

# ES 193DS Homework 3

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2024-06-04

Forked Repository: [https://github.com/ellastookey/stookey-ella\\_homework-03](https://github.com/ellastookey/stookey-ella_homework-03)

## Preparations

### Reading in packages

```
# hide messages and warnings
knitr::opts_chunk$set(echo = TRUE, message = FALSE, warning = FALSE)

suppressPackageStartupMessages({
  # general use
  library(tidyverse)
  library(readxl)
  library(here)
  library(janitor)

  # visualizing pairs
  library(GGally)

  # model selection
  library(MuMIn)

  # model predictions
  library(ggeffects)

  # model tables
  library(gtsummary)
  library(flextable)
```

```

library(modelsummary)
library(knitr)
})

drought_exp <- read_xlsx(path = here("data",
                                   "Valliere_etal_EcoApps_Data.xlsx"),
                        sheet = "First Harvest")

# quick look at data
str(drought_exp)

```

```

tibble [70 x 13] (S3: tbl_df/tbl/data.frame)
 $ Species      : chr [1:70] "ENCCAL" "ENCCAL" "ENCCAL" "ENCCAL" ...
 $ Water        : chr [1:70] "WW" "WW" "WW" "WW" ...
 $ Rep #        : num [1:70] 1 2 3 4 5 1 2 3 4 5 ...
 $ Height (cm)  : num [1:70] 5.8 4.9 8.4 6.5 7.1 3.2 4.4 4.2 4.5 3.9 ...
 $ Leaf #       : num [1:70] 11 8 11 12 10 7 7 10 8 6 ...
 $ Leaf dry weight (g): num [1:70] 0.0294 0.0185 0.0177 0.0178 0.0164 0.017 0.0193 0.0153 0.
 $ Leaf area (cm2) : num [1:70] 5.01 3.98 3.69 3.84 3.63 3.06 3.1 2.94 2.73 2.61 ...
 $ SLA          : num [1:70] 170 215 209 216 222 ...
 $ Total LA     : num [1:70] 55.1 31.8 40.6 46.1 36.3 ...
 $ Shoot (g)    : num [1:70] 0.253 0.164 0.241 0.213 0.232 ...
 $ Root (g)     : num [1:70] 0.202 0.165 0.209 0.146 0.12 ...
 $ Total (g)    : num [1:70] 0.455 0.329 0.45 0.359 0.352 ...
 $ R:S          : num [1:70] 0.8 1 0.9 0.7 0.5 0.8 1.2 3.1 0.9 1.2 ...

```

```

class(drought_exp)

```

```

[1] "tbl_df"      "tbl"        "data.frame"

```

## Cleaning

```

# cleaning
drought_exp_clean <- drought_exp %>%
  clean_names() %>% # nicer column names
  mutate(species_name = case_when( # adding column with species scientific names
    species == "ENCCAL" ~ "Encelia californica", # bush sunflower
    species == "ESCCAL" ~ "Eschscholzia californica", # California poppy
    species == "PENCEN" ~ "Penstemon centranthifolius", # Scarlet bugler
  ))

```

```

    species == "GRICAM" ~ "Grindelia camporum", # great valley gumweed
    species == "SALLEU" ~ "Salvia leucophylla", # Purple sage
    species == "STIPUL" ~ "Nasella pulchra", # Purple needlegrass
    species == "LOTSCO" ~ "Acmispon glaber" # deerweed
  )) %>%
  relocate(species_name, .after = species) %>% # moving species_name column after species
  mutate(water_treatment = case_when( # adding column with full treatment names
    water == "WW" ~ "Well watered",
    water == "DS" ~ "Drought stressed"
  )) %>%
  relocate(water_treatment, .after = water) # moving water_treatment column after water

drought_exp_clean

