# Coding Challenge. 4

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## Coding Challenge 3

Q.No.3 (a)

Readme file link

link to my Readme file

#### Clickable Link to the Manuscript

Manuscript Link

Q.No.6 (b)

File Tree of Github

```
fs::dir_tree()
```

```
## .
## +-- Coding challenge.4.Rmd
## +-- Coding Challenge.4.Rproj
## +-- Coding-challenge.4.docx
## +-- Coding-challenge.4.html
## +-- Coding-challenge.4.md
## +-- Coding-challenge.4.pdf
## +-- Coding-challenge.4.Rmd
## +-- Coding-challenge.4_files
     \-- figure-gfm
           +-- unnamed-chunk-5-1.png
## |
## |
           +-- unnamed-chunk-6-1.png
## |
          +-- unnamed-chunk-7-1.png
## |
           +-- unnamed-chunk-8-1.png
           \-- unnamed-chunk-9-1.png
## \-- MycotoxinData.csv
```

#### Libraries

```
library(readr)
library(ggplot2)
library(tidyverse)
## -- Attaching core tidyverse packages ----- tidyverse 2.0.0 --
## v dplyr
             1.1.4
                       v stringr
                                  1.5.1
## v forcats 1.0.0
                       v tibble
                                   3.2.1
                                   1.3.1
## v lubridate 1.9.4
                       v tidyr
             1.0.2
## v purrr
## -- Conflicts -----
                                           ## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                 masks stats::lag()
## i Use the conflicted package (<a href="http://conflicted.r-lib.org/">http://conflicted.r-lib.org/</a>) to force all conflicts to become error
library(ggpubr)
library(ggrepel)
cbbPalette <- c("#000000", "#E69F00", "#56B4E9", "#009E73",
               "#F0E442", "#0072B2", "#D55E00", "#CC79A7")
```

#### Q.No.3 (b)

Importing the excel file

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

#### Q.No.3 (c)

Changing the factor levels

```
Mycotoxin_Data$Treatment <- factor(Mycotoxin_Data$Treatment, levels = c("NTC", "Fg", "Fg + 37", "Fg + 4
```

## Appling Statistics using t test

a.

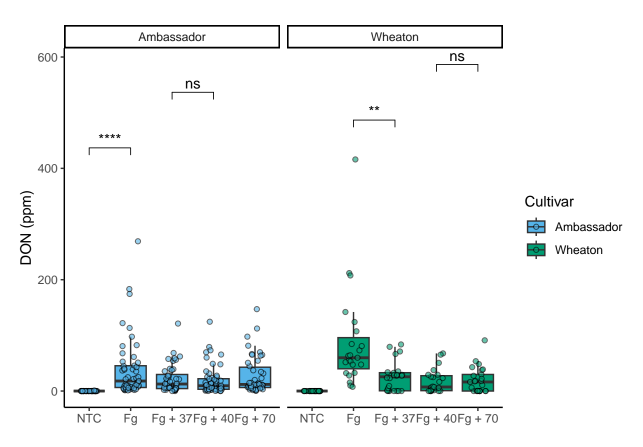
```
DONplot.pwc <- ggplot(Mycotoxin_Data, aes(x = Treatment, y = DON, fill = Cultivar)) +
  geom_boxplot(outliers = F) +
  xlab("") +
  ylab("DON (ppm)") +
  geom_point(alpha = 0.6, pch = 21, color = "black", position = position_jitterdodge()) +
  scale_color_manual(values = c(cbbPalette[3], cbbPalette[4])) +
  scale_fill_manual(values = c(cbbPalette[3], cbbPalette[4])) +</pre>
```

```
## Warning: Removed 8 rows containing non-finite outside the scale range
## ('stat_boxplot()').

## Warning: Removed 8 rows containing non-finite outside the scale range
## ('stat_signif()').

## Warning: Removed 8 rows containing missing values or values outside the scale range
```

