Coding Challenge 4

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The data used in this project was provided by: Noel, Z.A., Roze, L.V., Breunig, M., Trail, F. 2022. Endophytic fungi as promising biocontrol agent to protect wheat from Fusarium graminearum head blight. Plant Disease. See the paper here

```
#install.packages('tinytex')
#tinytex::install_tinytex()
library(ggplot2)
library(tidyverse)
library(ggpubr)
library(ggrepel)
library(ggrepel)
library(knitr)

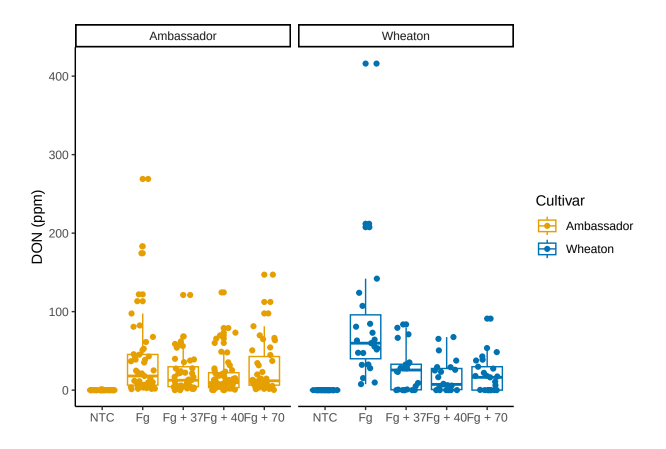
#getwd()
mycotoxin <- read.csv("MycotoxinData.csv", na.strings = "na")</pre>
```

#Question 2 - Change the factor order level so that the treatment "NTC" is first, followed by "Fg", "Fg + 37", "Fg + 40", and "Fg + 70".

```
 \label{thm:mycotoxin} $$\operatorname{Treatment, levels} = c("NTC", "Fg", "Fg + 37", "Fg + 40", "
```

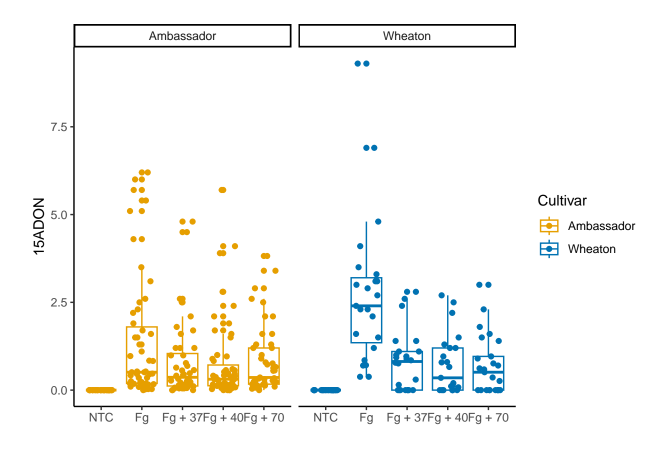
#Question 2 Boxplot

```
Question_2 <- ggplot(mycotoxin, aes(x = Treatment, y = DON, color = Cultivar)) +
    geom_boxplot() +
    geom_point(position = position_jitterdodge(jitter.width = 0.6)) +
    scale_color_manual(values = c("#E69F00", "#0072B2")) +
    xlab("") +
    ylab("DON (ppm)") +
    theme_classic() +
    facet_wrap(~Cultivar)
Question_2</pre>
```



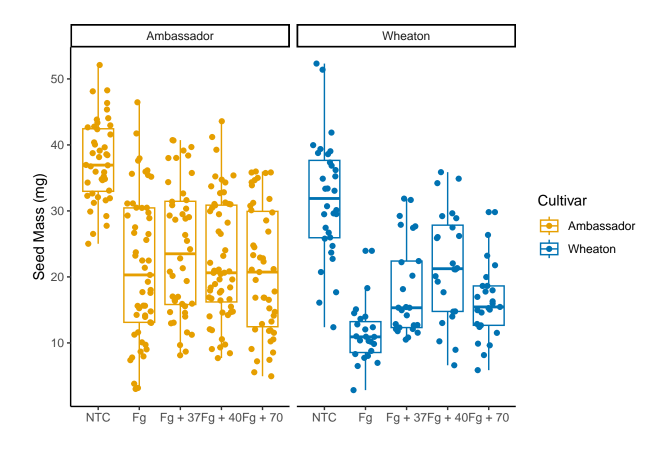
#Question 3a - Change the y-variable to plot X15ADON. The y-axis label should now be "15ADON".

```
Question_3a <- ggplot(mycotoxin, aes(x = Treatment, y = X15ADON, color = Cultivar)) +
    geom_boxplot() +
    geom_point(position = position_jitterdodge(jitter.width = 0.6)) +
    scale_color_manual(values = c("#E69F00", "#0072B2")) +
    xlab("") +
    ylab("15ADON") +
    theme_classic() +
    facet_wrap(~Cultivar)
Question_3a</pre>
```



#Question 3b - Change the y-variable to plot MassperSeed_mg. The y-axis label should now be "Seed Mass (mg)".

```
Question_3b <- ggplot(mycotoxin, aes(x = Treatment, y = MassperSeed_mg, color = Cultivar)) +
    geom_boxplot() +
    geom_point(position = position_jitterdodge(jitter.width = 0.6)) +
    scale_color_manual(values = c("#E69F00", "#0072B2")) +
    xlab("") +
    ylab("Seed Mass (mg)") +
    theme_classic() +
    facet_wrap(~Cultivar)
Question_3b</pre>
```

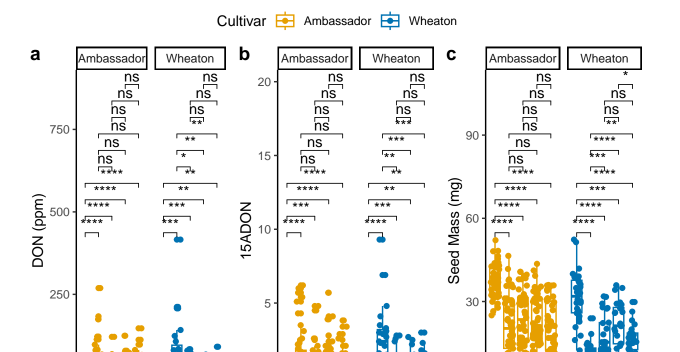


#Question 5 - Use geom_pwc() to add t.test pairwise comparisons to the three plots made above. Save each plot as a new R object, and combine them again with ggarange as you did in question 4.

```
Stats_Question_2 <- Question_2 +
    geom_pwc(aes(group = Treatment), method = "t_test", label = "p.adj.signif")
#Stats_Question_2

Stats_Question3a <- Question_3a +
    geom_pwc(aes(group = Treatment), method = "t_test", label = "p.adj.signif")
#Stats_Question3b <- Question_3b +
    geom_pwc(aes(group = Treatment), method = "t_test", label = "p.adj.signif")
#Stats_Question3b

#combination graph
Question_5Combo <- ggarrange(Stats_Question_2, Stats_Question3a, Stats_Question3b, labels = "auto",
    nrow = 1,
    ncol = 3,
    common.legend = T)
Question_5Combo</pre>
```



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