filter the Phylum into the cecum with the 3 groups that we will analyze

```
# Filter the samples so that you only have cecum samples and only PBS, 3 and 5 infected vehicele
Phylum.cecum <- Phylum %>%
  filter(SampleType == "Cecum") %>%
  filter(Group == "PBS" | Group == "Infected-3-Vehicle" | Group == "Infected-5-Vehicle")
```

First, lets plot all of the Phyla on a single bar graph.

```
Phylum.cecum.box <- Phylum.cecum
# Change phylum to a factor so I can change the order
Phylum.cecum.box$Phylum <- as.factor(Phylum.cecum.box$Phylum)</pre>
```

```
# Multiply the Abundance by 100 to show an easier percentage
Phylum.cecum.box$Abundance <- Phylum.cecum.box$Abundance*100

# Reorder the groups
Phylum.cecum.box$Group<- factor(Phylum.cecum.box$Group, levels = c('PBS', 'Infected-3-Vehicle', 'Infected-5-Vehicle'))

# Set the colors for the different groups
colorgroups = c("blue", "darkorange2", "black")

# Plot using ggplot using boxplots
ggplot(data=Phylum.cecum.box, aes(x=fct_rev(Phylum), y=Abundance, color=fct_rev(Group))) +
geom_boxplot()+
scale_color_manual(values=colorgroups)+
coord_flip()+
scale_y_continuous(breaks = seq(0, 100, by=25), limits=c(0, 100))+
theme_bw()+
theme(panel.grid = element_blank())</pre>
```

Since there are many phyla that have extremely low abundance, we are going to group all of the taxa that have less than 0.1% relative abundance since these taxa are very rare and we are more interested in the more prominant taxa to see trends

```
# Filter for only the most abundant phylum. Adding too many phyla will cause the graph to look muddled
Phylum.cecum.box <- Phylum.cecum
# Group taxa that have overall abundance of less than 0.1%
Phylum.cecum.box$Phylum[Phylum.cecum.box$Abundance < 0.01] <- "Taxa < 1% abund."
# Change phylum to a factor so I can change the order
Phylum.cecum.box$Phylum <- as.factor(Phylum.cecum.box$Phylum)</pre>
# Multiply the Abundance by 100 to show an easier percentage
Phylum.cecum.box$Abundance <- Phylum.cecum.box$Abundance*100
# Reorder the groups
Phylum.cecum.box$Group<- factor(Phylum.cecum.box$Group, levels = c('PBS', 'Infected-3-Vehicle',
'Infected-5-Vehicle'))
# Set the colors for the different groups
colorgroups = c("blue", "darkorange2", "black")
# Plot using ggplot using boxplots
ggplot(data=Phylum.cecum.box, aes(x=fct_rev(Phylum), y=Abundance, color=fct_rev(Group))) +
  geom_boxplot()+
  scale_color_manual(values=colorgroups)+
  coord_flip()+
  scale_y_continuous(breaks = seq(0, 100, by=25), limits=c(0, 100))+
  theme bw()+
 theme(panel.grid = element blank())
```

Since it is harder to see the less abundant taxa when all phyla are plotted in the same graph, lets split the phyla into 2 graphs.

First, lets calculate the abundances, plot the boxplots and perform statistical analysis for the more abundant phyla

```
# Filter for only the most abundant phylum. Adding too many phyla will cause the graph to look muddled
Phylum.cecum.box <- Phylum.cecum %>%
    filter(Phylum == "Bacteroidota" | Phylum == "Firmicutes" | Phylum == "Verrucomicrobiota")

# Change phylum to a factor so I can change the order
Phylum.cecum.box$Phylum <- as.factor(Phylum.cecum.box$Phylum)

# Multiply the Abundance by 100 to show an easier percentage
Phylum.cecum.box$Abundance <- Phylum.cecum.box$Abundance*100

# Reorder the groups
Phylum.cecum.box$Group<- factor(Phylum.cecum.box$Group, levels = c('PBS', 'Infected-3-Vehicle', 'Infected-5-Vehicle'))

# Set the colors for the different groups
colorgroups = c("blue", "darkorange2", "black")

# Plot using ggplot using boxplots
ggplot(data=Phylum.cecum.box, aes(x=fct_rev(Phylum), y=Abundance, color=fct_rev(Group))) +
geom_boxplot()+</pre>
```

```
scale_color_manual(values=colorgroups)+
  coord flip()+
  scale y continuous(breaks = seq(0, 100, by=25), limits=c(0, 100))+
  theme bw()+
  theme(panel.grid = element_blank())
\# Statistical Comparison using wilcox test using specific bacteria. I need to manually enter the
bacteria name. It gets too confusing when i add multiple lines with all phyla
Phylum.cecum.box.Special <- Phylum.cecum.box %>%
  filter(Phylum == "Bacteroidota")
compare_means(Abundance ~ Group, data = Phylum.cecum.box.Special, method = "wilcox.test")
Phylum.cecum.box.Special <- Phylum.cecum.box %>%
  filter(Phylum == "Firmicutes")
compare_means(Abundance ~ Group, data = Phylum.cecum.box.Special, method = "wilcox.test")
Phylum.cecum.box.Special <- Phylum.cecum.box %>%
  filter(Phylum == "Verrucomicrobiota")
compare_means(Abundance ~ Group, data = Phylum.cecum.box.Special, method = "wilcox.test")
```

Next, lets calculate the abundances, plot the boxplots and perform statistical analysis for the less abundant phyla

```
# Filter for only the most abundant phylum. Adding too many phyla will cause the graph to look muddled
Phylum.cecum.box <- Phylum.cecum %>%
  filter(Phylum == "Actinobacteriota" | Phylum == "Proteobacteria")
\# Change phylum to a factor so I can change the order
Phylum.cecum.box$Phylum <- as.factor(Phylum.cecum.box$Phylum)
# Multiply the Abundance by 100 to show an easier percentage
Phylum.cecum.box$Abundance <- Phylum.cecum.box$Abundance*100
# Reorder the groups
Phylum.cecum.box$Group<- factor(Phylum.cecum.box$Group, levels = c('PBS', 'Infected-3-Vehicle',
'Infected-5-Vehicle'))
# Set the colors for the different groups
colorgroups = c("blue", "darkorange2", "black")
# Plot using ggplot using boxplots
ggplot(data=Phylum.cecum.box, aes(x=fct_rev(Phylum), y=Abundance, color=fct_rev(Group))) +
  geom_boxplot()+
  scale_color_manual(values=colorgroups)+
  coord_flip()+
  scale_y_continuous(breaks = seq(0, 3, by=1), limits=c(0, 3))+
  theme bw()+
 theme(panel.grid = element_blank())
\# Statistical Comparison using wilcox test using specific bacteria. I need to manually enter the
bacteria name. It gets too confusing when i add multiple lines with all phyla
Phylum.cecum.box.Special <- Phylum.cecum.box %>%
  filter(Phylum == "Actinobacteriota")
compare_means(Abundance ~ Group, data = Phylum.cecum.box.Special, method = "wilcox.test")
Phylum.cecum.box.Special <- Phylum.cecum.box %>%
  filter(Phylum == "Proteobacteria")
compare_means(Abundance ~ Group, data = Phylum.cecum.box.Special, method = "wilcox.test")
```

## Figure 3B: Firmicutes/Bacteroidetes ratio in the cecum

Lets analyze the Firmicutes/Bacteroidetes ratio as this can be use as an indicator of dysbiosis.