```

# A tibble: 70 x 15

	species	species_name	water	water_treatment	rep_number	height_cm	leaf_number
	<chr>	<chr>	<chr>	<chr>	<dbl>	<dbl>	<dbl>
1	ENCCAL	Encelia calif~	WW	Well watered	1	5.8	11
2	ENCCAL	Encelia calif~	WW	Well watered	2	4.9	8
3	ENCCAL	Encelia calif~	WW	Well watered	3	8.4	11
4	ENCCAL	Encelia calif~	WW	Well watered	4	6.5	12
5	ENCCAL	Encelia calif~	WW	Well watered	5	7.1	10
6	ENCCAL	Encelia calif~	DS	Drought stress~	1	3.2	7
7	ENCCAL	Encelia calif~	DS	Drought stress~	2	4.4	7
8	ENCCAL	Encelia calif~	DS	Drought stress~	3	4.2	10
9	ENCCAL	Encelia calif~	DS	Drought stress~	4	4.5	8
10	ENCCAL	Encelia calif~	DS	Drought stress~	5	3.9	6

# i 60 more rows

# i 8 more variables: leaf\_dry\_weight\_g <dbl>, leaf\_area\_cm2 <dbl>, sla <dbl>,

# total\_la <dbl>, shoot\_g <dbl>, root\_g <dbl>, total\_g <dbl>, r\_s <dbl>

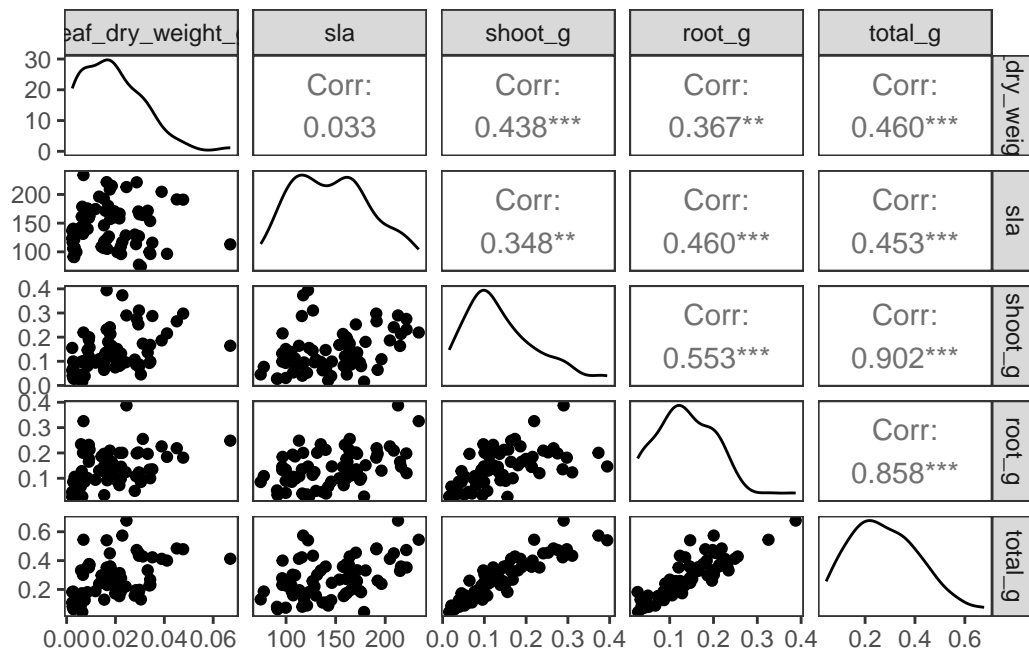
## correlations

```

ggpairs(drought_exp_clean, # data frame
  columns = c("leaf_dry_weight_g", # columns to visualize
    "sla",
    "shoot_g",
    "root_g",
    "total_g"),
  upper = list(method = "pearson")) + # calculating Pearson correlation coefficient

```

```
theme_bw() + # cleaner theme
theme(panel.grid = element_blank()) # getting rid of gridlines
```



```
# bottom left scatterplots of listed variables -- Leaf dry weight on x axis, y axis is total
# upper right shows Pearson's correlation -- positively correlated
```

