ENTMLGY 6707 Entomological Techniques and Data Analysis R Activity 2: Data Wrangling

1 Introduction

We are going to get some practice cleaning, exploring, and graphing data. Quality control checks are extremely important before you begin any formal analysis (e.g., are all the rows and columns loaded? are the missing values identified correctly?). For the actual activity, head to the last page. Otherwise, this document is comprised of advice on and examples of coding and analyses.

2 Loading in data and initial checks

Typically, we start by loading in some data (at which you all are now professionals)

It is a good idea to look at the summary(). I often use nrow(data_name) to examine the number of rows I have loaded.

```
summary(iris df)
     Sepal.Length
##
                      Sepal.Width
                                       Petal.Length
                                                        Petal.Width
           :4.300
##
    Min.
                     Min.
                            :2.000
                                      Min.
                                              :1.000
                                                       Min.
                                                              :0.100
##
    1st Qu.:5.100
                     1st Qu.:2.800
                                      1st Qu.:1.600
                                                       1st Qu.:0.300
   Median :5.800
                     Median :3.000
                                      Median :4.350
                                                       Median :1.300
##
    Mean
           :5.843
                     Mean
                            :3.057
                                      Mean
                                              :3.758
                                                       Mean
                                                              :1.199
##
    3rd Qu.:6.400
                     3rd Qu.:3.300
                                      3rd Qu.:5.100
                                                       3rd Qu.:1.800
##
           :7.900
                                              :6.900
                                                              :2.500
    Max.
                     Max.
                            :4.400
                                      Max.
                                                       Max.
##
      Species
##
    Length: 150
##
   Class : character
##
   Mode :character
##
##
##
nrow(iris_df)
```

[1] 150

You may also want to look at the structure of the data using str(), which provides insights into number of observations (= number of rows), types of variables (numeric vs. character vs. logical[TRUE/FALSE]), and the first few rows of data.

str(iris_df)

```
## tibble [150 x 5] (S3: tbl_df/tbl/data.frame)
## $ Sepal.Length: num [1:150] 5.1 4.9 4.7 4.6 5 5.4 4.6 5 4.4 4.9 ...
## $ Sepal.Width : num [1:150] 3.5 3 3.2 3.1 3.6 3.9 3.4 3.4 2.9 3.1 ...
## $ Petal.Length: num [1:150] 1.4 1.4 1.3 1.5 1.4 1.7 1.4 1.5 1.4 1.5 ...
## $ Petal.Width : num [1:150] 0.2 0.2 0.2 0.2 0.2 0.4 0.3 0.2 0.2 0.1 ...
## $ Species : chr [1:150] "setosa" "setosa" "setosa" "setosa" ...
```

It's a good idea to ensure you've correctly loaded in the top and bottom of the data (i.e., that your entire rectangular spreadsheet arrived safely into R).

```
head(iris_df)
```

```
## # A tibble: 6 x 5
##
     Sepal.Length Sepal.Width Petal.Length Petal.Width Species
##
             <dbl>
                          <dbl>
                                        <dbl>
                                                     <dbl> <chr>
## 1
               5.1
                            3.5
                                                       0.2 setosa
                                          1.4
## 2
               4.9
                            3
                                          1.4
                                                       0.2 setosa
## 3
               4.7
                            3.2
                                          1.3
                                                       0.2 setosa
## 4
               4.6
                            3.1
                                          1.5
                                                       0.2 setosa
## 5
               5
                            3.6
                                          1.4
                                                       0.2 setosa
## 6
               5.4
                            3.9
                                          1.7
                                                       0.4 setosa
tail(iris_df)
```

```
## # A tibble: 6 x 5
```

```
##
     Sepal.Length Sepal.Width Petal.Length Petal.Width Species
##
             <dbl>
                          <dbl>
                                        <dbl>
                                                     <dbl> <chr>
## 1
               6.7
                            3.3
                                          5.7
                                                       2.5 virginica
## 2
               6.7
                            3
                                          5.2
                                                       2.3 virginica
## 3
                            2.5
                                          5
                                                       1.9 virginica
               6.3
## 4
               6.5
                            3
                                          5.2
                                                       2
                                                            virginica
               6.2
                            3.4
## 5
                                          5.4
                                                       2.3 virginica
## 6
               5.9
                                          5.1
                                                       1.8 virginica
```