I tested calculating the ratio with raw counts and with relative abundances and the results were very similar. This script is for calculating the ration using relative abundances

```
# Create a data frame that only has the relative abundances of those classified as Firmicutes
Phylum.cecum_RA_Firm <- Phylum.cecum %>%
    filter(Phylum == "Firmicutes")

# Remove access columns that are not needed
# I will keep Abundance, SampleID, MouseID, Group and dpi
Phylum.cecum_RA_Firm.edit <- Phylum.cecum_RA_Firm[,c(3,4,8,9,10)]

# Change the first column header aka Abundance to Firmicutes
colnames(Phylum.cecum_RA_Firm.edit)[1] <- "Firmicutes"</pre>
```

```
# Export the data frame as a csv file
write.csv(Phylum.cecum RA Firm.edit, "Phylum.cecum RA Firm.csv")
# Create a data frame that only has the relative abundances of those classified as Bacteroidota
Phylum.cecum_RA_Bact <- Phylum.cecum %>%
  filter(Phylum == "Bacteroidota")
# Remove access columns that are not needed
# I will keep Abundance, SampleID, MouseID, Group and dpi
Phylum.cecum_RA_Bact.edit <- Phylum.cecum_RA_Bact[,c(3,4,8,9,10)]
# Change the first column header aka Abundance to Bacteroidota
colnames(Phylum.cecum_RA_Bact.edit)[1] <- "Bacteroidota"</pre>
# Export the data frame as a csv file
write.csv(Phylum.cecum RA Bact.edit, "Phylum.cecum RA Bact.csv")
# I calculated the ratios of the Firmicutes/Bacteroidota in excel and then imported it back into R
Phylum.cecum_RA_FBTOTAL <- read.csv("Phylum.cecum_RA_FBRatio.csv")
# Make Group a factor and relevel
Phylum.cecum_RA_FBTOTAL$Group <- factor(Phylum.cecum_RA_FBTOTAL$Group, levels = c("PBS", "Infected-3-
Vehicle", "Infected-5-Vehicle"))
# Set the colors for the different groups
colorgroups = c("black", "darkorange2", "blue")
# Plot the data with ggplot
ggplot(data=Phylum.cecum_RA_FBTOTAL, aes(x=FB_Ratio, y=Group, color=Group)) +
  geom_boxplot()+
   theme bw()+
  scale_color_manual(values=colorgroups)+
  scale_x_continuous(breaks = seq(0, 4, by=2), limits=c(0, 4))+
  coord_flip()+
  theme(panel.grid = element_blank())
# Statistical Comparison using wilcox test using specific bacteria. I need to manually enter the
bacteria name. It gets too confusing when i add multiple lines with all phyla
compare means(FB Ratio ~ Group, data = Phylum.cecum RA FBTOTAL, method = "wilcox.test")
```

## Figure 3C: Taxonomic boxplots of the cecum at the Family level

Filter the Family into the cecum with the 3 groups that we will analyze. We will group the taxa that have less than 1% relative abundance since we are interested in overall trends and those sequences that have extremely small abundances could be sequencing artifacts.

```
# Filter the samples so that you only have cecum samples and only PBS, 3 and 5 infected vehicele
Family.cecum <- Family %>%
  filter(SampleType == "Cecum") %>%
  filter(Group == "PBS" | Group == "Infected-3-Vehicle" | Group == "Infected-5-Vehicle")
```

Lets look at all of the families in the cecum

```
# Filter for the most abundant families
Family.cecum.box <- Family.cecum
# Change Family to a factor so i can change the order
Family.cecum.box$Family <- as.factor(Family.cecum.box$Family)</pre>
# Multiply the Abundance by 100 to show an easier percentage
Family.cecum.box$Abundance <- Family.cecum.box$Abundance*100
# Make Group a factor and relevel
Family.cecum.box$Group <- factor(Family.cecum.box$Group, levels = c("PBS", "Infected-3-Vehicle",
"Infected-5-Vehicle"))
# Set the colors for the different groups
colorgroups = c("blue", "darkorange2", "black")
# Plot the graph using ggplot
ggplot(data=Family.cecum.box, aes(x=fct rev(Family), y=Abundance, color=fct rev(Group))) +
 geom boxplot()+
  scale_color_manual(values=colorgroups)+
 coord flip()+
  scale_y_continuous(breaks = seq(0, 100, by=25), limits=c(0, 100))+
```

Since the abundances of all of the very rare families are making the trends of the bacterial families that are prominant hard to see, we are going to group the families that have less than 1% relative abundance in a single sample.

```
# I only want the families that have at least 1% relative abundance in a sample
Family.cecum_greater_1_abundance <- Family.cecum.box %>% filter(Family.cecum.box[,3] >= 1.0) #this is 1 since we already multiplied the abundances by 100 to
convert into %
unique(Family.cecum_greater_1_abundance$Family)
# Change phylum to a factor so I can change the order
Family.cecum_greater_1_abundance$Family <- as.factor(Family.cecum_greater_1_abundance$Family)
# Reorder the groups
Family.cecum_greater_1_abundance$Group<- factor(Family.cecum_greater_1_abundance$Group, levels =
c('PBS', 'Infected-3-Vehicle', 'Infected-5-Vehicle'))
# Set the colors for the different groups
colorgroups = c("blue", "darkorange2", "black")
# Plot using ggplot using boxplots
ggplot(data=Family.cecum_greater_1_abundance, aes(x=fct_rev(Family), y=Abundance,
color=fct_rev(Group))) +
  geom boxplot()+
  scale color manual(values=colorgroups)+
  coord flip()+
  scale_y_continuous(breaks = seq(0, 100, by=25), limits=c(0, 100))+
  theme bw()+
  theme(panel.grid = element_blank())
```

All families appeared in more than 1 sample so they were not filtered out.

Next, lets graph the most abundant families separately so that we can have different y axis for the less abundant and see the relationships among rare taxa more clear.

```
# Filter for the most abundant families
Family.cecum.box <- Family.cecum %>%
  filter(Family == "Lachnospiraceae" | Family == "Muribaculaceae" | Family == "Akkermansiaceae" |
Family == "Bacteroidaceae" | Family == "Erysipelotrichaceae" | Family == "Oscillospiraceae")
# Relevel the families to be alphabetical
Family.cecum.box$Family <- factor(Family.cecum.box$Family, levels = c("Bacteroidaceae",
"Muribaculaceae", "Erysipelotrichaceae", "Lachnospiraceae", "Oscillospiraceae", "Akkermansiaceae"))
# Change Family to a factor so i can change the order
Family.cecum.box$Family <- as.factor(Family.cecum.box$Family)</pre>
# Multiply the Abundance by 100 to show an easier percentage
Family.cecum.box$Abundance <- Family.cecum.box$Abundance*100
# Make Group a factor and relevel
Family.cecum.box$Group <- factor(Family.cecum.box$Group, levels = c("PBS", "Infected-3-Vehicle",
"Infected-5-Vehicle"))
# Set the colors for the different groups
colorgroups = c("blue", "darkorange2", "black")
# Plot the graph using ggplot
ggplot(data=Family.cecum.box, aes(x=fct_rev(Family), y=Abundance, color=fct_rev(Group))) +
  geom boxplot()+
  scale_color_manual(values=colorgroups)+
  coord flip()+
  scale y continuous(breaks = seq(0, 100, by=25), limits=c(0, 100))+
  theme bw()+
  theme(panel.grid = element_blank())
# Statistical Comparison using wilcox test using specific bacteria. I need to manually enter the
```

```
bacteria name. It gets too confusing when i add multiple lines with all abundant families
Family.cecum.box.Special <- Family.cecum.box %>%
 filter(Family == "Bacteroidaceae")
compare_means(Abundance ~ Group, data = Family.cecum.box.Special, method = "wilcox.test")
Family.cecum.box.Special <- Family.cecum.box %>%
  filter(Family == "Muribaculaceae")
compare means (Abundance ~ Group, data = Family.cecum.box.Special, method = "wilcox.test")
Family.cecum.box.Special <- Family.cecum.box %>%
filter(Family == "Erysipelotrichaceae")
compare_means(Abundance ~ Group, data = Family.cecum.box.Special, method = "wilcox.test")
Family.cecum.box.Special <- Family.cecum.box %>%
 filter(Family == "Lachnospiraceae")
compare means (Abundance ~ Group, data = Family.cecum.box.Special, method = "wilcox.test")
Family.cecum.box.Special <- Family.cecum.box %>%
 filter(Family == "Oscillospiraceae")
compare_means(Abundance ~ Group, data = Family.cecum.box.Special, method = "wilcox.test")
Family.cecum.box.Special <- Family.cecum.box %>%
  filter(Family == "Akkermansiaceae")
compare means (Abundance ~ Group, data = Family.cecum.box.Special, method = "wilcox.test")
After, lets graph the less abundant families in their own graph so the trends are easier to view
in the graph
# Filter for the less abundant families
Family.cecum.less.box <- Family.cecum %>%
filter(Family == "Acholeplasmataceae" | Family == "Clostridiaceae" | Family == "Lactobacillaceae" | Family == "Monoglobaceae" | Family == "Ruminococcaceae" | Family == "Sutterellaceae")
# Relevel the families so that they are grouped by phylum
Family.cecum.less.box$Family <- factor(Family.cecum.less.box$Family, levels = c("Acholeplasmataceae",
"Clostridiaceae","Lactobacillaceae", "Monoglobaceae", "Ruminococcaceae","Sutterellaceae"))
# Change family to a factor so I can change the order
Family.cecum.less.box$Family <- as.factor(Family.cecum.less.box$Family)</pre>
# Multiply the Abundance by 100 to show an easier percentage
Family.cecum.less.box$Abundance <- as.numeric(Family.cecum.less.box$Abundance)
Family.cecum.less.box$Abundance <- Family.cecum.less.box$Abundance*100
# Make Group a factor and relevel
Family.cecum.less.box$Group <- factor(Family.cecum.less.box$Group, levels = c("PBS", "Infected-3-
Vehicle", "Infected-5-Vehicle"))
# Set the colors for the different groups
colorgroups = c("blue", "darkorange2", "black")
# Plot the graph using ggplot
ggplot(data=Family.cecum.less.box, aes(x=fct rev(Family), y=Abundance, color=fct rev(Group))) +
  geom boxplot()+
  scale_color_manual(values=colorgroups)+
  coord flip()+
  scale_y_continuous(breaks = seq(0, 12, by=2), limits=c(0, 12))+
  theme bw()+
  theme(panel.grid = element_blank())
\# Statistical Comparison using wilcox test using specific bacteria. I need to manually enter the
bacteria name. It gets too confusing when i add multiple lines with all less abundant families
Family.cecum.less.box.Special <- Family.cecum.less.box %>%
filter(Family == "Acholeplasmataceae")
compare_means(Abundance ~ Group, data = Family.cecum.less.box.Special, method = "wilcox.test")
Family.cecum.less.box.Special <- Family.cecum.less.box %>%
 filter(Family == "Clostridiaceae")
compare_means(Abundance ~ Group, data = Family.cecum.less.box.Special, method = "wilcox.test")
Family.cecum.less.box.Special <- Family.cecum.less.box %>%
  filter(Family == "Lactobacillaceae")
compare means (Abundance ~ Group, data = Family.cecum.less.box.Special, method = "wilcox.test")
Family.cecum.less.box.Special <- Family.cecum.less.box %>%
  filter(Family == "Monoglobaceae")
compare_means(Abundance ~ Group, data = Family.cecum.less.box.Special, method = "wilcox.test")
```

```
Family.cecum.less.box.Special <- Family.cecum.less.box %>%
    filter(Family == "Ruminococcaceae")
compare_means(Abundance ~ Group, data = Family.cecum.less.box.Special, method = "wilcox.test")

Family.cecum.less.box.Special <- Family.cecum.less.box %>%
    filter(Family == "Sutterellaceae")
compare_means(Abundance ~ Group, data = Family.cecum.less.box.Special, method = "wilcox.test")
```