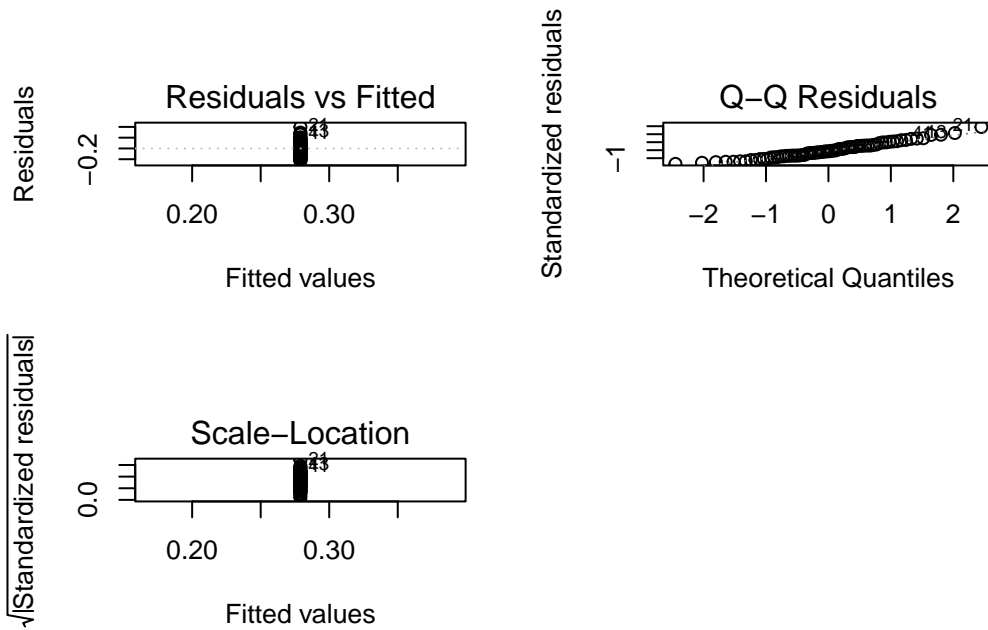
## Problem 1. Multiple linear regression: model selection and construction

### Part a

#### 0. Null model

```
model0 <- lm(total_g ~ 1, # formula
              data = drought_exp_clean) # data frame

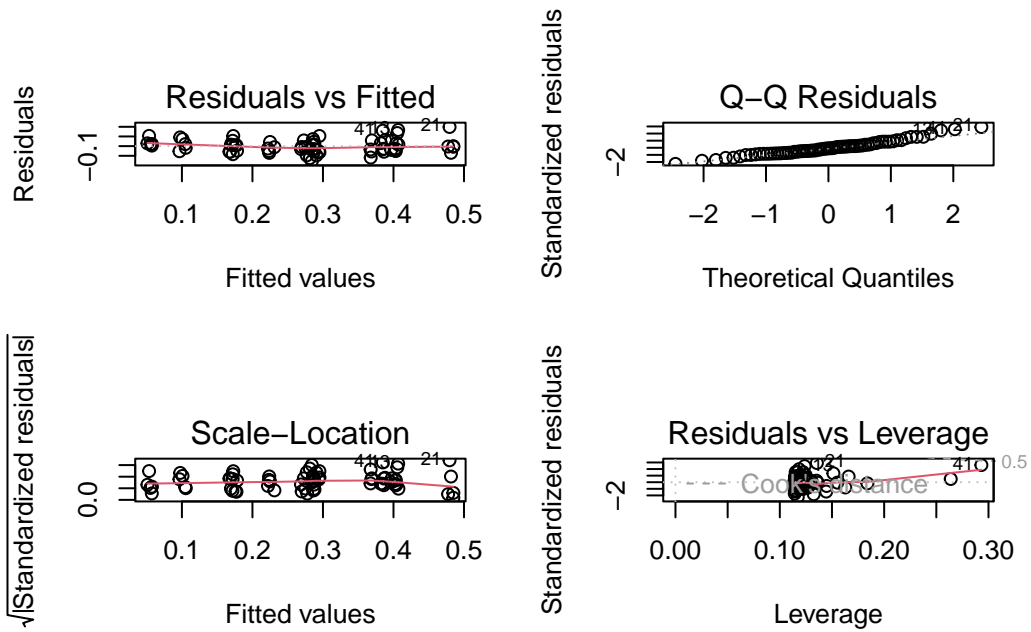
par(mfrow = c(2, 2)) # 2 x 2 format
plot(model0) # display
```