The below code opens up a new tab in R studio displaying your entire data frame. It is nice to view the data this way, but you typically still need the above functions to find erroneous NA values, typographical errors, missing rows/columns, etc.

```
View(iris_df)
```

3 Converting variables

As a reminder, you can access specific columns in data frames using a \$.

```
head(iris_df$Species)
```

```
## [1] "setosa" "setosa" "setosa" "setosa" "setosa" "setosa"
```

It is very common in practice to convert a variable from a number to a factor. For example, maybe you recorded your sites as 1,2,3...R will read that column (incorrectly, as far as we are concerned) as numeric. In the tidyverse (of which readxl is a part), some factors are read in as characters. You will notice chr in the str() output above for the Species column, indicating that Species is a character according to R.

You can quickly convert such variables to factors. Notice below how the output of summary() changes after the conversion. Specifying the variable type incorrectly can cause a lot of problems when it comes to graphing and analysis, so always check that R has correctly identified numbers as numeric and categorical variables as factors (or characters). The difference(s) between factors and characters in R will not cause problems when graphing and analyzing, as the main difference is just that factors have predefined levels. This distinction can cause problems if you are trying to add a new level to an existing factor, but that is so uncommon we are not going to worry about the differences in this course again.

```
summary(iris df$Species)
##
      Length
                  Class
                              Mode
##
         150 character character
is.character(iris_df$Species)
## [1] TRUE
iris_df$Species <- as.factor(iris_df$Species)</pre>
summary(iris_df$Species)
##
       setosa versicolor
                           virginica
##
           50
```

4 Logical operators

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R has several built-in logical operators that can make it easier to access subsets of data.

Let's create a logical vector called sepal_greater5 and feed it to table(). We are asking each value of iris_df\$Sepal.Length if it is greater than 5 and getting an answer of TRUE or FALSE.

```
sepal_greater5 <- iris_df$Sepal.Length > 5
table(sepal_greater5)

## sepal_greater5
## FALSE TRUE
## 32 118
```

Let's do something similar, asking if each value of Sepal.Lenth is equal to 5.1.

```
sepal_equal5.1 <- iris_df$Sepal.Length == 5.1
table(sepal_equal5.1)

## sepal_equal5.1
## FALSE TRUE</pre>
```

You should notice that our output is a string of TRUE and FALSE values exactly equal to the length of iris_df\$Sepal.Length.

Note the == means "exactly equal to" whereas = is often used as a substitute for <-. Below is a quick example. Also note the use of semicolons (;), which you can use to split up code on the same line. I use it when I make graphs or create vectors and want the output printed immediately every time I run the code. Use it sparingly.

```
x <- 5+5;x
## [1] 10
```

```
y = 7+3;y

## [1] 10

y==x

## [1] TRUE
```

One other operator I frequently use is %in%, which you read aloud as 'that occurs in'. It can come in really handy when extracting information from a data set. This code determines if each observation of Sepal.Length equals 5.1, 4.6 or 6.0, again returning a TRUE or FALSE.

```
head(iris_df$Sepal.Length %in% c(5.1,4.6,6.0))
```

```
## [1] TRUE FALSE FALSE TRUE FALSE FALSE
```

I will often use it with the which() command. Instead of returning TRUE or FALSE, this now returns the location of observations for which the operator is TRUE.

```
head(which(iris_df$Sepal.Length %in% c(5.1,4.6,6.0)))
## [1] 1 4 7 18 20 22
```

5 Subsetting data using base R

You will often need to inspect or extract subsets of your data, and here we'll cover several useful functions for doing so. We can extract specific rows and columns by entering the name of a data frame followed by brackets: <code>iris_df[,]</code>. There are two "arguments" inside the brackets, the first indicates which rows you want and the second indicates which columns you want:

```
iris_df[25,2:3] # row 25, columns 2 and 3
iris_df[c(26,31),] # rows 26 and 31 for all columns
iris_df[,] # when left blank, the entire dataframe is returned
iris_df[-5,1:2] # remove the 5th row, return columns 1 and 2 for everything left
iris_df[which(iris_df$Sepal.Length>5),] # returns observations with Sepal.Length > 5
iris_df[which(iris_df$Sepal.Length==5.1),] # returns observations with Sepal.Length = 5.1
```

I prefer to use which() because I like inputting numbers into the dataframe[,] arguments, rather than TRUE and FALSE values. Both should work, but ALWAYS ALWAYS ALWAYS check your output each time you run code to make sure R is doing exactly what you think it is doing. Likewise, if you have a nested function like I have above (i.e., which(iris_df\$Sepal.Length==5.1) nested in iris_df[,]), it is a good idea to check the output of that nested function before entering it into another function.