#### Figure 3D: Correlation analysis of the cecum at the Family level

We will now perform a correlation analysis when analyzing the factors of infection to bacterial families

First, lets create a function to flatten a matrix as this will be needed for the correlation matrix

```
flattenCorrMatrix <- function(cormat, pmat) {
  ut <- upper.tri(cormat)
  data.frame(
    row = rownames(cormat)[row(cormat)[ut]],
    column = rownames(cormat)[col(cormat)[ut]],
    cor =(cormat)[ut],
    p = pmat[ut]
  )
}</pre>
```

Next, we will perform the correlation analysis using rcorr from the package Hmisc

```
\# Filter the samples so that you only have cecum samples and only PBS, 3 and 5 infected vehicle
cecum_added_factors <- read.csv("Added_factors_cecum.csv") # File uploaded on github</pre>
Cecum total factors <- merge(Family.cecum, cecum added factors, by = "SampleID")
Cecum_total_factors_edited <- Cecum_total_factors[,-</pre>
c(1,2,3,5,6,7,8,9,11,12,13,14,15,16,17,18,19,21,23)]
#Acholeplasmataceae
Cecum_total_factors_edited_filter <- Cecum_total_factors_edited %>%
  filter(Cecum_total_factors_edited[,3] == "Acholeplasmataceae")
Cecum total factors edited filter <- as.matrix(Cecum total factors edited filter[,-3])
res <- rcorr(Cecum total_factors_edited_filter, type =</pre>
                                                             "spearman")
results <- flattenCorrMatrix(res$r, res$P)</pre>
resultsfilter <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter$row <- revalue(resultsfilter$row, c("Abundance"="Acholeplasmataceae"))
#Akkermansiaceae
Cecum_total_factors_edited_filter <- Cecum_total_factors_edited %>%
    filter(Cecum_total_factors_edited[,3] == "Akkermansiaceae")
Cecum_total_factors_edited_filter <- as.matrix(Cecum_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Cecum_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter2 <- results %>%
filter(results[,1] == "Abundance")
resultsfilter2$row <- revalue(resultsfilter2$row, c("Abundance"="Akkermansiaceae"))</pre>
resultsfilter_final <- rbind(resultsfilter, resultsfilter2)</pre>
#Bacteroidaceae
Cecum_total_factors_edited_filter <- Cecum_total_factors_edited %>%
  filter(Cecum_total_factors_edited[,3] == "Bacteroidaceae")
Cecum total factors edited filter <- as.matrix(Cecum total factors edited filter[,-3])
res <- rcorr(Cecum_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter2$row <- revalue(resultsfilter2$row, c("Abundance"="Bacteroidaceae"))</pre>
resultsfilter_final <- rbind(resultsfilter_final, resultsfilter2)</pre>
Cecum_total_factors_edited_filter <- Cecum_total_factors_edited %>%
  filter(Cecum_total_factors_edited[,3] == "Clostridiaceae")
Cecum_total_factors_edited_filter <- as.matrix(Cecum_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Cecum_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)</pre>
resultsfilter2 <- results %>%
 filter(results[,1] == "Abundance")
```

```
resultsfilter2$row <- revalue(resultsfilter2$row, c("Abundance"="Clostridiaceae"))
resultsfilter final <- rbind(resultsfilter final, resultsfilter2)</pre>
#Ervsipelotrichaceae
Cecum_total_factors_edited_filter <- Cecum_total_factors_edited %>%
  filter(Cecum_total_factors_edited[,3] == "Erysipelotrichaceae")
Cecum_total_factors_edited_filter <- as.matrix(Cecum_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Cecum total factors edited filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter2 <- results %>%
filter(results[,1] == "Abundance")
resultsfilter2$row <- revalue(resultsfilter2$row, c("Abundance"="Erysipelotrichaceae"))</pre>
resultsfilter_final <- rbind(resultsfilter_final, resultsfilter2)</pre>
Cecum total factors edited filter <- Cecum total factors edited %>%
 filter(Cecum_total_factors_edited[,3] == "Lachnospiraceae")
Cecum total factors edited filter <- as.matrix(Cecum total factors edited filter[,-3])
res <- rcorr(Cecum_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)</pre>
resultsfilter2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter2$row <- revalue(resultsfilter2$row, c("Abundance"="Lachnospiraceae"))
resultsfilter final <- rbind(resultsfilter final, resultsfilter2)</pre>
#Lactobacillaceae
Cecum_total_factors_edited_filter <- Cecum_total_factors_edited %>%
  filter(Cecum_total_factors_edited[,3] == "Lactobacillaceae")
Cecum_total_factors_edited_filter <- as.matrix(Cecum_total_factors_edited_filter[,-3])
res <- rcorr(Cecum_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter2$row <- revalue(resultsfilter2$row, c("Abundance"="Lactobacillaceae"))</pre>
resultsfilter_final <- rbind(resultsfilter_final, resultsfilter2)</pre>
Cecum total factors edited filter <- Cecum total factors edited %>%
  filter(Cecum_total_factors_edited[,3] == "Monoglobaceae")
Cecum_total_factors_edited_filter <- as.matrix(Cecum_total_factors_edited_filter[,-3])
res <- rcorr(Cecum_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)</pre>
resultsfilter2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter2$row <- revalue(resultsfilter2$row, c("Abundance"="Monoglobaceae"))
resultsfilter final <- rbind(resultsfilter final, resultsfilter2)</pre>
#Muribaculaceae
Cecum total_factors_edited_filter <- Cecum_total_factors_edited %>%
  filter(Cecum_total_factors_edited[,3] == "Muribaculaceae")
Cecum_total_factors_edited_filter <- as.matrix(Cecum_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Cecum_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)</pre>
resultsfilter2 <- results %>%
filter(results[,1] == "Abundance")
resultsfilter2$row <- revalue(resultsfilter2$row, c("Abundance"="Muribaculaceae"))</pre>
resultsfilter_final <- rbind(resultsfilter_final, resultsfilter2)</pre>
Cecum_total_factors_edited_filter <- Cecum_total_factors_edited %>%
  filter(Cecum_total_factors_edited[,3] == "Oscillospiraceae")
Cecum total factors edited filter <- as.matrix(Cecum total factors edited filter[,-3])
res <- rcorr(Cecum_total_factors_edited_filter, type = "spearman")</pre>
results <- flattenCorrMatrix(res$r, res$P)</pre>
resultsfilter2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter2$row <- revalue(resultsfilter2$row, c("Abundance"="Oscillospiraceae"))
resultsfilter final <- rbind(resultsfilter final, resultsfilter2)</pre>
#Ruminococcaceae
Cecum_total_factors_edited_filter <- Cecum_total_factors_edited %>%
  filter(Cecum_total_factors_edited[,3] == "Ruminococcaceae")
Cecum_total_factors_edited_filter <- as.matrix(Cecum_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Cecum_total_factors_edited_filter, type = "spearman")</pre>
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter2 <- results %>%
 filter(results[,1] == "Abundance")
resultsfilter2$row <- revalue(resultsfilter2$row, c("Abundance"="Ruminococcaceae"))</pre>
```

```
resultsfilter_final <- rbind(resultsfilter_final, resultsfilter2)</pre>
#Sutterellaceae
Cecum total_factors_edited_filter <- Cecum_total_factors_edited %>%
  filter(Cecum_total_factors_edited[,3] == "Sutterellaceae")
Cecum_total_factors_edited_filter <- as.matrix(Cecum_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Cecum_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter2$row <- revalue(resultsfilter2$row, c("Abundance"="Sutterellaceae"))
resultsfilter_final <- rbind(resultsfilter_final, resultsfilter2)
resultsfilter_final$row <- factor(resultsfilter_final$row, levels = c("Bacteroidaceae", "Muribaculaceae", "Acholeplasmataceae", "Clostridiaceae", "Erysipelotrichaceae", "Lacthospiraceae", "Lactobacillaceae", "Monoglobaceae", "Oscillospiraceae", "Ruminococcaceae", "Sutterellaceae",
"Akkermansiaceae"))
resultsfilter_final$column <- factor(resultsfilter_final$column, levels = c("Inoculum", "dpi.x",
"Acitivity", "Viral.NT.Titer", "Viral.Lung.Titer", "Viral.Brain.Titer"))</pre>
ggplot(resultsfilter_final, aes(column, fct_rev(row))) +
  geom point(aes(colour = cor),
                         size = 8) +
  scale\_color\_gradient2(limits = c(-0.7, 0.7), breaks = c(-0.6, -0.3, 0, 0.3, 0.6),
                              low = "royalblue1",
                              mid = "brown3",
                              high = "yellow")+
  theme bw()
# Produce a table with those correlation results that have are significant or have a p value < 0.05
resultsfilter_final_significant <- resultsfilter_final %>%
 filter(resultsfilter_final[,4] < 0.05)</pre>
```