## Warning: Removed 8 rows containing missing values or values outside the scale range
## ('geom\_point()').

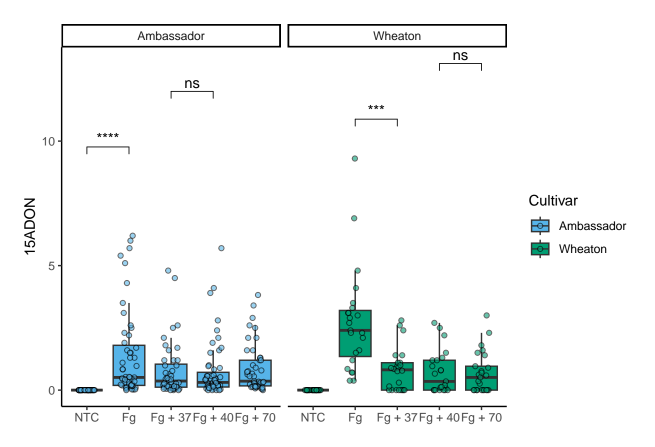


b.

## Warning: Removed 10 rows containing non-finite outside the scale range
## ('stat\_boxplot()').

## Warning: Removed 10 rows containing non-finite outside the scale range
## ('stat\_signif()').

## Warning: Removed 10 rows containing missing values or values outside the scale range
## ('geom\_point()').



#### Combining all plots with statistics

```
Combineplot.pwc <- ggarrange( DONplot.pwc,</pre>
                                   X15ADONplot.pwc,
                       Seed.massplot.pwc,
                       labels = c("A", "B", "C"),
                       nrow = 1,
                       ncol = 3,
                       common.legend = TRUE,
                       legend = "bottom")
## Warning: Removed 8 rows containing non-finite outside the scale range
## ('stat boxplot()').
## Warning: Removed 8 rows containing non-finite outside the scale range
## ('stat signif()').
## Warning: Removed 8 rows containing missing values or values outside the scale range
## ('geom_point()').
## Warning: Removed 8 rows containing non-finite outside the scale range
## ('stat_boxplot()').
## Warning: Removed 8 rows containing non-finite outside the scale range
## ('stat_signif()').
## Warning: Removed 8 rows containing missing values or values outside the scale range
## ('geom_point()').
## Warning: Removed 10 rows containing non-finite outside the scale range
## ('stat_boxplot()').
```

```
## Warning: Removed 10 rows containing non-finite outside the scale range ## ('stat_signif()').
```

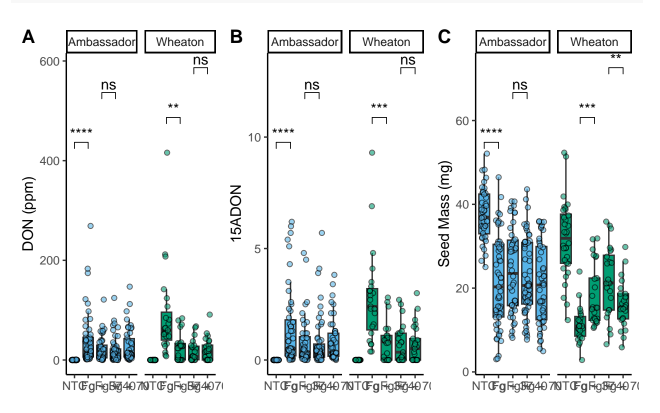
## Warning: Removed 10 rows containing missing values or values outside the scale range
## ('geom\_point()').

## Warning: Removed 2 rows containing non-finite outside the scale range
## ('stat\_boxplot()').

## Warning: Removed 2 rows containing non-finite outside the scale range
## ('stat\_signif()').

## Warning: Removed 2 rows containing missing values or values outside the scale range
## ('geom\_point()').

#### Combineplot.pwc



Cultivar Ambassador Wheaton