## Datawrangling

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## load required library

```
library(tidyverse)
## Warning: package 'tidyverse' was built under R version 4.4.2
## Warning: package 'ggplot2' was built under R version 4.4.2
## Warning: package 'readr' was built under R version 4.4.2
## Warning: package 'forcats' was built under R version 4.4.2
## Warning: package 'lubridate' was built under R version 4.4.2
## -- Attaching core tidyverse packages ----- tidyverse 2.0.0 --
## v dplyr
             1.1.4
                       v readr
                                    2.1.5
## v forcats 1.0.0
                                    1.5.1
                       v stringr
## v ggplot2 3.5.1
                       v tibble
                                    3.2.1
## v lubridate 1.9.3
                        v tidyr
                                    1.3.1
## v purrr
              1.0.2
## -- Conflicts ----- tidyverse_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(dplyr)
```

1. 3 pts. Download two .csv files from Canvas called DiversityData.csv and Metadata.csv, and read them into R using relative file paths.

```
diversitydata=read.csv("DiversityData.csv")
metadata=read.csv("Metadata.csv",na.strings="na")
```

2. 4 pts. Join the two dataframes together by the common column 'Code'. Name the resulting dataframe alpha.

```
alpha = metadata %>%
full_join(diversitydata, by = "Code")
```

- 3. 4 pts. Calculate Pielou's evenness index: Pielou's evenness is an ecological parameter calculated by the Shannon diversity index (column Shannon) divided by the log of the richness column.
- a. Using mutate, create a new column to calculate Pielou's evenness index.
- b. Name the resulting dataframe alpha even.

```
alpha_even <- alpha %>%
  mutate(Pielou_evenness = shannon / log(richness))
```

- 4. Pts. Using tidyverse language of functions and the pipe, use the summarise function and tell me the mean and standard error evenness grouped by crop over time.
- a. Start with the alpha even dataframe
- b. Group the data: group the data by Crop and Time Point.
- c. Summarize the data: Calculate the mean, count, standard deviation, and standard error for the even variable within each group.
- d. Name the resulting dataframe alpha average

```
alpha_average <- alpha_even %>%
group_by(Crop, Time_Point) %>%
summarise(
   mean_evenness = mean(Pielou_evenness, na.rm = TRUE),
   count = n(),
   sd_evenness = sd(Pielou_evenness, na.rm = TRUE),
   se_evenness = sd_evenness / sqrt(count)
)
```

```
## 'summarise()' has grouped output by 'Crop'. You can override using the
## '.groups' argument.
```

- 5. 4. Pts. Calculate the difference between the soybean column, the soil column, and the difference between the cotton column and the soil column
- a. Start with the alpha\_average dataframe
- b. Select relevant columns: select the columns Time Point, Crop, and mean.even.
- c. Reshape the data: Use the pivot\_wider function to transform the data from long to wide format, creating new columns for each Crop with values from mean.even.
- d. Calculate differences: Create new columns named diff.cotton.even and diff.soybean.even by calculating the difference between Soil and Cotton, and Soil and Soybean, respectively.
- e. Name the resulting data frame alpha\_average2  $\,$

```
alpha_average2 <- alpha_average %>%
  select(Time_Point, Crop, mean_evenness) %>%
  pivot_wider(names_from = Crop, values_from = mean_evenness) %>%
  mutate(
    diff_cotton_even = Soil-Cotton,
    diff_soybean_even = Soil-Soybean
)
```

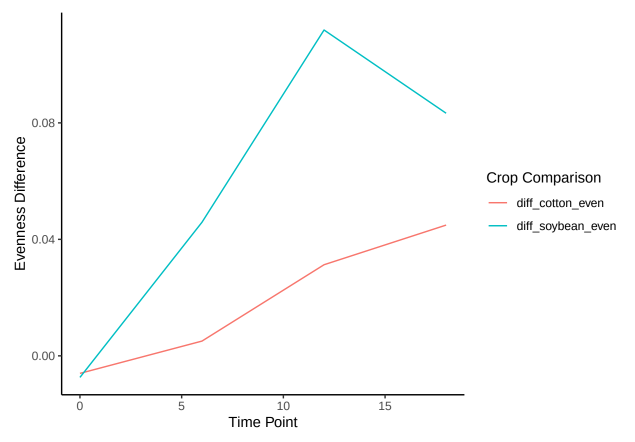
- 6. 4 pts. Connecting it to plots
- a. Start with the alpha\_average2 dataframe
- b. Select relevant columns: select the columns Time\_Point, diff.cotton.even, and diff.soybean.even.
- c. Reshape the data: Use the pivot\_longer function to transform the data from wide to long format, creating a new column named diff that contains the values from diff.cotton.even and diff.soybean.even.
- d. This might be challenging, so I'll give you a break. The code is below.

pivot\_longer(c(diff.cotton.even, diff.soybean.even), names\_to = "diff")

d. Create the plot: Use ggplot and geom\_line() with 'Time\_Point' on the x-axis, the column 'values' on the y-axis, and different colors for each 'diff' category. The column named 'values' come from the pivot\_longer. The resulting plot should look like the one to the right.

```
alpha_long <- alpha_average2 %>%
    select(Time_Point, diff_cotton_even, diff_soybean_even) %>%
    pivot_longer(cols = c(diff_cotton_even, diff_soybean_even), names_to = "diff", values_to = "values")

# Create the plot
ggplot(alpha_long, aes(x = Time_Point, y = values, color = diff)) +
    geom_line() +
    labs(x = "Time_Point", y = "Evenness_Difference", color = "Crop_Comparison") +
    theme_classic()
```



7. 2 pts. Commit and push a gfm .md file to GitHub inside a directory called Coding Challenge 5. Provide me a link to your github written as a clickable link in your .pdf or .docx

Challenge 5 GitHub Link