The figure was further edited in Adobe Illustrator to indicate the relationships that were considered statistically significant.

## Figure 4: Taxonomic barplot of the cecum at the Family level

Now lets produce a graph for the taxonomic barplot

```
# Combine the Taxa that are less than 1% relative abundance together
Family.cecum.barplot <- Family.cecum
Family.cecum.barplot$Family[Family.cecum.barplot$Abundance < 0.01] <- "Taxa < 1% abund."
ColorTest = c("gray31", "yellow", "forestgreen", "midnightblue", "blue", "steelblue3", "lightblue2",
"royalblue1", "cyan", "mediumblue", "deepskyblue1", "darkorchid4", "orchid1")
Family.cecum.barplot$MouseID <- factor(Family.cecum.barplot$MouseID, levels = c("5", "6", "23", "24", "25", "34", "35", "36", "3", "4", "20", "21", "22", "31", "32", "33", "1", "2", "15", "17", "19"))
Family.cecum.barplot$Family <- factor(Family.cecum.barplot$Family, levels = c("Muribaculaceae",
"Bacteroidaceae", "Lachnospiraceae", "Oscillospiraceae", "Erysipelotrichaceae", "Acholeplasmataceae", "Lactbacillaceae", "Ruminococcaceae", "Clostridiaceae", "Monoglobaceae", "Sutterellaceae", "Akkermansiaceae", "Taxa < 1% abund."))
#plot using ggplot
ggplot(data=Family.cecum.barplot, aes(x=MouseID, y=Abundance, fill=fct_rev(Family))) +
  geom_bar(aes(), colour = "black", stat="identity", position="stack", width = 0.8) +
   scale_fill_manual(values=ColorTest)+
  facet_grid(~dpi, scales = "free", space = "free")+
theme(legend.position="bottom") +
   guides(fill=guide_legend(nrow=5))+
   theme(legend.position="right") + guides(fill=guide_legend(ncol =1))+
   theme_bw()+
   theme(panel.grid = element_blank())+
  theme(strip.text = element text(size=11, face="bold"))
```

The figure is then combined with activity score and viral titers in Adobe Illustrator

## Figure 7A: Taxonomic boxplots of the lung at the Phylum level

Filter the Phylum into the cecum with the 3 groups that we will analyze

```
# Filter the samples so that you only have cecum samples and only PBS, 3 and 5 infected vehicele
Phylum.lung.GC376 <- Phylum %>%
    filter(SampleType == "Lung") %>%
    filter(Group == "Mock-GC376" | Group == "Infected-3-GC376" | Group == "Infected-5-GC376")

First, lets plot all of the Phyla on a single bar graph.

Phylum.lung.box <- Phylum.lung.GC376

# Change phylum to a factor so I can change the order</pre>
```

```
# Change phylum to a factor so I can change the order
Phylum.lung.box$Phylum <- as.factor(Phylum.lung.box$Phylum)

# Multiply the Abundance by 100 to show an easier percentage
Phylum.lung.box$Abundance <- Phylum.lung.box$Abundance*100

# Reorder the groups
Phylum.lung.box$Group<- factor(Phylum.lung.box$Group, levels = c("Mock-GC376", "Infected-3-GC376",
    "Infected-5-GC376"))

# Set the colors for the different groups
colorgroups = c("maroon1", "forestgreen", "chocolate4")

# Plot using ggplot using boxplots
ggplot(data=Phylum.lung.box, aes(x=fct_rev(Phylum), y=Abundance, color=fct_rev(Group))) +
geom_boxplot()+
scale_color_manual(values=colorgroups)+
coord_flip()+
scale_y_continuous(breaks = seq(0, 100, by=25), limits=c(0, 100))+
theme_bw()+
theme_pw()+
theme(panel.grid = element_blank())</pre>
```

Like the cecum, there are many phyla that have extremely low abundance, we are going to group all of the taxa that have less than 0.1% relative abundance since these taxa are very rare and we are more interested in the more prominant taxa to see trends

```
# Filter for only the most abundant phylum as seen in the bargraph above. Adding too many phyla will
cause the graph to look muddled
Phylum.lung.GC376.box <- Phylum.lung.GC376
# Group taxa that have overall abundance of less than 1%
Phylum.lung.GC376.box$Phylum[Phylum.lung.GC376.box$Abundance < 0.01] <- "Taxa < 1% abund."
# Change phylum to a factor so i can change the order
Phylum.lung.GC376.box$Phylum <- as.factor(Phylum.lung.GC376.box$Phylum)
# Multiply the Abundance by 100 to show an easier percentage
Phylum.lung.GC376.box$Abundance <- Phylum.lung.GC376.box$Abundance*100
# Relevel the groups
Phylum.lung.GC376.box$Group <- factor(Phylum.lung.GC376.box$Group, levels = c("Mock-GC376", "Infected-
3-GC376", "Infected-5-GC376"))
# Set the colors for the different groups
colorgroups = c("maroon1", "forestgreen", "chocolate4")
# Graph the plot using ggplot
ggplot(data=Phylum.lung.GC376.box, aes(x=fct_rev(Phylum), y=Abundance, color=fct_rev(Group))) +
  geom boxplot()+
  scale_color_manual(values=colorgroups)+
  coord_flip()+
  scale_y_continuous(breaks = seq(0, 100, by=25), limits=c(0, 100))+
  theme_bw()+
theme(panel.grid = element_blank())
```

Since it is harder to see the less abundant taxa when all phyla are plotted in the same graph, lets split the phyla into 2 graphs.