# 1. total biomass as a function of SLA, water treatment, and species

```
# saturated model
model1 <- lm(total_g ~ sla + water_treatment + species_name,
              data = drought_exp_clean)

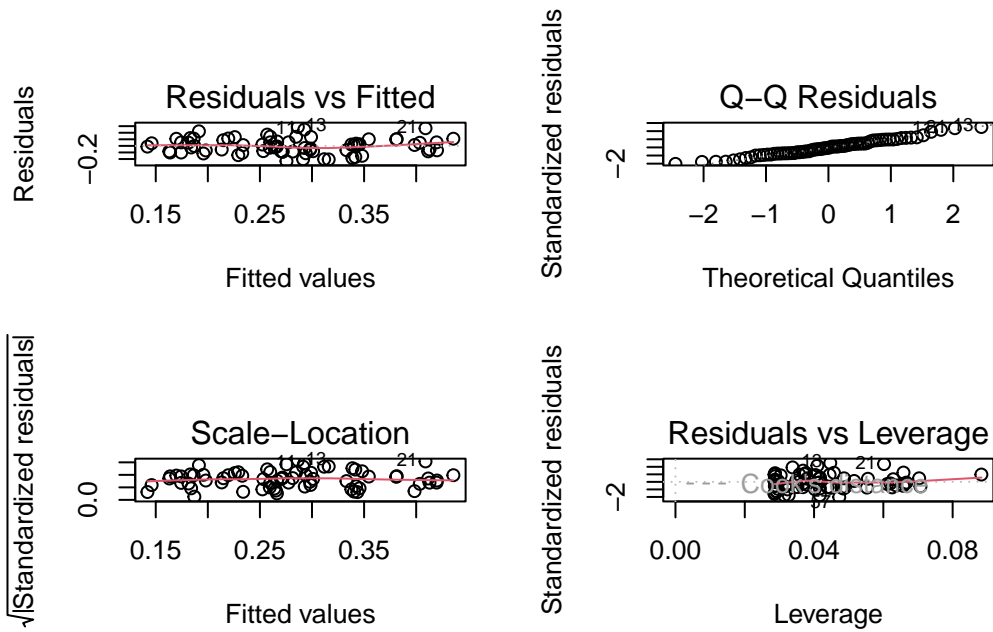
par(mfrow = c(2, 2))
plot(model1)
```



