Data analysis of Arabidopsis experiment B1

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Loading the packages

```
library(readxl)
library(tidyverse)
library(gdata)
library(car)
library(cowplot)
library(magick)
library(here)
```

The first step is to load in the data from the Excel file and combine it with the metadata.

Loading file and generating complete dataframe

```
data_raw <- read_excel("data/arabidopsis_data.xlsx", col_types = c("text", rep("numeric", 8), "text"))</pre>
metadata <- tibble("Group" = c("L1", "L2", "S3", "S4", "L5", "L6", "S7", "S8"),
                    "Population" = rep(c("Bovra", "Smadalen", "Helin", "Spiterstulen"), each = 2),
                    "Size" = rep(c("Large", "Small", "Large", "Small"), each = 2),
                    "Elevation" = rep("High", each = 8),
                    "Condition" = rep(c("Stable", "Declining", "Declining", "Declining"), each = 2), #difference bet
                    "Soil" = rep(c("Riverbed", "Riverbed", "Scree", "Rocks"), each = 2),
                    "Treatment" = c("Unsalted", "Salted", "Unsalted", "Salted", "Unsalted", "Salted", "Sal
                    "Conc_NaCl" = c(0, 50, 0, 50, 0, 50, 0, 50),
                    "Conc_unit" = rep("mM", each = 8),
                    "Length_unit" = rep("cm", each = 8),
                    "Weight_unit" = rep("grams"), each = 8)
data <- mutate(data_raw,</pre>
                                           "Length_dif" = data_raw$Length_longest_leaf_a - data_raw$Length_longest_leaf_b,
                                           "Number_dif" = data_raw$Number_leaves_a - data_raw$Number_leaves_b) #calculating differe
data_complete <- left_join(data, metadata, by = "Group") #create one complete dataframe
```