First, lets calculate the abundances, plot the boxplots and perform statistical analysis for the more abundant phyla

```
# Filter for only the most abundant phylum as seen in the bargraph above. Adding too many phyla will
cause the graph to look muddled
Phylum.lung.GC376.box <- Phylum.lung.GC376 %>%
  filter(Phylum == "Firmicutes" | Phylum == "Bacteroidota" | Phylum == "Proteobacteria")
# Change phylum to a factor so i can change the order
Phylum.lung.GC376.box$Phylum <- as.factor(Phylum.lung.GC376.box$Phylum)
# Multiply the Abundance by 100 to show an easier percentage
Phylum.lung.GC376.box$Abundance <- Phylum.lung.GC376.box$Abundance*100
# Relevel the groups
Phylum.lung.GC376.box$Group <- factor(Phylum.lung.GC376.box$Group, levels = c("Mock-GC376", "Infected-
          "Infected-5-GC376"))
3-GC376",
# Set the colors for the different groups
colorgroups = c("maroon1", "forestgreen", "chocolate4")
# Graph the plot using ggplot
ggplot(data=Phylum.lung.GC376.box, aes(x=fct_rev(Phylum), y=Abundance, color=fct_rev(Group))) +
  geom_boxplot()+
  scale color manual(values=colorgroups)+
  coord_flip()+
  scale y continuous(breaks = seq(0, 100, by=25), limits=c(0, 100))+
  theme bw()+
  theme(panel.grid = element blank())
# Statistical Comparison using wilcox test using specific bacteria. I need to manually enter the
bacteria name. It gets too confusing when i add multiple lines with all phyla
Phylum.lung.GC376.box.Special <- Phylum.lung.GC376.box %>%
  filter(Phylum == "Firmicutes")
compare means(Abundance ~ Group, data = Phylum.lung.GC376.box.Special, method = "wilcox.test")
Phylum.lung.GC376.box.Special <- Phylum.lung.GC376.box %>%
 filter(Phylum == "Bacteroidota")
compare_means(Abundance ~ Group, data = Phylum.lung.GC376.box.Special, method = "wilcox.test")
Phylum.lung.GC376.box.Special <- Phylum.lung.GC376.box %>%
  filter(Phylum == "Proteobacteria")
compare means(Abundance ~ Group, data = Phylum.lung.GC376.box.Special, method = "wilcox.test")
Next, lets calculate the abundances, plot the boxplots and perform statistical analysis for the less
abundant phyla
# Filter for only the most abundant phylum as seen in the bargraph above. Adding too many phyla will
cause the graph to look muddled
Phylum.lung.GC376.box <- Phylum.lung.GC376 %>%
 filter(Phylum == "Actinobacteriota" | Phylum == "Deinococcota" | Phylum == "Verrucomicrobiota")
# Change phylum to a factor so i can change the order
Phylum.lung.GC376.box$Phylum <- as.factor(Phylum.lung.GC376.box$Phylum)
# Multiply the Abundance by 100 to show an easier percentage
Phylum.lung.GC376.box$Abundance <- Phylum.lung.GC376.box$Abundance*100
# Relevel the groups
Phylum.lung.GC376.box$Group <- factor(Phylum.lung.GC376.box$Group, levels = c("Mock-GC376", "Infected-
3-GC376", "Infected-5-GC376"))
# Set the colors for the different groups
colorgroups = c("maroon1", "forestgreen", "chocolate4")
# Graph the plot using ggplot
ggplot(data=Phylum.lung.GC376.box, aes(x=fct_rev(Phylum), y=Abundance, color=fct_rev(Group))) +
  geom_boxplot()+
  scale_color_manual(values=colorgroups)+
  coord_flip()+
  scale_y_continuous(breaks = seq(0, 15, by=3), limits=c(0, 15))+
  theme bw()+
  theme(panel.grid = element blank())
# Statistical Comparison using wilcox test using specific bacteria. I need to manually enter the
bacteria name. It gets too confusing when i add multiple lines with all phyla
Phylum.lung.GC376.box.Special <- Phylum.lung.GC376.box %>%
  filter(Phylum == "Actinobacteriota")
compare_means(Abundance ~ Group, data = Phylum.lung.GC376.box.Special, method = "wilcox.test")
```

```
Phylum.lung.GC376.box.Special <- Phylum.lung.GC376.box %>%
    filter(Phylum == "Deinococcota")
compare_means(Abundance ~ Group, data = Phylum.lung.GC376.box.Special, method = "wilcox.test")
Phylum.lung.GC376.box.Special <- Phylum.lung.GC376.box %>%
    filter(Phylum == "Verrucomicrobiota")
compare_means(Abundance ~ Group, data = Phylum.lung.GC376.box.Special, method = "wilcox.test")
```

#### Figure 7B: Firmicutes/Bacteroidetes ration in the lung

Lets analyze the Firmicutes/Bacteroidetes ratio as this can be use as an indicator of dysbiosis.

I tested calculating the ratio with raw counts and with relative abundances and the results were very similar. This script is for calculating the ration using relative abundances

```
\# Create a data frame that only has the relative abundances of those classified as Firmicutes
Phylum.lung_RA_Firm <- Phylum.lung.GC376 %>%
filter(Phylum == "Firmicutes")
# Remove access columns that are not needed
# I will keep Abundance, SampleID, MouseID, Group and dpi
Phylum.lung_RA_Firm.edit <- Phylum.lung_RA_Firm[,c(3,4,8,9,10)]
# Change the first column header aka Abundance to Firmicutes
colnames(Phylum.lung_RA_Firm.edit)[1] <- "Firmicutes"</pre>
# Export the data frame as a csv file
write.csv(Phylum.lung_RA_Firm.edit, "Phylum.lung_RA_Firm.csv")
\# Create a data frame that only has the relative abundances of those classified as Bacteroidota
Phylum.lung_RA_Bact <- Phylum.lung.GC376 %>%
filter(Phylum == "Bacteroidota")
# Remove access columns that are not needed
# I will keep Abundance, SampleID, MouseID, Group and dpi
Phylum.lung_RA_Bact.edit <- Phylum.lung_RA_Bact[,c(3,4,8,9,10)]</pre>
# Change the first column header aka Abundance to Bacteroidota
colnames(Phylum.lung_RA_Bact.edit)[1] <- "Bacteroidota"</pre>
# Export the data frame as a csv file
write.csv(Phylum.lung_RA_Bact.edit, "Phylum.lung_RA_Bact.csv")
# I calculated the ratios of the Firmicutes/Bacteroidota in excel and then imported it back into R
Phylum.lung_RA_FBTOTAL <- read.csv("Phylum.lung.RA_FBRatio.csv")</pre>
# Make Group a factor
Phylum.lung_RA_FBTOTAL$Group <- factor(Phylum.lung_RA_FBTOTAL$Group, levels = c("Mock-GC376",
"Infected-3-GC376", "Infected-5-GC376"))
# Set the colors for the different groups
colorgroups = c("chocolate4", "forestgreen", "maroon1" )
# Graph the plot using ggplot
ggplot(data=Phylum.lung_RA_FBTOTAL, aes(x=FB_Ratio, y=Group, color=Group)) +
  geom boxplot()+
   theme bw()+
  scale color manual(values=colorgroups)+
  scale_x_continuous(breaks = seq(0, 6, by=2), limits=c(0, 6))+
  coord flip()+
  theme(panel.grid = element_blank())
# Statistical Comparison using wilcox test using specific bacteria. I need to manually enter the
bacteria name. It gets too confusing when i add multiple lines with all phyla
compare_means(FB_Ratio ~ Group, data = Phylum.lung_RA_FBTOTAL, method = "wilcox.test")
```

There is one outlier that is not shown in the Infected-5-GC376 group that is not shown (FB Ratio = 18)

#### Fig 7C: Taxonomic boxplots of the lung at the Family level

Filter the Family into the lung with the 3 groups that we will analyze. We will group the taxa that have less than 2% relative abundance since we are interested in overall trends and there were more families with lower relative abundances in the lungs compared to the ceca

```
# Filter the samples so that you only have lung samples and only GC376
Family.lung.GC376 <- Family %>%
  filter(SampleType == "Lung") %>%
  filter(Group == "Mock-GC376" | Group == "Infected-3-GC376" | Group == "Infected-5-GC376")
write.csv(Family.lung.GC376, "Family lung GC376.csv")
Lets look at all of the families in the lungs
# Filter the most abundant bacteria
Family.lung.GC376.box <- Family.lung.GC376
# Multipy abundances by 100
Family.lung.GC376.box$Abundance <- as.numeric(Family.lung.GC376.box$Abundance)
Family.lung.GC376.box$Abundance <- Family.lung.GC376.box$Abundance*100
# Assign the colors for the graphs
colorgroups = c("maroon1", "forestgreen", "chocolate4")
# Relevel the groups
Family.lung.GC376.box$Group <- factor(Family.lung.GC376.box$Group, levels = c("Mock-GC376", "Infected-
3-GC376",
          "Infected-5-GC376"))
# Graph plot using ggplot
ggplot(data=Family.lung.GC376.box, aes(x=fct rev(Family), y=Abundance, color=fct rev(Group))) +
  geom boxplot()+
  scale color manual(values=colorgroups)+
  coord flip()+
  scale_y_continuous(breaks = seq(0, 100, by=25), limits=c(0, 100))+
  theme_bw()+
 theme(panel.grid = element blank())
Since the abundances of all of the very rare families are making the trends of the bacterial families
```