The tapply() function is useful for viewing information in one column by levels of another. This gives us the mean of Sepal.Width by Species. I use this all the time when I am writing papers and need to type out the means and standard errors of treatment groups.

```
tapply(iris_df$Sepal.Width, iris_df$Species, mean)

## setosa versicolor virginica
## 3.428 2.770 2.974
```

6 Cleaning/viewing data with tidyverse

Install the tidyverse package and load it into R

```
install.packages("tidyverse")
library(tidyverse)
```

We will now use the tidyverse functions and its pipe operators. First, we will use the filter() function to subset rows for sepal length > 5.

```
iris_df_subset_1 <- iris_df %>% filter(Sepal.Length > 5)
summary(iris_df_subset_1)
```

```
##
     Sepal.Length
                     Sepal.Width
                                      Petal.Length
                                                      Petal.Width
##
           :5.100
                           :2.200
                                            :1.200
                                                             :0.100
   Min.
                    Min.
                                     Min.
                                                     Min.
##
    1st Qu.:5.600
                    1st Qu.:2.800
                                     1st Qu.:3.925
                                                     1st Qu.:1.200
   Median :6.100
                    Median :3.000
                                     Median :4.700
                                                     Median :1.500
##
##
  Mean
           :6.130
                    Mean
                           :3.048
                                     Mean
                                            :4.315
                                                     Mean
                                                             :1.432
##
    3rd Qu.:6.575
                    3rd Qu.:3.300
                                     3rd Qu.:5.375
                                                     3rd Qu.:1.900
           :7.900
                           :4.400
                                     Max.
                                            :6.900
                                                             :2.500
##
   Max.
                    Max.
                                                     Max.
##
          Species
##
  setosa
              :22
##
   versicolor:47
##
   virginica:49
##
##
##
```

Let's also get the subset of observations with Sepal.Length values equal to 5.

```
iris_df_subset_2 <- filter(iris_df, Sepal.Length == 5)
summary(iris_df_subset_2)</pre>
```

```
##
     Sepal.Length Sepal.Width
                                   Petal.Length
                                                   Petal.Width
                                                                        Species
##
   Min.
           :5
                  Min.
                         :2.000
                                  Min.
                                          :1.20
                                                  Min.
                                                         :0.20
                                                                            :8
                                                                 setosa
                                   1st Qu.:1.40
   1st Qu.:5
                  1st Qu.:3.050
                                                  1st Qu.:0.20
                                                                 versicolor:2
## Median:5
                  Median :3.350
                                  Median:1.55
                                                  Median:0.25
                                                                 virginica:0
##
   Mean
           :5
                  Mean
                         :3.120
                                  Mean
                                          :1.84
                                                  Mean
                                                         :0.43
##
   3rd Qu.:5
                  3rd Qu.:3.475
                                   3rd Qu.:1.60
                                                  3rd Qu.:0.55
##
   Max.
           :5
                  Max.
                         :3.600
                                   Max.
                                          :3.50
                                                  Max.
                                                         :1.00
```

Let's get the mean, standard deviation, and max value of Sepal. Width by Species. Notice it is pretty straightforward to get multiple pieces of information for each level of a factor.

```
## # A tibble: 3 x 4
##
     Species
                Means
                         SD max_sep
##
     <fct>
                <dbl> <dbl>
                               <dbl>
## 1 setosa
                 3.43 0.379
                                 4.4
## 2 versicolor 2.77 0.314
                                 3.4
## 3 virginica
                 2.97 0.322
                                 3.8
```

Do the same thing as the previous example, but arrange the output by mean Sepal.Width.

```
iris_df %>%
group_by(Species) %>%
summarise(Means = mean(Sepal.Width),
SD = sd(Sepal.Width),
max_sep = max(Sepal.Width)) %>%
arrange(Means)
```

```
## # A tibble: 3 x 4
##
    Species
               Means
                         SD max_sep
##
     <fct>
                <dbl> <dbl>
                              <dbl>
## 1 versicolor 2.77 0.314
                                3.4
## 2 virginica 2.97 0.322
                                3.8
## 3 setosa
                 3.43 0.379
                                4.4
```