Now to begin the data analysis, the Shapiro-Wilk values of each group are calculated too see whether the values are normally distributed.

```
names <- c("Wet_weight", "Length_dif", "Number_dif") #collect the names of the columns of interest
sw_results <- vector("list", 3) #prepare an empty list

for (i in 1:3) { #create a list of dataframes where every dataframe contains its Shapiro-Wilk p-values
result <- data_complete %>% group_by(Size, Treatment) %>% summarize(sw = shapiro.test(!!sym(names[i]))$
result <- unite(result, "Treatment", Size, Treatment)
sw_results[[i]] <- result</pre>
```

```
merged <- sw_results %>% reduce(full_join, by= "Treatment") #merge all the dataframes from the list
tidy_sw_results <- pivot_longer(data = merged, cols = names,</pre>
                              names_to = "Info", values_to = "ShapiroWilk_p.value") #make the dataframe
tidy_sw_results %>% filter(ShapiroWilk_p.value > 0.05) #filter for which groups are normally distribute
tidy_sw_results %>% filter(ShapiroWilk_p.value < 0.05) #filter for which groups are NOT normally distri
All groups of interest are normally distributed, except for the difference in number of leaves
To check whether there is a difference between the wet weights and growth (difference in length of the longest
leaf) between the salted and unsalted groups, a t-test is used after checking for equal variance:
leveneTest(data_complete$Wet_weight ~ as.factor(Treatment), data = data_complete) #Pr(>F) = 0.4168, whi
## Levene's Test for Homogeneity of Variance (center = median)
        Df F value Pr(>F)
              0.667 0.4168
## group 1
leveneTest(data_complete$Length_dif ~ as.factor(Treatment), data = data_complete) #Pr(>F) = 0.4793, whi
## Levene's Test for Homogeneity of Variance (center = median)
         Df F value Pr(>F)
## group 1 0.5058 0.4793
##
         72
t.test(formula = data_complete$Length_dif ~ data_complete$Treatment,
       paired = FALSE, var.equal = TRUE)$p.value %>% round(.,3) #t-test, p = 0.005, which means there i
## [1] 0.005
t.test(formula = data_complete$Wet_weight ~ data_complete$Treatment,
       paired = FALSE, var.equal = TRUE) p.value \%\% round(.,3) #t-test, p = 0.011, which means there i
## [1] 0.011
wilcox.test(Number_dif ~ Treatment, data = data_complete) #Wilcoxon test, p = 0.5845, which means there
## Warning in wilcox.test.default(x = DATA[[1L]], y = DATA[[2L]], ...): cannot
## compute exact p-value with ties
##
## Wilcoxon rank sum test with continuity correction
##
## data: Number_dif by Treatment
## W = 715, p-value = 0.5845
## alternative hypothesis: true location shift is not equal to 0
These differences are interesting to visualize as they are statistically significant:
Generating the first plots
summary <- data_complete %>%
  group_by(Treatment) %>%
  summarize(mean_noleaves.increase = mean(Number_dif, na.rm = TRUE),
            mean_lengthleaf.increase = mean(Length_dif, na.rm = TRUE),
            mean_wetweight = mean(Wet_weight, na.rm = TRUE),
```

```
stdevLength = sd(Length_dif, na.rm = TRUE),
            stdevWetweight = sd(Wet_weight, na.rm = TRUE),
            stdevNumber = sd(Number_dif, na.rm = TRUE))
plot_lengthdiffull <- summary %>% #plot the increase in longest leaf length grouped by population size
  ggplot(aes(x = Treatment, y = mean_lengthleaf.increase))+
  geom_col(aes(fill = Treatment))+
  geom_errorbar(aes(ymin = mean_lengthleaf.increase - stdevLength,
                    ymax = mean_lengthleaf.increase + stdevLength), width=.2)+
  theme light()+
  labs(title = "Increase in length of the longest\nleaf, grouped by treatment",
      subtitle = "Error bars depict 1 standard deviation",
       y="Mean increase in the length\nof the longest leaf in cm")+
  theme(legend.position = "none", text = element_text(size=12), plot.title = element_text(size=14),
       plot.subtitle = element_text(size=12))
plot_wetweightfull <- summary %>% #plot the wet weight grouped by population size
  ggplot(aes(x = Treatment, y = mean_wetweight)) +
  geom_col(aes(fill = Treatment))+
  geom_errorbar(aes(ymin = mean_wetweight - stdevWetweight,
                    ymax = mean_wetweight + stdevWetweight), width=.2)+
  theme_light()+
  labs(title = "Wet weight, grouped by treatment",
      subtitle = "Error bars depict 1 standard deviation",
      x="Treatment",
      y="Wet weight in grams")+
  theme(legend.position = "none", text = element_text(size=12), plot.title = element_text(size=14),
       plot.subtitle = element_text(size=12))
plot_numberdiffull <- summary %>% #plot the increase in number of leaves grouped by population size
  ggplot(aes(x = Treatment, y = mean_noleaves.increase)) +
  geom_col(aes(fill = Treatment))+
  geom_errorbar(aes(ymin = mean_noleaves.increase - stdevNumber,
                    ymax = mean_noleaves.increase + stdevNumber), width=.2)+
  theme_light()+
  labs(title = "Increase in number of leaves, \ngrouped by treatment",
       subtitle = "Error bars depict 1 standard deviation",
       x="Treatment",
       y="Mean increase in the number of leaves")+
  theme(legend.position = "none", text = element_text(size=12), plot.title = element_text(size=14),
       plot.subtitle = element_text(size=12))
plotgrid0 <- plot_grid(plot_lengthdiffull, plot_wetweightfull, plot_numberdiffull, #combine the 3 plots
                       labels = c("A", "B", "C"),
                       ncol = 3, nrow = 1)
```

Zooming in, the large population-salted and small population-salted groups will be compared using a t-test for the wet weight and length-increase and a Wilcoxon test for the increase in number of leaves (as this group's values were not normally distributed):

```
data_salted <- subset(data_complete, Treatment == "Salted")
leveneTest(data_salted$Wet_weight ~ as.factor(Size), data = data_salted) #Pr(>F) = 0.3754 which means e
```

