Group Project 1

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R for Dummies (BIOL208 Edition)

Welcome to this R tutorial for BIOL208! This is designed for someone who is brand new to R, so we are going to start with the very basics. To make this interesting, we'll be looking at a dataset on irises since plants are extremely important to the study of ecology and evolution. To best follow along, you should have a new markdown file open in R studio where you can copy and paste line sof code from this tutorial for you to practice running on your own.

Typically, when doing data analysis in R, we want to begin by loading in the data and packages that we will use.

Loading Libraries and Data

R has many built in functions that will allow us to perform all kinds of actions. However, sometimes it is useful to load some additional functions that other R users have created. We do this by loading in libraries. In the code chunk below (the grey box; this is where code is typed and run) you can see that I have loaded in some libraries.

```
# Install packages and load their libraries here.
if (!require("tidyverse")) install.packages("tidyverse"); library(tidyverse)
if (!require("cowplot")) install.packages("cowplot"); library(cowplot)
if (!require("UsingR")) install.packages("UsingR"); library(UsingR)
if (!require("ggpubr")) install.packages("ggpubr"); library(ggpubr)
if (!require("readxl")) install.packages("readxl"); library(readxl)
```

Libraries

For some examples, one library is called tidyverse. It contains a ton of useful tools, but the one most relevant to this tutorial is called **ggplot**. **ggplot** allows us to create all kinds of plots to visualize our data, which you will learn more about below. You can also see the library called **cowplot**. **cowplot** is useful as it allows us to easily change the aesthetics of our graphs, and keep the designs consistent between different graphs. Finally, you can see the library **readxl**. This is useful for exactly what it sounds like... reading Microsoft Excel files! This bring us to the next step, entering data into R.

Data

In the code chunk below I am going to read in some data regarding features of different species of iris (the flower). There are a few main parts of the code to think about when loading in data. First, the **read_csv()** function tells R to load in a comma separated values (csv) file with the name of the file in the parentheses. Here, you can see that our data is called "iris.csv". It is very important to type the file name exactly as it is saved on your computer, otherwise R will not be able to find it. Once the data is loaded in, we need to call it something. That is what the "IrisData <-" part of the code means. In English, it translates to:

IrisData is the data found in the csv file "iris.csv". When you read in a data file, you will see it appear in your "Environment" on the right of R studio. If you click on "IrisData" in your Environment, a new tab will appear with all of the data.

```
IrisData <- read_csv ("iris.csv")

## Rows: 150 Columns: 5

## -- Column specification --------
## Delimiter: ","

## chr (1): variety

## dbl (4): sepal.length, sepal.width, petal.length, petal.width

##

## i Use `spec()` to retrieve the full column specification for this data.

## i Specify the column types or set `show_col_types = FALSE` to quiet this message.</pre>
```

What happens if your data is in an excel file instead of a csv file? Fear not! Data can also be loaded in from an excel file using the **read_excel()** function instead of **read_csv()**. In a normal data analysis, you would only read in the data once using *either* the **read_csv()** or **read_excel()** functions. Your data will only read in properly if the file type matches the function you're using, so pay attention to how your data is saved before you read it in!

Using R (The Basics)

Now that we have the pieces we need, we can start learning about R. You have already seen a few aspects of R in the last section, but here we will go into some more detail about the basic functions of R to get you started with analyzing data.

This document is an R markdown file. The highlights of a markdown file is that you can have mostly regular text (not code) but still include code and their output in specific sections. This is especially useful as it allows for lots of explanation of what the code does, like we need to do here for a tutorial. Markdowns are also convenient because they can be converted to many different formats such as a PDF (like you're reading), an html, and others.

As mentioned before, the grey sections are called code chunks. This is where the actual code is written and ran. You can insert a code chunk to your markdown file by clicking the green box with "+C" at the top of R studio. When you do this, a drop-down menu appears where you can choose a coding language (since this is about R, we will only use that). Refer to Figure 1 if you can't find the code chunk button.

In the top line of your new code chunk, you'll see "'{r}. This can be used to name your code chunks if you wish. This can help keep your thoughts organized and is actually an important part of knitting your file... Say what??? We're going to be knitting? Don't worry, we'll explain *all* the details about knitting in the later sections of this tutorial. For now just know that naming your code chunks can help organize your thoughts and can save you problems with knitting later down the road if you make sure to name your each of your code chunks something unique.

```
#Example code chunk
4+1

## [1] 5

#This line will not run
```

Code within chunks can be run in a few ways:

- 1. In the top right of a chunk is a green play button that will run every line of code in the chunk.
- 2. Placing your cursor on a line you want to run, or highlight it, and hit command + enter (or control + enter).

