# $Coding\_Challenge4\_Markdown\_mer0127\_kat0084$

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#### **Data Information**

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. DOI

# Coding Challenge 3

#### Quesiton Five

- 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

```
library(ggplot2) #version 3.5.1
library(ggpubr) #version 0.6.0
library(ggrepel) #version 0.9.6
```

#### Data Set

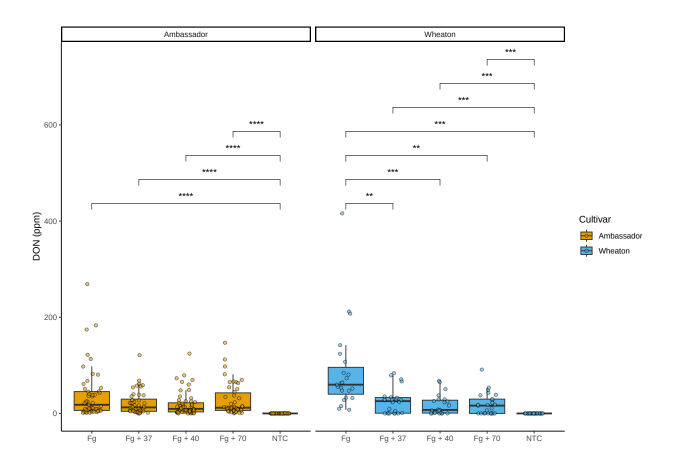
```
Mycotoxin<- read.csv("MycotoxinData.csv", na.strings = "na")
head(Mycotoxin)</pre>
```

```
Treatment Cultivar BioRep MassperSeed_mg
##
                                                 DON X15ADON
## 1
                              2
                                     10.291304 107.3
                                                         3.00
            Fg
                Wheaton
## 2
                Wheaton
                                     12.803226 32.6
                                                         0.85
## 3
                              2
                                      2.846667 416.0
                                                         3.50
                Wheaton
                Wheaton
                              2
                                      6.500000 211.9
                                                         3.10
            Fg
                              2
                                     10.179167 124.0
## 5
                                                         4.80
            Fg
                Wheaton
## 6
            Fg
                Wheaton
                                     12.044444 73.1
                                                         3.30
```

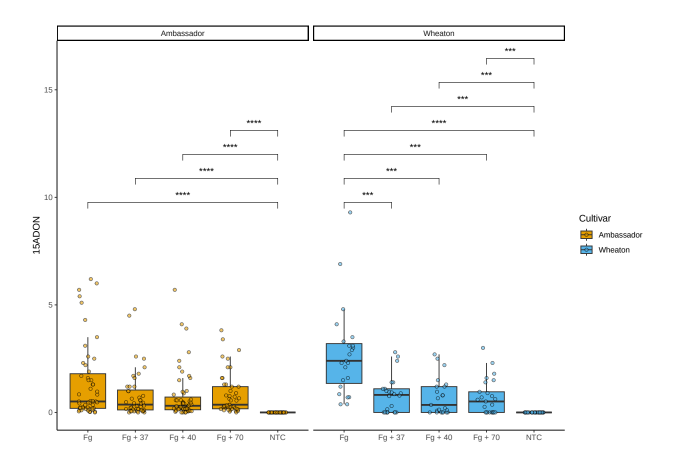
```
## 'data.frame': 375 obs. of 6 variables:
## $ Treatment : chr "Fg" "Fg" "Fg" "Fg" ...
## $ Cultivar : chr "Wheaton" "Wheaton" "Wheaton" "Wheaton" ...
## $ BioRep : int 2 2 2 2 2 2 2 2 2 3 ...
## $ MassperSeed_mg: num 10.29 12.8 2.85 6.5 10.18 ...
## $ DON : num 107.3 32.6 416 211.9 124 ...
## $ X15ADON : num 3 0.85 3.5 3.1 4.8 3.3 6.9 2.9 2.1 0.71 ...
New Color Palette

cbbPalette <- c( "#E69F00", "#56B4E9", "#009E73", "#F0E442", "#0072B2", "#D55E00", "#CC79A7", "#000000")</pre>
```

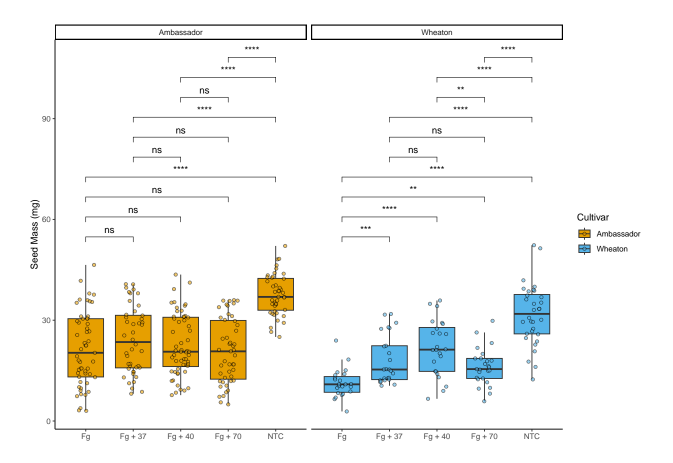
#### DON with pairwise comparisons



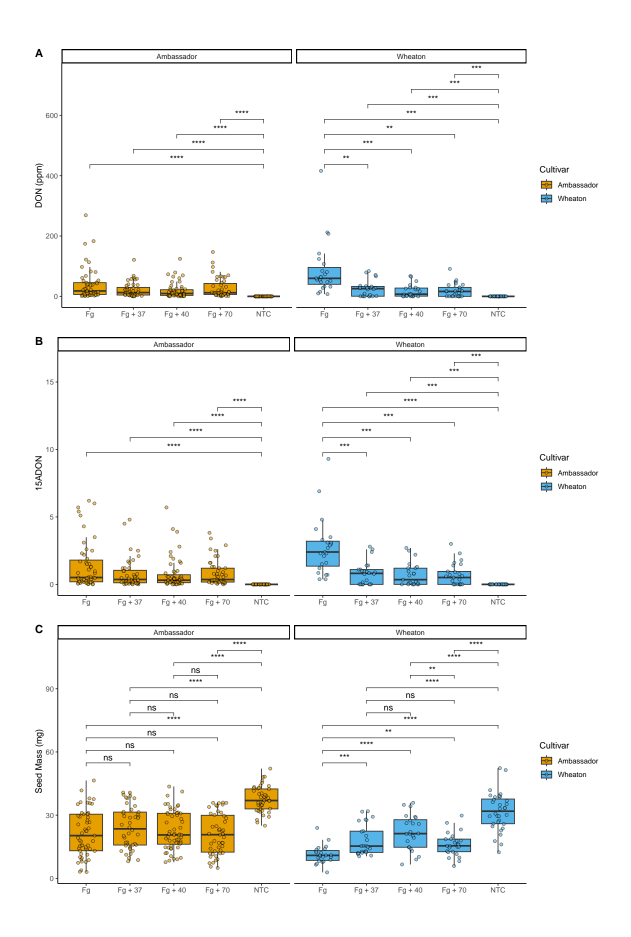
## X15ADON with pairwise comparisons



### ${\bf Massper Seed\_mg\ with\ pairwise\ comparisons}$



### Combining the three plots into one file



ggsave("combined\_plot.jpeg", plot = combined\_plots, width =15 , height = 10)
#Saving Plot to Rproject Folder