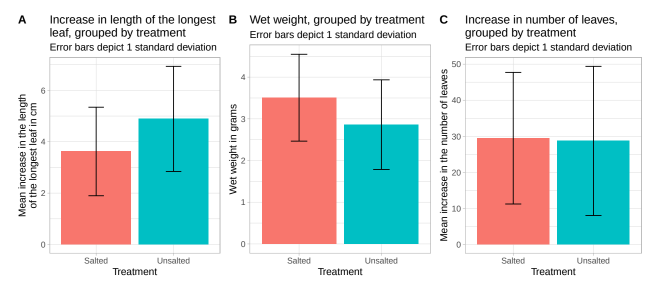


Figure 1: Figure 1: Bar graphs showing A) the increase in longest leaf length grouped by treatment, B) the increase in number of leaves grouped by treatment, and C) the increase in number of leaves grouped by treatment, with error bars depicting 1 standard deviation.

```
## Levene's Test for Homogeneity of Variance (center = median)
##
         Df F value Pr(>F)
  group
         1
            0.8061 0.3754
##
         35
t.test(formula = data_salted$Wet_weight ~ data_salted$Size,
       paired = FALSE, var.equal = TRUE) p.value \%\% round(.,3) #t-test, p = 0.848, which means there i
## [1] 0.848
data_salted <- subset(data_complete, Treatment == "Salted")</pre>
leveneTest(data_salted$Length_dif ~ as.factor(Size), data = data_salted) #Pr(>F) = 0.7135 which means e
## Levene's Test for Homogeneity of Variance (center = median)
##
         Df F value Pr(>F)
## group 1
             0.1369 0.7135
##
         36
t.test(formula = data_salted$Length_dif ~ data_salted$Size,
       paired = FALSE, var.equal = TRUE)p.value \%\% round(.,3) #t-test, p = 0.563, which means there i
## [1] 0.563
wilcox.test(Number_dif ~ Size, data = data_salted) #Wilcoxon test, p = 0.4735, which means there is no
##
   Wilcoxon rank sum test with continuity correction
##
##
## data: Number_dif by Size
## W = 205, p-value = 0.4735
## alternative hypothesis: true location shift is not equal to 0
The data can be plotted, where it's visualized how little difference there is:
```