## 2. total biomass as a function of SLA and water treatment

```
model2 <- lm(total_g ~ sla + water_treatment,
              data = drought_exp_clean)

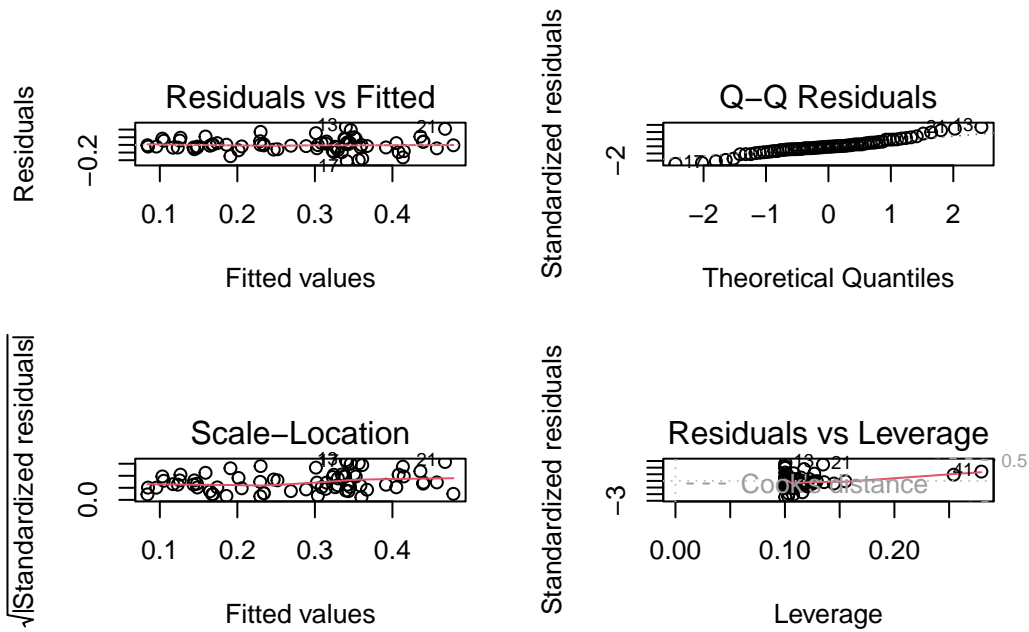
par(mfrow = c(2, 2))
plot(model2)
```



### 3. total biomass as a function of SLA and species

```
model3 <- lm(total_g ~ sla + species_name,
              data = drought_exp_clean)

par(mfrow = c(2, 2))
plot(model3)
```

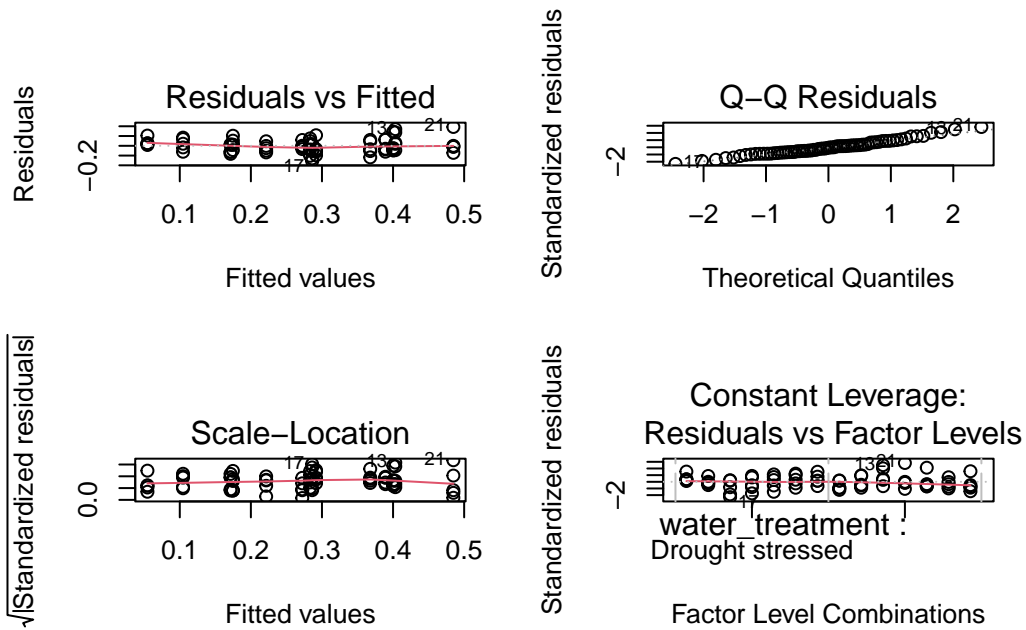


#### 4. total biomass as a function of water treatment and species

```
model4 <- lm(total_g ~ water_treatment + species_name,
              data = drought_exp_clean)

par(mfrow = c(2, 2))
plot(model4)
```





```
# Model selection table to see AIC and delta values
```

```
model.sel(model0,
           model1,
           model2,
           model3,
           model4)
```

