# Homework 5

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# **Loading Packages**

```
# should haves (from last week)
library(tidyverse)
## -- Attaching core tidyverse packages ----- tidyverse 2.0.0 --
## v dplyr
             1.1.1
                        v readr
                                     2.1.4
## v forcats 1.0.0
                                     1.5.0
                         v stringr
## v ggplot2 3.4.2
                        v tibble
                                     3.2.1
## v lubridate 1.9.2
                         v tidyr
                                     1.3.0
## v purrr
              1.0.1
                                         ------tidyverse_conflicts() --
## -- 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(here)
## here() starts at /Users/luketodd/Desktop/Rprojects/ES 193DS/ES 193DS Homework 5
library(janitor)
##
## Attaching package: 'janitor'
## The following objects are masked from 'package:stats':
##
##
       chisq.test, fisher.test
library(ggeffects)
library(performance)
library(naniar) # or equivalent
library(flextable) # or equivalent
##
## Attaching package: 'flextable'
## The following object is masked from 'package:purrr':
##
##
       compose
```

```
library(car)
## Loading required package: carData
##
## Attaching package: 'car'
## The following object is masked from 'package:dplyr':
##
##
       recode
##
## The following object is masked from 'package:purrr':
##
##
       some
library(broom)
# would be nice to have
library(corrplot)
## corrplot 0.92 loaded
library(AICcmodavg)
library(GGally)
\mbox{\tt \#\#} Registered S3 method overwritten by 'GGally':
##
     method from
##
            ggplot2
     +.gg
Loading Data
sar <- read.csv(here("Data/hf109-01-sarracenia.csv")) %>%
  clean_names() %>%
  select(totmass, species, feedlevel, sla, chlorophyll, amass, num_lvs, num_phylls)
7a
7b
7c
gg_miss_var(sar)
# creating a dataset without the missing values
sar_nona <- sar %>%
  drop_na(sla, chlorophyll, amass, num_lvs, num_phylls)
```

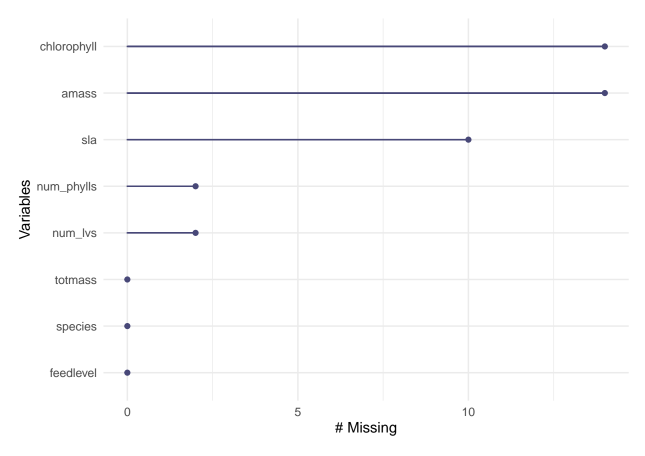


Figure 1: This figure displays the number of missing values for each variable in the sar data frame. Based on this figure, we can see that chlorophyll (Chlorophyll content), amass (Photosynthetic Rate), and sla (Specific Leaf Area) are missing the most values with 14, 14, and 10 missing values, respectively. Num\_phylls and num\_lvs are missing 2 values, and the rest are missing none.

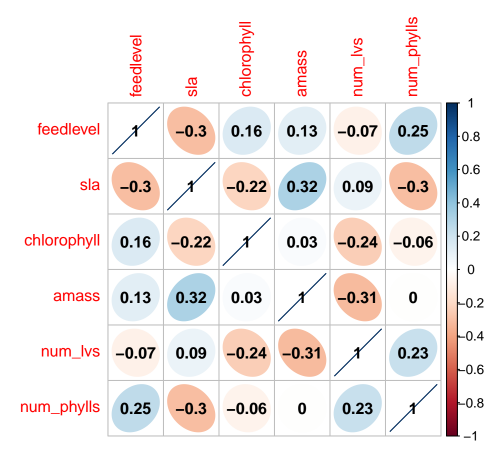
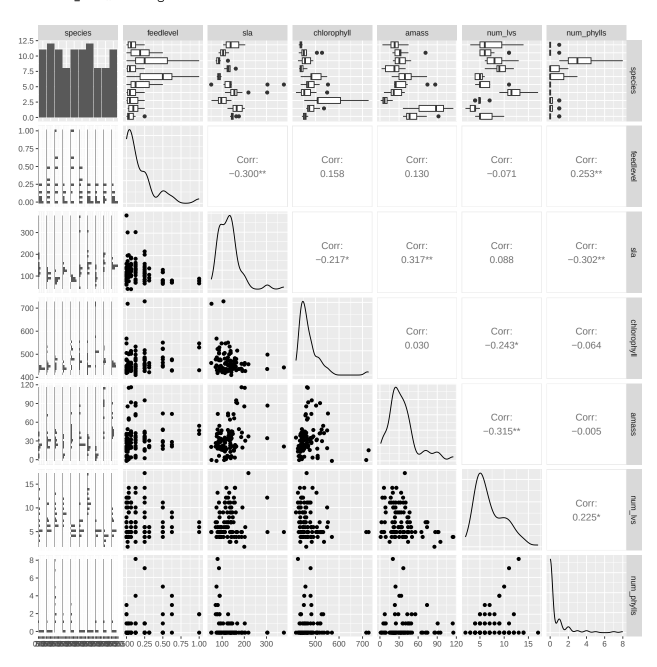