Generating the figures

```
summary_salted <- data_salted %>%
  group_by(Size) %>%
  summarize(mean_noleaves.increase = mean(Number_dif, na.rm = TRUE),
            mean_lengthleaves.increase = mean(Length_dif, na.rm = TRUE),
            mean_wetweight = mean(Wet_weight, na.rm = TRUE),
            stdevlengthleave.increase = sd(Length_dif, na.rm = TRUE),
            stdevwetweight = sd(Wet_weight, na.rm = TRUE),
            stdevNumber = sd(Number dif, na.rm = TRUE))
plot_length <- summary_salted %% #plot the increase in longest leaf length grouped by population size
  ggplot(aes(x = Size, y = mean_lengthleaves.increase))+
  geom_col(aes(fill = Size))+
  geom_errorbar(aes(ymin = mean_lengthleaves.increase - stdevlengthleave.increase,
                    ymax = mean_lengthleaves.increase + stdevlengthleave.increase), width=.2)+
  theme_light()+
  labs(title = "Increase in length of the longest\nleaf, grouped by population",
       subtitle = "Error bars depict 1 standard deviation, \nplot only showing salted plants",
       x="Population size",
       y="Mean increase in the length\nof the longest leaf in cm")+
  theme(legend.position = "none", text = element_text(size=12), plot.title = element_text(size=14),
        plot.subtitle = element_text(size=12))+
  scale_fill_manual(values=c("#e457b5", "#57e486"))
plot_noleaves <- summary_salted %>% #plot the increase in number of leaves grouped by population
  ggplot(aes(x = Size, y = mean_noleaves.increase))+
  geom col(aes(fill = Size))+
  geom_errorbar(aes(ymin = mean_noleaves.increase - stdevNumber,
                    ymax = mean_noleaves.increase + stdevNumber), width=.2)+
  theme light()+
  labs(title = "Increase in number of leaves, \ngrouped by population size",
      subtitle = "Error bars depict 1 standard deviation,\nplot only showing salted plants",
       x="Population size",
       y="Mean increase in the number of leaves")+
  theme(legend.position = "none", text = element_text(size=12), plot.title = element_text(size=15),
        plot.subtitle = element_text(size=12))+
  scale_fill_manual(values=c("#e457b5", "#57e486"))
plot_wetweight <- summary_salted %>% #plot the increase in number of leaves grouped by population
  ggplot(aes(x = Size, y = mean_wetweight)) +
  geom_col(aes(fill = Size))+
  geom_errorbar(aes(ymin = mean_wetweight - stdevwetweight,
                    ymax = mean_wetweight + stdevwetweight), width=.2)+
  theme light()+
  labs(title = "Wet weight of the plants, \ngrouped by population size",
       subtitle = "Error bars depict 1 standard deviation, \nplot only showing salted plants",
      x="Population size",
       y="Mean wet weight in grams")+
  theme(legend.position = "none", text = element_text(size=12), plot.title = element_text(size=15),
       plot.subtitle = element_text(size=12))+
  scale_fill_manual(values=c("#e457b5", "#57e486"))
plotgrid1 <- plot_grid(plot_length, plot_wetweight, plot_noleaves, #combine the 3 plots into 1 figure
          labels = c("A", "B", "C"),
```

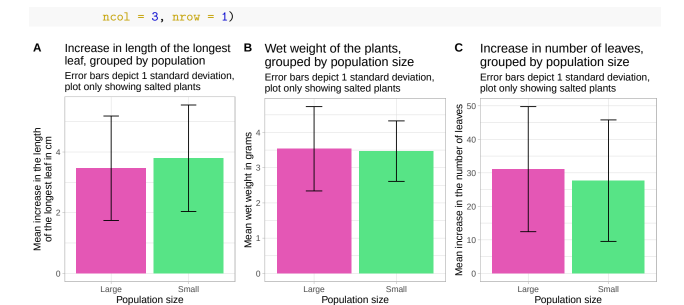


Figure 2: Figure 2: Bar graphs showing A) the increase in longest leaf length grouped by population size, B) the increase in number of leaves grouped by population size, and C) the increase in number of leaves grouped by population size, with error bars depicting 1 standard deviation, and the data filtered to only show the salted groups.