Model selection table

	(Int)	sla	spc_nam	wtr_trt	df	logLik	AICc	delta	weight
model4	0.05455			+	9	88.598	-156.2	0.00	0.772
model11	0.07994	-0.0002475		+	10	88.741	-153.8	2.44	0.228
model13	-0.03315	0.0012900		+	9	72.538	-124.1	32.12	0.000
model12	0.04670	0.0012810		+	4	52.220	-95.8	60.37	0.000
model10	0.27900				2	39.580	-75.0	81.22	0.000

Models ranked by AICc(x)

```
# See model 4 coefficients
```

```
summary(model4)
```

Call:

```
lm(formula = total_g ~ water_treatment + species_name, data = drought_exp_clean)
```

Residuals:

Min	1Q	Median	3Q	Max
-0.157087	-0.046953	-0.003733	0.041244	0.192657

Coefficients:

	Estimate	Std. Error	t value	Pr(> t )
(Intercept)	0.05455	0.02451	2.225	0.02973 *
water_treatmentWell watered	0.11695	0.01733	6.746	5.90e-09 ***
species_nameEncelia californica	0.21774	0.03243	6.714	6.70e-09 ***
species_nameEschscholzia californica	0.23164	0.03243	7.143	1.22e-09 ***
species_nameGrindelia camporum	0.31335	0.03243	9.662	5.53e-14 ***
species_nameNasella pulchra	0.22881	0.03243	7.055	1.72e-09 ***
species_namePenstemon centranthifolius	0.05003	0.03243	1.543	0.12799
species_nameSalvia leucophylla	0.12020	0.03243	3.706	0.00045 ***

---

Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 0.07252 on 62 degrees of freedom

Multiple R-squared: 0.7535, Adjusted R-squared: 0.7257

F-statistic: 27.08 on 7 and 62 DF, p-value: < 2.2e-16

## Table presentation

```
# Making a table
model_info <- data.frame(
  Model_Number = c("0 (null)", "1 (saturated)", 2, 3, 4), # label rows by model numbers
  Predictors = c("None", "SLA, Treatment, Species", "SLA, Treatment", "SLA, Species", "Treatment, Species")
) # add predictors to rows
```

```
kable(model_info,
  caption = "Models and their Predictors for Total Plant Biomass. This table shows five models.",
  col.names = c("Model", "Predictors")) # label columns
```

Table 1: Models and their Predictors for Total Plant Biomass. This table shows five linear regression models (0-4) and their predictors. Model 0 is a null model (contains no predictors) and model 1 is saturated (contains all three possible predictors). The other three models consist of a combination of predictors. The model numbers and the predictors are in the rows, while the columns serve as labels for these two categories

Model	Predictors
0 (null)	None
1 (saturated)	SLA, Treatment, Species
2	SLA, Treatment
3	SLA, Species
4	Treatment, Species

## Part b

To examine the influence of specific leaf area, water treatment, and plant species type on total plant biomass, I have constructed and analyzed five linear regression models. The first model (model 0) was a null model meaning that there were no predictors. On the other hand, model 1 was a saturated model and all predictors were present. Models 2-4 each had some combination of predictors to examine their effect on total plant biomass. To determine the model that best described the impact these potential influences have on plant biomass, a model selection table was used. This revealed model 4, which examined the water treatment and plant species variables, to have the lowest Akaike Information Criterion (AIC) of -156.2 and a delta of 0. These values suggest that model 4 is best because it is descriptive, but not too complex. To confirm this, I had to ensure that model 4 conformed to the assumptions of a linear model by examining its diagnostic plots. Beginning with the homoscedastic models, the residuals are scattered along a straight line and have an even distribution above and below the red line indicating constant variance. The Q-Q plot tests if the data is normally distributed, which it appears to be as it follows a linear path. The last plot shows no outliers with significant influence because if there were, there would be a red dashed line with points falling on the outside. Overall, the diagnostic plots confirm the model selection tables' suggestion that model 4 is best. Lastly, model 4's coefficient summary was examined to further solidify the conclusion that it is the best model. After looking at the slope, intercept, and level estimates, model 4 was confirmed best as they showed the significant differences between water treatments and the varying biomasses across plant species.