Figure 2: This figure displays the Person's correlation values between different variables. High absolute values means that there is a greater correlation between the variables. For example, sla and amass have the greatest positive correlation between each other with a value of 0.32. On the otherhand, num\_lvs and amass have the greatest negative correlation between each other with a value of -0.31.

**7e** 

```
sar_nona %>%
  select(species:num_phylls) %>%
  ggpairs()
```

```
## 'stat_bin()' using 'bins = 30'. Pick better value with 'binwidth'.
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```

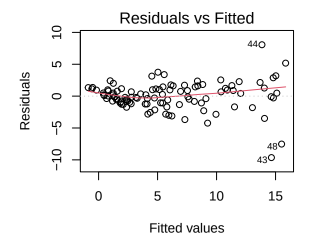


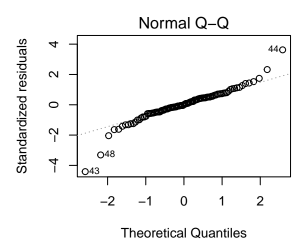
### 7f

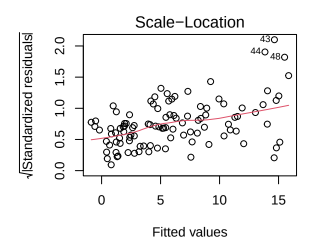
As we are trying to predict totmass, the y-value is set to totmass. For the null model, we set the x-value to 1, as this selects for just the intercept. For the full model, we select every variable that we are interested in.

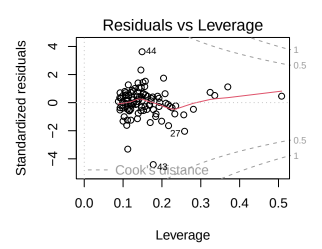
# 7g/h

```
# visual diagnostics for full model
par(mfrow = c(2, 2))
plot(full)
```









# statistical diagnostic checks
check\_normality(full)

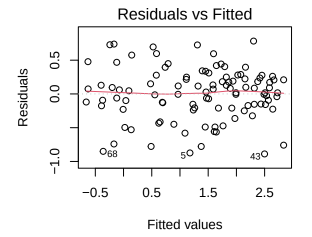
## Warning: Non-normality of residuals detected (p < .001).

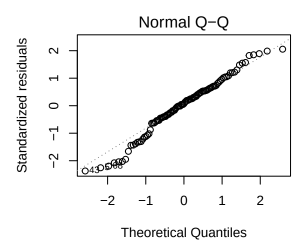
```
check_heteroscedasticity(full)
```

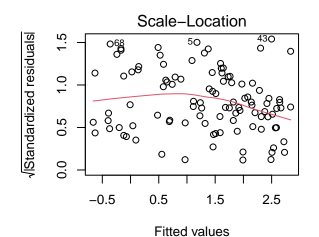
## Warning: Heteroscedasticity (non-constant error variance) detected (p < .001).

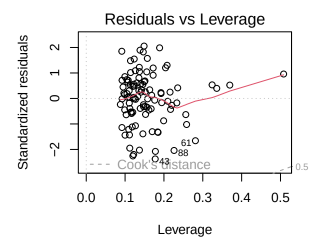
Checking normality and homoscedasticity assumptions both gave p-values less than 0.001, indicating that our current full model is non-normal and heteroscedastic. Because of this, we will log-transform our model and retest the diagnostics.