Population size is only used as a proxy for inbreeding, but the genotypes of these plants are still unknown. This is why the populations need to be looked at individually as well:

```
data_complete %>%
  group_by(Population) %>%
  summarise(ShapiroWilk_p.value = shapiro.test(Length_dif)$p.value) #Shapiro-Wilk_test, all p-values ar
## # A tibble: 4 x 2
##
     Population
                  ShapiroWilk_p.value
##
     <chr>>
                                 <dbl>
## 1 Bovra
                                 0.717
## 2 Helin
                                 0.442
                                 0.945
## 3 Smadalen
                                 0.909
## 4 Spiterstulen
data_complete %>%
  group_by(Population) %>%
  summarise(ShapiroWilk_p.value = shapiro.test(Wet_weight)$p.value) #Shapiro-Wilk test, all p-values ar
## # A tibble: 4 x 2
##
     Population
                  ShapiroWilk_p.value
     <chr>
##
                                 <dbl>
## 1 Bovra
                                 0.532
## 2 Helin
                                 0.646
## 3 Smadalen
                                 0.301
                                 0.276
## 4 Spiterstulen
data_complete %>%
  group_by(Population) %>%
  summarise(ShapiroWilk_p.value = shapiro.test(Number_dif)$p.value) #Shapiro-Wilk test, one of the p-va
```

```
## # A tibble: 4 x 2
##
    Population ShapiroWilk_p.value
##
     <chr>
                                <dbl>
                               0.955
## 1 Bovra
## 2 Helin
                               0.316
## 3 Smadalen
                               0.124
## 4 Spiterstulen
                               0.0147
leveneTest(data_complete$Length_dif ~ as.factor(Population), data = data_complete) #Pr(>F) = 0.5092, wh
## Levene's Test for Homogeneity of Variance (center = median)
        Df F value Pr(>F)
## group 3 0.7798 0.5092
##
        70
leveneTest(data_complete$Wet_weight ~ as.factor(Population), data = data_complete) #Pr(>F) = 0.5227, wh
## Levene's Test for Homogeneity of Variance (center = median)
        Df F value Pr(>F)
## group 3
             0.756 0.5227
##
         69
aov(Length_dif ~ Population, data_complete) %% summary.aov() #Pr(>F) = 0.0635, which means there is no
              Df Sum Sq Mean Sq F value Pr(>F)
               3 28.06
                           9.354
                                  2.538 0.0635 .
## Population
## Residuals
              70 257.95
                           3.685
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
aov(Wet_weight ~ Population, data_complete) %>% summary.aov() #Pr(>F) = 0.397, which means there is no
##
              Df Sum Sq Mean Sq F value Pr(>F)
## Population
               3 3.64
                          1.214
                                 1.003 0.397
## Residuals
              69 83.47
                           1.210
## 1 observation deleted due to missingness
kruskal.test(Number_dif ~ Population, data = data_complete) #Kruskal-Wallis test, p = 0.0002492, which
##
## Kruskal-Wallis rank sum test
##
## data: Number_dif by Population
## Kruskal-Wallis chi-squared = 19.194, df = 3, p-value = 0.0002492
summary_complete_pop <- data_complete %>%
  group_by(Population) %>%