The tidyverse has its own command for converting variables into factors. If I have not convinced you yet, R has a LOT of ways of doing the same thing. Find the commands you need, make sure they are working as you intend (i.e, always check your output!), and try not to get bogged down in all the options.

```
iris_df$Species <- as_factor(iris_df$Species)</pre>
```

7 Dealing with NA values

Now let's add some NA values to the column Sepal.Length.

```
set.seed(123) # set location of random number generator
iris_df[sample(1:nrow(iris_df),10),"Sepal.Length"] <- NA</pre>
```

Here is a logical operator for NA values. The NA values of Sepal.Length are indicated by the values of TRUE. tail(is.na(iris_df\$Sepal.Length))

```
## [1] FALSE FALSE TRUE FALSE TRUE
```

Some functions in R do not know how to handle NA values.

```
mean(iris_df$"Sepal.Length") # oof

## [1] NA

mean(na.omit(iris_df$"Sepal.Length")) # yay

## [1] 5.855714

mean(iris_df$"Sepal.Length", na.rm=T) # also yay

## [1] 5.855714
```

Let's create a subset of data with only observations of Sepal.Length that are greater than the mean Sepal.Length. Notice that NA values can cause problems.

```
try1 <- iris_df %>% filter(Sepal.Length > mean(Sepal.Length))
summary(try1) # oof
```

Sepal.Length Sepal.Width Petal.Length Petal.Width Species

```
Min.
           : NA
                          : NA
                                         : NA
                                                        : NA
##
                   Min.
                                 Min.
                                                Min.
                                                               setosa
##
    1st Qu.: NA
                   1st Qu.: NA
                                  1st Qu.: NA
                                                1st Qu.: NA
                                                               versicolor:0
    Median: NA
                   Median: NA
                                 Median: NA
                                                Median : NA
                                                               virginica:0
##
    Mean
           :NaN
                   Mean
                          :NaN
                                 Mean
                                         :NaN
                                                Mean
                                                        :NaN
##
    3rd Qu.: NA
                   3rd Qu.: NA
                                  3rd Qu.: NA
                                                3rd Qu.: NA
           : NA
                                         : NA
##
    Max.
                   Max.
                          : NA
                                 Max.
                                                Max.
                                                        : NA
```

So, tell the mean() function how to handle NA values...

```
try2 <- iris_df %% filter(Sepal.Length > mean(Sepal.Length, na.rm = TRUE))
summary(try2) # yay!
##
     Sepal.Length
                      Sepal.Width
                                       Petal.Length
                                                        Petal.Width
##
    Min.
           :5.900
                     Min.
                            :2.200
                                      Min.
                                             :4.000
                                                       Min.
                                                              :1.000
##
    1st Qu.:6.225
                     1st Qu.:2.800
                                      1st Qu.:4.700
                                                       1st Qu.:1.500
##
    Median :6.450
                     Median :3.000
                                      Median :5.100
                                                       Median :1.800
##
    Mean
           :6.582
                            :2.956
                                             :5.229
                                                              :1.809
                     Mean
                                      Mean
                                                       Mean
##
    3rd Qu.:6.800
                     3rd Qu.:3.175
                                      3rd Qu.:5.700
                                                       3rd Qu.:2.100
##
           :7.900
                            :3.800
                                                               :2.500
    Max.
                     Max.
                                      Max.
                                             :6.900
                                                       Max.
##
          Species
##
              : 0
    setosa
##
    versicolor:25
##
    virginica:41
##
##
##
```

8 Proof of concept on the random number generator.

The random number generator in R can be very helpful for assigning treatments, running simulations, and calling on students during class. Run the following code all at once. Then just run the sample() function a few times. Then run the all the code all at once again. You should notice that the output of sample() is exactly the same when you "set the seed."

```
set.seed(123) # set random number generator to same place
sample(1:100,3)
```

[1] 31 79 51

9 R Activity

- 1. Load in the breakfast data set. Remember to check the file type so that you can identify the right command for loading in the data.
- 2. Use a single command to count the number of columns in the breakfast data.
- 3. Use base R (i.e., no external packages should be used for this question) to create a subset of the data that has only observations with values of Calories greater than 130.
- 4. Use the pipe operators in tidyverse to create a subset of the data that has only observations with values of Calories greater than 130.
- 5. Use the pipe operators in tidyverse to calculate the mean Sugars by Company.