```
# visual diagnostic checks
par(mfrow = c(2, 2))
plot(full_log)
```









```
# statistical diagnostic checks
check_normality(full_log)

## OK: residuals appear as normally distributed (p = 0.107).
check_heteroscedasticity(full_log)
```

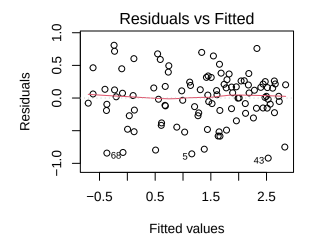
## OK: Error variance appears to be homoscedastic (p = 0.071).

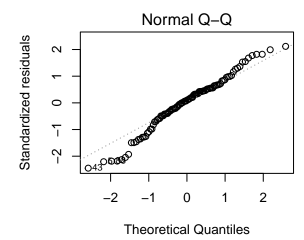
The statistical diagnostic checks for normality and homoscedasticity passed, so we will continue using the log-transformed model.

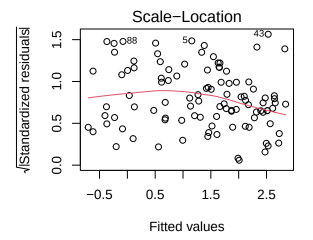
### 7i

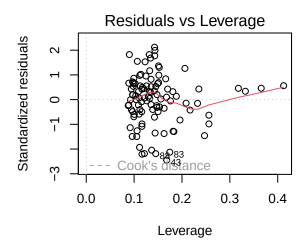
# using ANOVA tables to create new models, eliminating one non-significant variable at a time anova(full\_log)

```
## Analysis of Variance Table
##
## Response: log(totmass)
##
              Df Sum Sq Mean Sq F value
                                          Pr(>F)
               9 91.175 10.1305 59.3944 < 2.2e-16 ***
## species
## feedlevel
             1 0.110 0.1099 0.6442 0.4243716
              1 1.300 1.3003 7.6236 0.0070259 **
## sla
## chlorophyll 1 2.564 2.5636 15.0301 0.0002049 ***
## amass
               1 0.033 0.0330 0.1937 0.6609205
## num_lvs
              1 2.921 2.9205 17.1227 8.068e-05 ***
## num_phylls 1 0.100 0.0999 0.5859 0.4460675
## Residuals 87 14.839 0.1706
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
# based on this ANOVA, I will create a model without "amass" since it had the highest p-value
model2 <- lm(log(totmass) ~ species + feedlevel + sla + chlorophyll + num_lvs + num_phylls,</pre>
            data = sar_nona)
# checking model2 visual diagnostics
par(mfrow = c(2, 2))
plot(model2)
```









# checking model statistical diagnostics
check\_normality(model2)

## OK: residuals appear as normally distributed (p = 0.075).

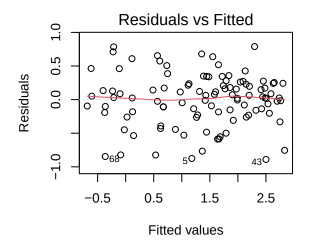
check\_heteroscedasticity(model2)

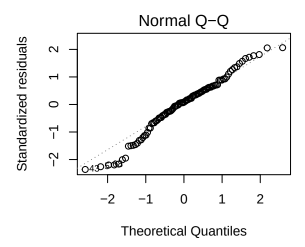
## OK: Error variance appears to be homoscedastic (p = 0.054).

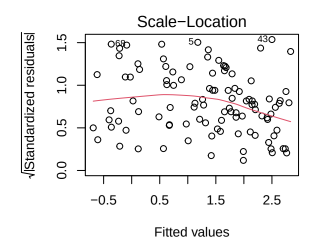
Based on the visual and statistical diagnostic test, model2 passes all tests. Next, I will eliminate another variable and see if it still passes these tests.

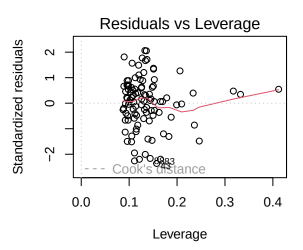
## anova(model2)