  summarize(mean_noleaves.increase = mean(Number_dif, na.rm = TRUE),
           mean_lengthleaves.increase = mean(Length_dif, na.rm = TRUE),
           mean_wetweight = mean(Wet_weight, na.rm = TRUE),
           stdevlengthleave.increase = sd(Length_dif, na.rm = TRUE),
            stdevwetweight = sd(Wet_weight, na.rm = TRUE),
            stdevNumber = sd(Number_dif, na.rm = TRUE))
```

And again, these can be plotted:

Plotting more figures

```
ggplot(aes(x = Population, y = mean_lengthleaves.increase))+
  geom_col(aes(fill = Population), color = c("#e457b5", "#e457b5", "#57e486", "#57e486"), linewidth = 2
  geom_errorbar(aes(ymin = mean_lengthleaves.increase - stdevlengthleave.increase,
                    ymax = mean_lengthleaves.increase + stdevlengthleave.increase), width=.2)+
  theme light()+
  labs(title = "Increase in length of the longest\nleaf, grouped by population",
       subtitle = "Error bars depict 1 standard deviation",
       x="Population",
       y="Mean increase in the length\nof the longest leaf in cm")+
  theme(legend.position = "none", text = element_text(size=12), plot.title = element_text(size=14),
        plot.subtitle = element_text(size=12))+
  scale_fill_manual(values=c("#8111ee", "#7eee11", "#f0860f", "#0f79f0"))
plot_noleaves2 <- summary_complete_pop %>% #plot the increase in number of leaves grouped by population
  ggplot(aes(x = Population, y = mean_noleaves.increase))+
  geom_col(aes(fill = Population), color = c("#e457b5", "#e457b5", "#57e486", "#57e486"), linewidth = 2
  geom_errorbar(aes(ymin = mean_noleaves.increase - stdevNumber,
                    ymax = mean_noleaves.increase + stdevNumber), width=.2)+
  theme_light()+
  labs(title = "Increase in number of leaves, \ngrouped by population",
       subtitle = "Error bars depict 1 standard deviation",
       x="Population",
       y="Mean increase in the number of leaves")+
  theme(legend.position = "none", text = element_text(size=12), plot.title = element_text(size=15),
        plot.subtitle = element_text(size=12))+
  scale_fill_manual(values=c("#8111ee", "#7eee11", "#f0860f", "#0f79f0"))
plot_wetweight2 <- summary_complete_pop %>% #plot the increase in number of leaves grouped by populatio
  ggplot(aes(x = Population, y = mean_wetweight))+
  geom_col(aes(fill = Population), color = c("#e457b5", "#e457b5", "#57e486", "#57e486"), linewidth = 2
  geom_errorbar(aes(ymin = mean_wetweight - stdevwetweight,
                    ymax = mean_wetweight + stdevwetweight), width=.2)+
  theme_light()+
  labs(title = "Wet weight of the plants,\ngrouped by population",
       subtitle = "Error bars depict 1 standard deviation",
       x="Population",
      y="Mean wet weight in grams")+
  theme(legend.position = "none", text = element_text(size=12), plot.title = element_text(size=15),
        plot.subtitle = element_text(size=12))+
  scale_fill_manual(values=c("#8111ee", "#7eee11", "#f0860f", "#0f79f0"))
plotgrid2 <- plot_grid(plot_length2, plot_wetweight2, plot_noleaves2, #combine the 3 plots into 1 figur
          labels = c("A", "B", "C"),
          ncol = 3, nrow = 1)
Different types of phenotypes were scored, like the color and shape of the leaves. These scores can be
```