## Part c

### Model predictions

```
model4_preds <- ggpredict(model4,
                           terms = c(
                               "water_treatment",
                               "species_name"))

# use View(model_preds) to see the predictions as a data frame
# use model_preds to see the predictions formatted nicely

# creating new data frame of model predictions for plotting
model_preds_for_plotting <- model4_preds %>%
  rename(water_treatment = x, # renaming columns to make this easier to use
         species_name = group)

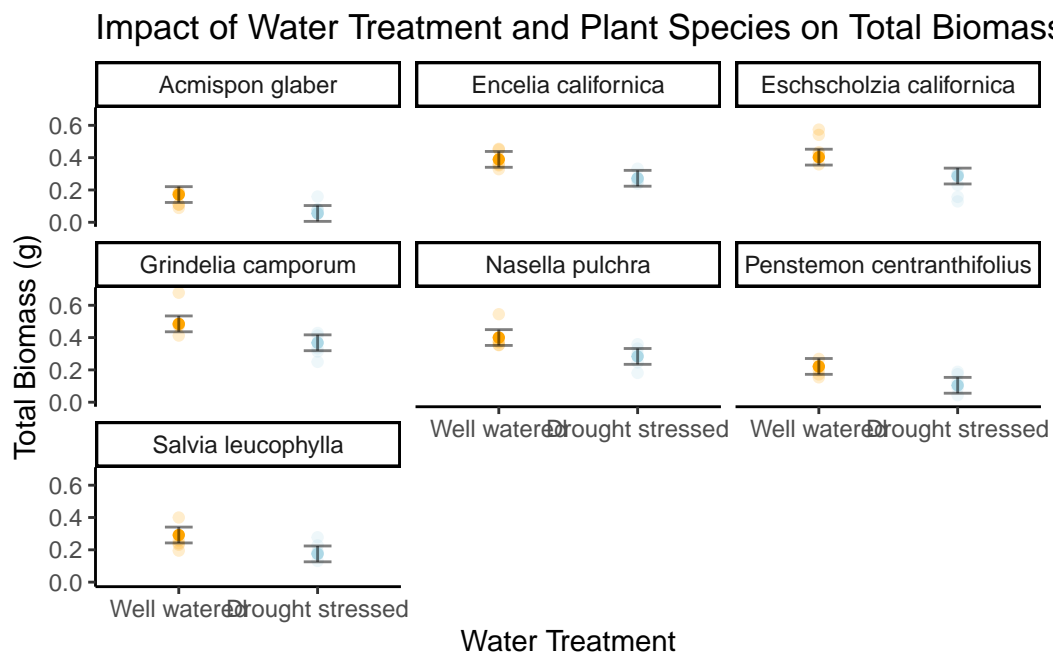
# use View(model_preds_for_plotting)
# to compare this to the original model_preds data frame

ggplot() +
  # underlying data
  geom_point(data = drought_exp_clean,
            alpha = 0.2, # making underlying data transparent
            aes(x = water_treatment,
                y = total_g,
                color = water_treatment)) +
  # add prediction data
  geom_point(data = model_preds_for_plotting,
            aes(x = water_treatment,
                y = predicted,
                color = water_treatment)) +
  # model prediction 95% CI errorbar
  geom_errorbar(data = model_preds_for_plotting,
               aes(x = water_treatment,
                   y = predicted,
                   ymin = conf.low,
                   ymax = conf.high,
                   fill = water_treatment),
               width = 0.2, alpha = 0.5) +
  # cleaner theme
```

```

theme_classic() +
# creating different panels for species
facet_wrap(~species_name) +
theme(legend.position = "none") + # removed legend
labs(title = "Impact of Water Treatment and Plant Species on Total Biomass", # add title
      x = "Water Treatment",
      y = "Total Biomass (g)") + # change plot title and axes labels
scale_color_manual(values = c("Well watered" = "orange",
                              "Drought stressed" = "lightblue")) # change colors

```



## Part d

This figure shows the predicted (darker points) and underlying data (lighter points) of changes in water treatment and varying plant species on total plant biomass. Each plot represents a different species' data, all having well watered (orange) and drought stressed (blue) on the x-axis and total biomass (in grams) on the y-axis. The error bars represent a 95% confidence interval for the predicted values of each species.