Since the abundances of all of the very rare families are making the trends of the bacterial families that are prominant hard to see, we are going to group the families that have less than 1% relative abundance in a single sample.

```
\# I only want the families that have at least 1\$ relative abundance in a sample
Family.lung_greater_1_abundance <- Family.lung.GC376.box %>%
 filter(Family.lung.GC376.box[,3] >= 1.0) #this is 1 since we already multiplied the abundances by 100
to convert into %
unique(Family.lung greater 1 abundance$Family)
# Change phylum to a factor so I can change the order
Family.lung_greater_1_abundance$Family <- as.factor(Family.lung_greater_1_abundance$Family)
Family.lung_greater_1_abundance$Group<- factor(Family.lung_greater_1_abundance$Group, levels = c("Mock-
GC376", "Infected-3-GC376", "Infected-5-GC376"))
# Set the colors for the different groups
colorgroups = c("maroon1", "forestgreen", "chocolate4")
# Plot using ggplot using boxplots
ggplot(data=Family.lung greater 1 abundance, aes(x=fct rev(Family), y=Abundance, color=fct rev(Group)))
  geom boxplot()+
  scale_color_manual(values=colorgroups)+
  coord_flip()+
  scale_y_continuous(breaks = seq(0, 100, by=25), limits=c(0, 100))+
  theme_bw()+
 theme(panel.grid = element blank())
```

Then, I filtered out families that only appeared in a single sample (most likely contamination) Blattabacteriaceae, Caulobacteraceae, Halomonadaceae, Hydrogenophilaceae, Microbacteriaceae, Pasteurellaceae

Next, I will graph the most abundant families separately so that we can have different y axis for the less abundant and see the relationships among rare taxa more clear.

```
# Filter the most abundant bacteria
Family.lung.GC376.box <- Family.lung.GC376 %>%
    filter(Family == "Muribaculaceae" | Family == "Lachnospiraceae")
# Order the families
Family.lung.GC376.box$Family <- factor(Family.lung.GC376.box$Family, levels = c("Muribaculaceae",</pre>
```

```
"Lachnospiraceae"))
# Multipy abundances by 100
Family.lung.GC376.box$Abundance <- as.numeric(Family.lung.GC376.box$Abundance)
Family.lung.GC376.box$Abundance <- Family.lung.GC376.box$Abundance*100
# Assign the colors for the graphs
colorgroups = c("maroon1", "forestgreen", "chocolate4")
# Relevel the groups
Family.lung.GC376.box$Group <- factor(Family.lung.GC376.box$Group, levels = c("Mock-GC376", "Infected-3-GC376", "Infected-5-GC376"))
# Graph plot using ggplot
ggplot(data=Family.lung.GC376.box, aes(x=fct_rev(Family), y=Abundance, color=fct_rev(Group))) +
      geom boxplot()+
      scale color manual(values=colorgroups)+
     coord flip()+
     scale_y_continuous(breaks = seq(0, 100, by=25), limits=c(0, 100))+
     theme bw()+
     theme(panel.grid = element_blank())
 # Perform statistical
Family.lung.GC376.box.Special <- Family.lung.GC376.box %>%
     filter(Family == "Muribaculaceae")
compare means(Abundance ~ Group, data = Family.lung.GC376.box.Special, method = "wilcox.test")
Family.lung.GC376.box.Special <- Family.lung.GC376.box %>%
     filter(Family == "Lachnospiraceae")
compare means(Abundance ~ Group, data = Family.lung.GC376.box.Special, method = "wilcox.test")
Now, I will graph the less abundant families
# Filter the families that have lower abundance
Family.lung.GC376.less.box <- Family.lung.GC376 %>%
filter(Family == "Acholeplasmataceae" | Family == "Aerococcaceae" | Family == "Akkermansiaceae" | Family == "Bacteroidaceae" | Family == "Beijerinckiaceae" | Family == "Burkholderiaceae" | Family ==
"Butyricicoccaceae" | Family == "Caulobacteraceae" | Family == "Clostridiaceae" | Family == "Corynebacteriaceae" | Family == "Enterobacteriaceae" | Family == "Entero
                                                                                                                                                                                                                                               | Family ==
"Lactobacillaceae" | Family == "Leuconostocaceae" | Family == "Microbacteriaceae" | Family == "Moraxellaceae" | Family == "Moraxellaceae" | Family == "Secillospiraceae" | Family == "Peptococcaceae" | Family == "Peptococcaceae" | Family == "Peptococcaceae" | Family == "Ruminococcaceae" | Family == "Staphylococcaceae" | Family == "Sutterellaceae" | Family == "Staphylococcaceae" | Family == "Sutterellaceae" | Family == "Weeksellaceae" | Family == "W
Family == "Xanthomonadaceae")
# Relevel the families so that they are grouped by phylum
Family.lung.GC376.less.box$Family <- factor(Family.lung.GC376.less.box$Family, levels = c("Corynebacteriaceae", "Eggerthellaceae", "Microbacteriaceae",
  'Bacteroidaceae", "Weeksellaceae",
 "Thermaceae".
 "Acholeplasmataceae", "Aerococcaceae", "Butyricicoccaceae", "Clostridiaceae", "Enterococcaceae",
 "Erysipelatoclostridiaceae", "Erysipelotrichaceae", "Lactobacillaceae",
  Leuconostocaceae", "Oscillospiraceae", "Peptococcaceae", "Peptostreptococcaceae", "Ruminococcaceae",
 "Staphylococcaceae",
"Akkermansiaceae",
"Beijerinckiaceae", "Burkholderiaceae", "Caulobacteraceae", "Enterobacteriaceae", "Moraxellaceae", "Morganellaceae", "Pseudomonadaceae", "Rhodanobacteraceae", "Sphingomonadaceae", "Sutterellaceae", "Xanthomonadaceae"))
# Multipy abundances by 100
Family.lung.GC376.less.box$Abundance <- as.numeric(Family.lung.GC376.less.box$Abundance)
Family.lung.GC376.less.box$Abundance <- Family.lung.GC376.less.box$Abundance*100
# Assign the colors for the graphs
colorgroups = c("maroon1", "forestgreen", "chocolate4")
# Relevel the groups
Family.lung.GC376.less.box$Group <- factor(Family.lung.GC376.less.box$Group, levels = c("Mock-GC376",
 "Infected-3-GC376", "Infected-5-GC376"))
# Graph the plot using ggplot
ggplot(data=Family.lung.GC376.less.box, aes(x=fct_rev(Family), y=Abundance, color=fct_rev(Group))) +
     geom boxplot()+
```

```
scale_color_manual(values=colorgroups)+
coord_flip()+
scale_y_continuous(breaks = seq(0, 22, by=5), limits=c(0, 22))+
theme_bw()+
theme(panel.grid = element_blank())
```