plot_length2 <- summary_complete_pop %>% #plot the increase in longest leaf length grouped by populatio

Different types of phenotypes were scored, like the color and shape of the leaves. These scores can be visualized as well:

Generating even more figures

```
data_color <- data_complete %>% #prepare color data for plotting
mutate(Color_graph = case_when(
   Color == 1 ~ "Green",
   Color == 2 ~ "Yellow",
```

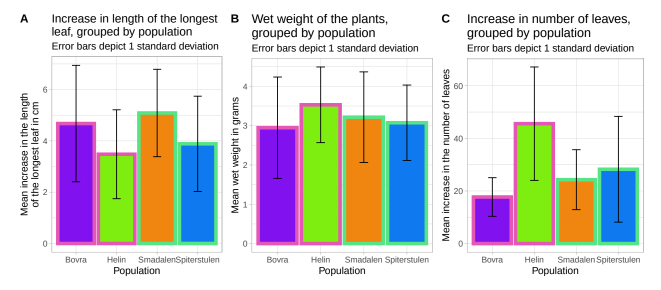


Figure 3: Figure 3: Bar graphs showing A) the increase in longest leaf length grouped by population, B) the increase in number of leaves grouped by population, and C) the increase in number of leaves grouped by population, with error bars depicting 1 standard deviation and the bar borders showing wether the population is large (pink) or small (green).

```
Color == 3 ~ "Red",
    Color == 4 ~ "Red and\nyellow"))
plot_color <- data_color %>% #plot number of leaves of each color, grouped by treatment
  ggplot()+
  geom_bar(aes(x = Color_graph, fill = Treatment), position = position_dodge2(width = 0.9, preserve = "
  theme_light()+
  labs(title = "Number of leaves of each color, \ngrouped by treatment",
       x="Color",
       y="Count")
plot color2 <- data color %>% #plot number of leaves of each color, grouped by population
  ggplot()+
  geom_bar(aes(x = Color_graph, fill = Population, color = Size), linewidth = 1,
           position = position_dodge2(width = 0.9, preserve = "single"))+
  theme light()+
  labs(title = "Number of leaves of each color, \ngrouped by population",
       x="Color",
       v="Count")+
  scale_fill_manual(values=c("#8111ee", "#7eee11", "#f0860f", "#0f79f0"))+
  scale_color_manual(values = c("#e457b5", "#57e486"))+
  guides(color=guide_legend(title="Population size"))
plot_shape_salt <- data_complete %>% #plot number of leaves of each shape, grouped by treatment
  ggplot()+
  geom_bar(aes(x = Leaf_shape, fill = Treatment), position = position_dodge2(width = 0.9, preserve = "s
  theme light() +
  labs(title = "Number of leaves of each shape, \ngrouped by treatment",
       x="Leaf shape",
       y="Count")
```

```
plot_shape_popsize <- data_complete %% #plot number of leaves of each shape, grouped by population siz
  ggplot()+
  geom_bar(aes(x = Leaf_shape, fill = Size), position = position_dodge2(width = 0.9, preserve = "single")
  labs(title = "Number of leaves of each shape, \ngrouped by population size",
       x="Leaf shape",
       y="Count")+
  scale fill manual(values=c("#e457b5", "#57e486"))+
  guides(fill=guide_legend(title="Population size"))
plot_shape_pop <- data_complete %>% #plot number of leaves of each shape, grouped by population
  ggplot()+
  geom_bar(aes(x = Leaf_shape, fill = Population, color = Size), linewidth = 2,
           position = position_dodge2(width = 0.9, preserve = "single"))+
  theme_light() +
  labs(title = "Number of leaves of each shape, \ngrouped by population",
       x="Leaf shape",
       y="Count")+
  scale_fill_manual(values=c("#8111ee", "#7eee11", "#f0860f", "#0f79f0"))+
  scale_color_manual(values = c("#e457b5", "#57e486"))+
  guides(color=guide legend(title="Population size"))
img_leafshapes <- image_read(here("images/leaf_shapes.png")) %% image_ggplot() #import the picture sho
plotgrid3 <- plot_grid(plot_color, plot_color2, plot_shape_salt, img_leafshapes, plot_shape_popsize, pl
          labels = c("A", "B", "C", "D", "E", "F"),
          ncol = 2, nrow = 3)
```

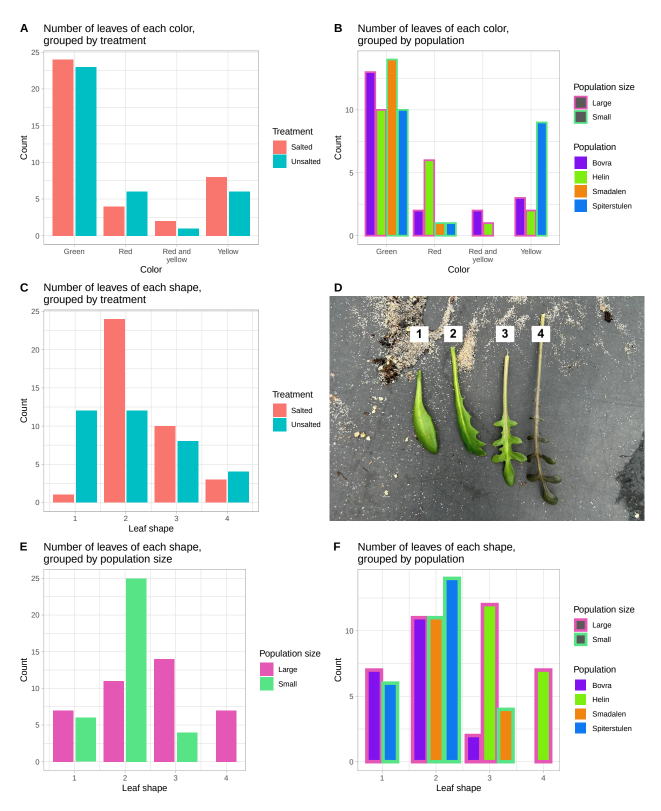


Figure 4: Figure 4: Bar graphs and an image showing A) the number of leaves of each color, grouped by treatment, B) the number of leaves of each color, grouped by population, C) the number of leaves of each shape, grouped by treatment D) the picture showing which shape is equal to which number, E) the number of leaves of each shape, grouped by population size, and F) the number of leaves of each shape, grouped by population.