Valliere, Justin; Zhang, Jacqueline; Sharifi, M.; Rundel, Philip (2019). Data from: Can we condition native plants to increase drought tolerance and improve restoration success? [Dataset]. Dryad. <https://doi.org/10.5061/dryad.v0861f7>

## Part e

Model 4, the best model and shown above in the figure, has the predictors of water treatment and plant species, both of which best describe total mass. Model 4 has the lowest AIC value ( $AIC = -156.2$ ) of all the models and a delta of 0, suggesting it is the best and most significant model ( $F(7,62) = 27.08$ ,  $p < 0.001$ ,  $\alpha = 0.05$ , adjusted  $R^2 = 0.73$ ). On average between water treatments, those that were well watered tended to have a higher total biomass than those that were drought stressed. *Grindelia camporum* was the species with the highest total biomass for both well watered and drought stressed treatments, however all species had around the same amount of total biomass on average.

## Problem 2. Affective visualization

### Part a

For my personal data set, where I am examining the distance traveled each day, I could use a bar graph and outline the perimeter of each peak. In doing so, the graph will appear to be “hilly”. Since my data is about driving, I will turn this into a scene with a car driving over hills (ie the bar graph).

### Part b

see at end

### Part c

see at end

### Part d

For my visualization, I have created a scene of a car traveling on a hilly road. The hills represent the distance traveled (in miles) each day, some days peaking while others are flatter. This work was done on a digital coloring platform called Notability. I began by importing a screenshot of my data (bar graph) and from there I traced and colored it. I finished it off by adding details, such as the road and cars. During this process I was unsure whether to keep each date and numerical value, however ultimately did decide to keep them because I thought it helped the viewer see the information clearer.

### **Problem 3. Statistical critique**

#### **Part a**

To examine the long-term effects of a wildfire on soil nutrients and makeup, the researchers used a two-way ANOVA test and if significant differences were found ( $p < 0.05$ ), a Tukey HSD post-hoc test was applied. The authors represented these statistical tests in three tables. Table 1 shows the results of the ANOVA test and Tables 2 & 3 show the descriptive statistics for certain nutrients. In addition to the tables, there were two figures. The first was a map that illustrated the study location and areas with varying fire severity. The second figure was an RDA for the relation between factors 1 and 2.

#### **Part b**

All three tables were very clear, with descriptive captions and column and row labels. Figure 1 was also simple to understand because it consisted of images and maps for context. However, figure 2 was significantly more confusing to understand because I have never looked at a redundancy analysis (RDA) plot before. There are no units on the x and y axis and at first glance the numbers seem quite arbitrary. After researching how to read the plot, it made more sense and I could see how the variables' summary statistics (means and standard deviations) were being shown. No model predictions were in the matrix, but rather just the collected data.

#### **Part c**

The tables all hold a lot of information and data making them seem a bit visually cluttered. However, this was crucial information for the researchers to show so it was necessary to include it all. Figure 2, the RDA plot, had a very good data to ink ratio, only consisting of a few lines, two colors, and minimal lettering.

#### **Part d**

For the tables, I think they could have been made clearer if titles/ labels were bolded or a larger font. This would help differentiate the organizational aspects from the large amounts of data. Additionally having lines in the table would have helped section off different nutrients' information instead of it all blending together. As for the figure, I wish the caption included more information about what each percent on the sides of the plot and the axes represent. Although information about this is included in the text of the paper, I think including it on the figure is crucial so the reader can get a clear picture of the results from the experiment.