Since there are still many families to place in the graph and it is hard to see. I will split the graphs into 2 separate graphs.

```
# Filter the families that have lower abundance
Family.lung.GC376.less.box <- Family.lung.GC376 %>%
filter(Family == "Acholeplasmataceae" | Family == "Aerococcaceae" | Family == "Bacteroidaceae" |
Family == "Butyricicoccaceae" | Family == "Clostridiaceae" | Family == "Enterococcaceae" | Family ==
"Erysipelatoclostridiaceae" | Family == "Erysipelotrichaceae" | Family == "Lactobacillaceae" | Family
== "Leuconostocaceae" | Family == "Oscillospiraceae" | Family == "Peptococcaceae" | Family == "Peptostreptococcaceae" | Family == "Ruminococcaceae" | Family == "Staphylococcaceae" | Family == "Peptostreptococcaceae" | Family == "Staphylococcaceae" | Family == "Staphylococcaceae" | Family == "Peptostreptococcaceae" | Family == "Staphylococcaceae" | Family == "Staphylococcaceae" | Family == "Peptostreptococcaceae" | Family == "Staphylococcaceae" | Family == "Staphylococcaceae
"Weeksellaceae")
# Relevel the families so that they are grouped by phylum
Family.lung.GC376.less.box$Family <- factor(Family.lung.GC376.less.box$Family, levels =
c("Bacteroidaceae", "Weeksellaceae",
"Acholeplasmataceae", "Aerococcaceae", "Butyricicoccaceae", "Clostridiaceae", "Enterococcaceae",
"Erysipelatoclostridiaceae", "Erysipelotrichaceae", "Lactobacillaceae", "Leuconostocaceae", "Oscillospiraceae", "Peptococcaceae", "Peptostreptococcaceae", "Ruminococcaceae", "Staphylococcaceae"))
# Multipy abundances by 100
Family.lung.GC376.less.box$Abundance <- as.numeric(Family.lung.GC376.less.box$Abundance)
Family.lung.GC376.less.box$Abundance <- Family.lung.GC376.less.box$Abundance*100
# Assign the colors for the graphs
colorgroups = c("maroon1", "forestgreen", "chocolate4")
# Relevel the groups
Family.lung.GC376.less.box$Group <- factor(Family.lung.GC376.less.box$Group, levels = c("Mock-GC376",
 "Infected-3-GC376", "Infected-5-GC376"))
# Graph the plot using ggplot
ggplot(data=Family.lung.GC376.less.box, aes(x=fct_rev(Family), y=Abundance, color=fct_rev(Group))) +
      geom_boxplot()+
      scale_color_manual(values=colorgroups)+
     coord_flip()+
     scale_y_continuous(breaks = seq(0, 22, by=5), limits=c(0, 22))+
     theme bw()+
     theme(panel.grid = element blank())
\# Statistical Comparison using wilcox test using specific bacteria. I need to manually enter the
bacteria name. It gets too confusing when i add multiple lines with all phyla
Family.lung.GC376.less.box.Special <- Family.lung.GC376.less.box %>%
     filter(Family == "Staphylococcaceae")
compare means (Abundance ~ Group, data = Family.lung.GC376.less.box.Special, method = "wilcox.test")
# Filter the families that have lower abundance
Family.lung.GC376.less.box <- Family.lung.GC376 %>%
filter(Family == "Akkermansiaceae" | Family == "Beijerinckiaceae" | Family == "Burkholderiaceae" |
Family == "Caulobacteraceae" | Family == "Corynebacteriaceae" | Family == "Enterobacteriaceae" | Family
== "Eggerthellaceae" | Family == "Microbacteriaceae" | Family == "Moraxellaceae" | Family == "Morganellaceae" | Family == "Morganellaceae" | Family == "Sphingomonadaceae" | Family == "Sphingomonadaceae" | Family == "Sphingomonadaceae" | Family == "Sphingomonadaceae" | Family == "Sutterellaceae" | Family == "Thermaceae" | Family == "Sphingomonadaceae" | Family == "Sutterellaceae" | Family == "Thermaceae" | Family == "Sphingomonadaceae" | Family == "Thermaceae" | Family == "Sphingomonadaceae" | Family == "Thermaceae" | Family == "Sphingomonadaceae" | Family == "Sphingomonadaceae" | Family == "Sphingomonadaceae" | Family == "Thermaceae" | Family == "Thermaceae" | Family == "Thermaceae" | Family == "Sphingomonadaceae" | Family == "Thermaceae" | Family == "Thermaceaee" | Family == "Thermaceaee" | Family == "Thermaceaeeeeeeeeeeeee
 "Xanthomonadaceae")
# Relevel the families so that they are grouped by phylum
Family.lung.GC376.less.box$Family <- factor(Family.lung.GC376.less.box$Family, levels =
c("Corynebacteriaceae", "Eggerthellaceae", "Microbacteriaceae",
  'Thermaceae",
"Akkermansiaceae",
"Beijerinckiaceae", "Burkholderiaceae", "Caulobacteraceae", "Enterobacteriaceae", "Moraxellaceae",
"Morganellaceae", "Pseudomonadaceae", "Rhodanobacteraceae", "Sphingomonadaceae", "Sutterellaceae",
"Xanthomonadaceae"))
# Multipy abundances by 100
Family.lung.GC376.less.box$Abundance <- as.numeric(Family.lung.GC376.less.box$Abundance)
Family.lung.GC376.less.box$Abundance <- Family.lung.GC376.less.box$Abundance*100
# Assign the colors for the graphs
```

```
colorgroups = c("maroon1", "forestgreen", "chocolate4")
# Relevel the groups
Family.lung.GC376.less.box$Group <- factor(Family.lung.GC376.less.box$Group, levels = c("Mock-GC376",
"Infected-3-GC376", "Infected-5-GC376"))
# Graph the plot using ggplot
ggplot(data=Family.lung.GC376.less.box, aes(x=fct_rev(Family), y=Abundance, color=fct_rev(Group))) +
  geom boxplot()+
  scale color manual(values=colorgroups)+
  coord flip()+
  scale_y_continuous(breaks = seq(0, 22, by=5), limits=c(0, 22))+
  theme_bw()+
  theme(panel.grid = element blank())
# Statistical Comparison using wilcox test using specific bacteria. I need to manually enter the
bacteria name. It gets too confusing when i add multiple lines with all phyla
Family.lung.GC376.less.box.Special <- Family.lung.GC376.less.box %>%
 filter(Family == "Xanthomonadaceae")
compare_means(Abundance ~ Group, data = Family.lung.GC376.less.box.Special, method = "wilcox.test")
```

One outlier in low group for Lactobacillaceae (~35%) and High dose Staphyloccoccacae (40%)

## Figure 7D: Correlation analysis of the lungs at the Family level

We will now perform a correlation analysis when analyzing the factors of infection to bacterial families

First, lets create a function to flatten a matrix as this will be needed for the correlation matrix

```
flattenCorrMatrix <- function(cormat, pmat) {
  ut <- upper.tri(cormat)
  data.frame(
    row = rownames(cormat)[row(cormat)[ut]],
    column = rownames(cormat)[col(cormat)[ut]],
    cor =(cormat)[ut],
    p = pmat[ut]
  )
}</pre>
```