```
0.110 0.1099 0.6471 0.4233265
## feedlevel
## sla
                   1.300
                          1.3003 7.6574 0.0068914 **
                1
                          2.5636 15.0966 0.0001975 ***
## chlorophyll
                   2.564
                   2.866
                          2.8665 16.8803 8.903e-05 ***
## num_lvs
## num_phylls
                1
                   0.083
                          0.0827
                                  0.4868 0.4872174
## Residuals
               88 14.943
                          0.1698
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
# based on the above ANOVA, I will create a model without "num_phylls" since it had the highest p-value
model3 <- lm(log(totmass) ~ species + feedlevel + sla + chlorophyll + num_lvs,</pre>
             data = sar_nona)
# checking model2 visual diagnostics
par(mfrow = c(2, 2))
plot(model3)
```









# checking model statistical diagnostics
check\_normality(model3)

## OK: residuals appear as normally distributed (p = 0.062).

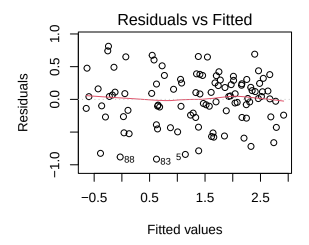
```
check_heteroscedasticity(model3)
```

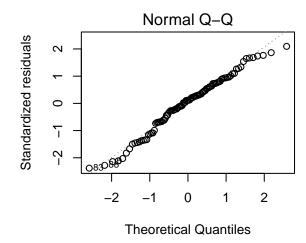
## OK: Error variance appears to be homoscedastic (p = 0.067).

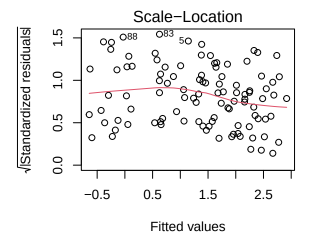
Based on the visual and statistical diagnostic test, model3 passes all tests. Next, I will eliminate another variable and see if it still passes these tests.

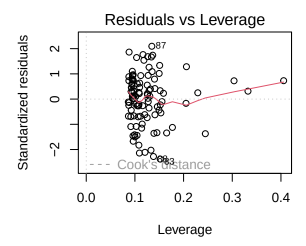
#### anova(model3)

```
## Analysis of Variance Table
##
## Response: log(totmass)
##
              Df Sum Sq Mean Sq F value
                                          Pr(>F)
## species
               9 91.175 10.1305 60.0035 < 2.2e-16 ***
## feedlevel
             1 0.110 0.1099 0.6508 0.4219656
              1 1.300 1.3003 7.7018 0.0067230 **
## sla
## chlorophyll 1 2.564 2.5636 15.1842 0.0001885 ***
## num_lvs
              1 2.866 2.8665 16.9782 8.457e-05 ***
## Residuals 89 15.026 0.1688
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
# based on the above ANOVA, I will create a model without "feedlevel" since it had the highest p-value
model4 <- lm(log(totmass) ~ species + sla + chlorophyll + num_lvs,</pre>
            data = sar_nona)
# checking model2 visual diagnostics
par(mfrow = c(2, 2))
plot(model4)
```









# checking model statistical diagnostics
check\_normality(model4)

## OK: residuals appear as normally distributed (p = 0.169).

check\_heteroscedasticity(model4)

## Warning: Heteroscedasticity (non-constant error variance) detected (p = 0.036).

The statistical tests show that model appears to have heteroscedasticity, or non-constant error variance, with a p-value of 0.036.

The type of model selection that I used above is called backward model selection. It begins at a full model and you slowly eliminate the least significant variable, until you are left with a model that passes all diagnostics, but has the least amount of predictor variables. Most of the time, it is better to have the least amount of predictor variables as possible since it increases the interpretability of your model. Based on the above method, "model3" appears to be our best model.

# # full model variance inflation factor check car::vif(full\_log)

```
GVIF Df GVIF^(1/(2*Df))
##
## species
               42.351675
                                    1.231351
## feedlevel
                1.621993
                                    1.273575
                          1
## sla
                1.999989
                          1
                                    1.414210
## chlorophyll
                1.949828
                                    1.396362
                          1
## amass
                2.872084
                                    1.694722
## num_lvs
                2.813855
                                    1.677455
                          1
## num_phylls
                2.995510
                                    1.730754
```

Based on the results, we can see that every variable has a GVIF value greater than 1, indicating that there is some multicollinearity. Ideally, you want to be as close as possible to 1.

#### 7k

```
# comparing models using AIC
MuMIn::AICc(full_log, null, model2, model3, model4)

## df AICc
## full_log 17 133.9424
## null 2 630.4028
## model2 16 131.7899
## model3 15 129.5498
```

The above chart shows that model3 has the lowest AICc value. Therefore, we will choose model3 as the best model.

#### 8a

## model4

14 130.8121