Next, we will perform the correlation analysis using rcorr from the package Hmisc

```
# Filter the samples so that you only have cecum samples and only PBS, 3 and 5 infected vehicele
lung_added_factors <- read.csv("factors_lung.csv"). # file on github</pre>
lung_total_factors <- merge(Family.lung.GC376, lung_added_factors, by = "SampleID")</pre>
lung total factors edited \leftarrow lung total factors[,-c(1,2,3,5,6,7,8,9,11,12,13,14,15,16,17,18,19,21,23)]
#Corvnebacteriaceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Corynebacteriaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter_Lung <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter_Lung$row <- revalue(resultsfilter_Lung$row, c("Abundance"="Corynebacteriaceae"))
#Eggerthellaceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
filter(lung_total_factors_edited[,3] == "Eggerthellaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")</pre>
results <- flattenCorrMatrix(res$r, res$P)</pre>
resultsfilter_Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Eggerthellaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_Lung)</pre>
#Microbacteriaceae
Lung total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Microbacteriaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")</pre>
results <- flattenCorrMatrix(res$r, res$P)</pre>
```

```
resultsfilter_Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter Lung2$row <- revalue(resultsfilter Lung2$row, c("Abundance"="Microbacteriaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
#Bacteroidaceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Bacteroidaceae")
Lung_total_factors_edited_filter(,-3])
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter_Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Bacteroidaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Muribaculaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")</pre>
results <- flattenCorrMatrix(res$r, res$P)</pre>
resultsfilter_Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Muribaculaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
#Weeksellaceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Weeksellaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter Lung2 <- results %>%
filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Weeksellaceae"))</pre>
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
#Thermaceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Thermaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)</pre>
resultsfilter_Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Thermaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
#Acholeplasmataceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Acholeplasmataceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])
res <- rcorr(Lung total factors edited filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Acholeplasmataceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
#Aerococcaceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Aerococcaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")</pre>
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter_Lung2 <- results %>%
 filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Aerococcaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
#Butyricicoccaceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Butyricicoccaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)</pre>
resultsfilter_Lung2 <- results %>%
```

```
filter(results[,1] == "Abundance")
resultsfilter Lung2$row <- revalue(resultsfilter Lung2$row, c("Abundance"="Butyricicoccaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
#Clostridiaceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
filter(lung_total_factors_edited[,3] == "Clostridiaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung total factors edited filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter Lung2 <- results %>%
 filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Clostridiaceae"))
resultsfilter final <- rbind(resultsfilter Lung2, resultsfilter final)
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Enterococcaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)</pre>
resultsfilter_Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter Lung2$row <- revalue(resultsfilter Lung2$row, c("Abundance"="Enterococcaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
#Erysipelatoclostridiaceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Erysipelatoclostridiaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter_Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Erysipelatoclostridiaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
#Erysipelotrichaceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
filter(lung_total_factors_edited[,3] == "Erysipelotrichaceae")
Lung_total_factors_edited_filter (.-3])
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)</pre>
resultsfilter_Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter Lung2$row <- revalue(resultsfilter Lung2$row, c("Abundance"="Erysipelotrichaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
#Lachnospiraceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
filter(lung_total_factors_edited[,3] == "Lachnospiraceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter_Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Lachnospiraceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Lactobacillaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")</pre>
results <- flattenCorrMatrix(res$r, res$P)</pre>
resultsfilter_Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter Lung2$row <- revalue(resultsfilter Lung2$row, c("Abundance"="Lactobacillaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
#Leuconostocaceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Leuconostocaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")</pre>
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter_Lung2 <- results %>%
  filter(results[,1] == "Abundance")
```

```
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Leuconostocaceae"))
resultsfilter final <- rbind(resultsfilter Lung2, resultsfilter final)
#Oscillospiraceae
Lung total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Oscillospiraceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Oscillospiraceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
Lung total factors edited filter <- lung total factors edited %>%
 filter(lung_total_factors_edited[,3] == "Peptococcaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)</pre>
resultsfilter_Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Peptococcaceae"))
resultsfilter final <- rbind(resultsfilter Lung2, resultsfilter final)
#Peptostreptococcaceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
filter(lung_total_factors_edited[,3] == "Peptostreptococcaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Peptostreptococcaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
Lung total factors edited filter <- lung total factors edited %>%
  filter(lung_total_factors_edited[,3] == "Ruminococcaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)</pre>
resultsfilter_Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Ruminococcaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
#Staphylococcaceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Staphylococcaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)</pre>
resultsfilter Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Staphylococcaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Beijerinckiaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")</pre>
results <- flattenCorrMatrix(res$r, res$P)</pre>
resultsfilter_Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Beijerinckiaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
#Burkholderiaceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Burkholderiaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter_Lung2 <- results %>%
 filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Burkholderiaceae"))
```

```
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)
#Caulobacteraceae
Lung total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Caulobacteraceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")</pre>
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Caulobacteraceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Enterobacteriaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung total factors edited filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter_Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Enterobacteriaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
#Moraxellaceae
Lung total factors edited filter <- lung total factors edited %>%
  filter(lung_total_factors_edited[,3] == "Enterococcaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter_Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Moraxellaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
#Morganellaceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Morganellaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter_Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Morganellaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
#Pseudomonadaceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
 filter(lung_total_factors_edited[,3] == "Pseudomonadaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")</pre>
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter_Lung2 <- results %>%
 filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Pseudomonadaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
#Rhodanobacteraceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Rhodanobacteraceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)</pre>
resultsfilter_Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Rhodanobacteraceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
#Sphingomonadaceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Sphingomonadaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")</pre>
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter_Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Sphingomonadaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
```

```
#Sutterellaceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
   filter(lung_total_factors_edited[,3] == "Sutterellaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)</pre>
resultsfilter_Lung2 <- results %>%
   filter(results[,1] == "Abundance")
resultsfilter Lung2$row <- revalue(resultsfilter Lung2$row, c("Abundance"="Sutterellaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
#Xanthobacteraceae
Lung_total_factors_edited_filter <- lung_total_factors_edited %>%
   filter(lung_total_factors_edited[,3] == "Xanthobacteraceae")
Lung total factors edited filter <- as.matrix(Lung total factors edited filter[,-3])
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")</pre>
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter_Lung2 <- results %>%
   filter(results[,1] == "Abundance")
resultsfilter_Lung2$row <- revalue(resultsfilter_Lung2$row, c("Abundance"="Xanthobacteraceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
#Akkermansiaceae
Lung total_factors_edited_filter <- lung_total_factors_edited %>%
  filter(lung_total_factors_edited[,3] == "Akkermansiaceae")
Lung_total_factors_edited_filter <- as.matrix(Lung_total_factors_edited_filter[,-3])</pre>
res <- rcorr(Lung_total_factors_edited_filter, type = "spearman")
results <- flattenCorrMatrix(res$r, res$P)
resultsfilter_Lung2 <- results %>%
  filter(results[,1] == "Abundance")
resultsfilter Lung2$row <- revalue(resultsfilter Lung2$row, c("Abundance"="Akkermansiaceae"))
resultsfilter_final <- rbind(resultsfilter_Lung2, resultsfilter_final)</pre>
# Heatmap
resultsfilter_final$row <- factor(resultsfilter_final$row, levels = c("Corynebacteriaceae",
"Eggerthellaceae", "Microbacteriaceae", "Bacteroidaceae", "Muribaculaceae", "Weeksellaceae",
"Thermaceae", "Acholeplasmataceae", "Aerococcaceae", "Butyricicoccaceae", "Clostridiaceae",
"Enterococcaceae", "Erysipelatoclostridiaceae", "Erysipelotrichaceae", "Lachnospiraceae",
"Lactobacillaceae", "Leuconostocaceae", "Oscillospiraceae", "Peptococcaceae", "Peptostreptococcaceae",
"Ruminococcaceae", "Staphylococcaceae", "Beijerinckiaceae", "Burkholderiaceae", "Caulobacteraceae",
"Enterobacteriaceae", "Moraxellaceae", "Morganellaceae", "Pseudomonadaceae", "Rhodanobacteraceae",
"Sphingomonadaceae", "Sutterellaceae", "Xanthobacteraceae", "Akkermansiaceae"))</pre>
resultsfilter_final$column <- factor(resultsfilter_final$column, levels = c("Inoculum", "dpi.x",
                  "Viral.NT.Titer", "Viral.Lung.Titer",
                                                                   "Viral.Brain.Titer"))
ggplot(resultsfilter_final, aes(row, fct_rev(column))) +
   #geom_tile(aes(fill = p))+
   #scale_fill_gradient(low = "snow",
                              high = "snow")+
   geom_point(aes(colour = cor),
                         size = 8) +
   scale\_color\_gradient2(limits = c(-0.7, 0.7), breaks = c(-0.6, -0.3, 0, 0.3, 0.6),
                              low = "royalblue1",
                              mid = "brown3".
                              high = "yellow")+
   theme_bw()
resultsfilter_final_significant <- resultsfilter_final %>%
 filter(resultsfilter_final[,4] < 0.05)</pre>
```

The figure was further edited in Adobe Illustrator to indicate the relationships that were considered statistically significant.

## Figure 8: Taxonomic barplot of the lungs at the Family level

Now lets produce a graph for the taxonomic barplot

```
# Combine the Taxa that are less than 1% relative abundance together
Family.lung.barplot <- Family.lung.GC376
Family.lung.barplot$Family[Family.lung.barplot$Abundance < 0.02] <- "Taxa < 2% abund."

# Set the MouseID to a factor and assign the order
Family.lung.barplot$MouseID <- factor(Family.lung.barplot$MouseID, levels = c("13", "14", "45", "46", "47", "56", "57", "58", "11", "12", "42", "43", "44", "53", "54", "55", "7", "8", "9", "37", "38", "39", "51"))</pre>
```

```
# Set the level of the families
Family.lung.barplot$Family <- factor(Family.lung.barplot$Family, levels = c("Eggerthellaceae",
Family.lung.barplot$Family <- factor(Family.lung.barplot$Family, levels = c("Eggerthellaceae", "Corynebacteriaceae", "Muribaculaceae", "Bacteroidaceae", "Weeksellaceae", "Lachnospiraceae", "Staphylococcaceae", "Erysipelatoclostridiaceae", "Peptostreptococcaceae", "Clostridiaceae", "Ruminococcaceae", "Oscillospiraceae", "Erysipelotrichaceae", "Lactobacillaceae", "Leuconostocaceae", "Enterococcaceae", "Aerococcaceae", "Butyricicoccaceae", "Pseudomonadaceae", "Enterobacteriaceae", "Burkholderiaceae", "Xanthomonadaceae", "Caulobacteraceae", "Sphingomonadaceae", "Moraxellaceae", "Akkermansiaceae", "Taxa < 2% abund."))
# Set color palette to accommodate the number of families
ColorTest = c("gray31", "yellow",
"springgreen4", "olivedrab1", "olivedrab4", "yellowgreen", "darkolivegreen", "chartreuse", "forestgreen", "dodgerblue", "lightblue3", "navy", "cornflowerblue", "lightblue2", "cyan", "mediumblue", "steelblue3", "blue", #these are same colors tak
                                                                                                                                          #these are same colors taken
from ceca
                          "dodgerblue3","skyblue2","royalblue1",
"deepskyblue1", #these
                                                                                     #these are same colors taken from ceca
                           "purple1",
                          "darkorchid4", "orchid1", "darkorange3", "sandybrown")
                                                                                     #these are same colors taken from ceca
# Plot using ggplot
ggplot(data=Family.lung.barplot, aes(x=MouseID, y=Abundance, fill=fct_rev(Family))) +
geom_bar(aes(), colour = "black", stat="identity", position="stack", width = 0.8) +
    scale_fill_manual(values=ColorTest)+
    facet_grid(~dpi, scales = "free", space = "free")+
theme(legend.position="bottom") +
    guides(fill=guide_legend(nrow=5))+
    theme(legend.position="right") + guides(fill=guide_legend(ncol =1))+
    theme bw()+
    theme(panel.grid = element_blank())+
   theme(strip.text = element_text(size=11, face="bold"))
```

The figure is then combined with activity score and viral titers in Adobe Illustrator