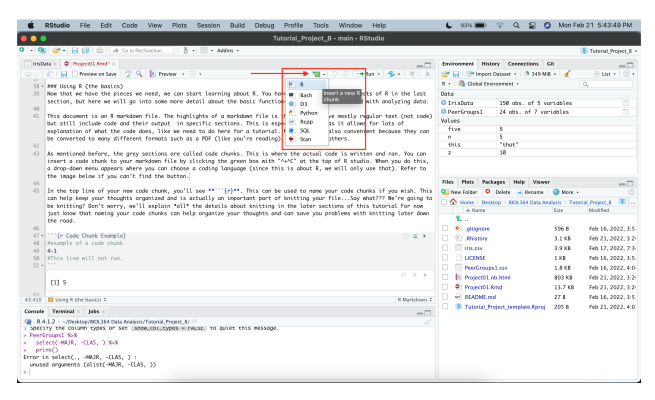


Figure 1: Adding a New Code Chunk

3. Click the run button in the top of the screen, which will bring up a few options for running code. You can run all of the code chunks in your document at once (in order from top to bottom) with this option. This is useful for ensuring your code is functional when finalizing a document. Figure 2 outlines where you can find each of these.

Looking at the example code chunk above, there a few points to make about coding in R. First, you can see a green line that starts with a "#". The hashtag/number/pound symbol tells R that this line is a comment, and that it should not be run as code. If you run the code chunk you will see that that line is skipped, and the only output comes from the addition found in the next line (the output should be 5 since 4+1=5). Comments are very useful for explaining your code as you write it. This is helpful for yourself in the future, so you can remind yourself how different code works, and for other people reading through or referencing your code.

Another useful bit of information is how to assign values to objects. Essentially, this is a way of naming things. You have actually already seen this in action, where we assigned our iris data to the object IrisData using the assign operator <-. Lets go to the next code chunk to see an example and notice how all of these values/variables will get stored in the Environment.

```
#Essentially, the <- means "store this value under this name," so you can think of the symbol as the wo this <- "that"

#In the line above we are storing the word that under the name this. So the code would read in English this

## [1] "that"

#Other examples 
n <-5 
z <-10
```

n+z

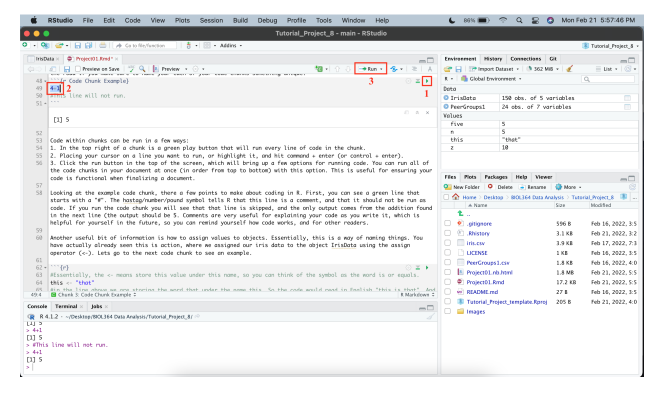


Figure 2: Running Code

```
## [1] 15
z/n
## [1] 2
(z+n)/3 -> five
five * 10
```

This is especially useful for storing data under a specific name, like we did at the beginning with the iris data. If you run the line of code below, it will print out all of the iris data.

IrisData

[1] 50

```
## # A tibble: 150 x 5
##
      sepal.length sepal.width petal.length petal.width variety
##
              dbl>
                           <dbl>
                                         <dbl>
                                                      <dbl> <chr>
##
                5.1
                             3.5
                                           1.4
                                                        0.2 Setosa
    1
##
    2
                4.9
                             3
                                                        0.2 Setosa
                                           1.4
    3
                4.7
                                                        0.2 Setosa
##
                             3.2
                                           1.3
##
    4
                4.6
                             3.1
                                           1.5
                                                        0.2 Setosa
                                           1.4
##
    5
                5
                             3.6
                                                        0.2 Setosa
##
    6
                5.4
                             3.9
                                           1.7
                                                        0.4 Setosa
    7
##
                4.6
                             3.4
                                           1.4
                                                        0.3 Setosa
##
    8
                5
                             3.4
                                           1.5
                                                        0.2 Setosa
    9
##
                4.4
                             2.9
                                           1.4
                                                        0.2 Setosa
## 10
                4.9
                                           1.5
                                                        0.1 Setosa
                             3.1
## # ... with 140 more rows
```

We can also access specific variables within the data using the \$ symbol. So if we only wanted to see the data about petal length, we can run the following code:

IrisData\$petal.length

```
## [1] 1.4 1.4 1.3 1.5 1.4 1.7 1.4 1.5 1.4 1.5 1.6 1.4 1.1 1.2 1.5 1.3 1.4 ## [19] 1.7 1.5 1.7 1.5 1.0 1.7 1.9 1.6 1.6 1.5 1.4 1.6 1.6 1.5 1.5 1.4 1.5 1.2 ## [37] 1.3 1.4 1.3 1.5 1.3 1.3 1.3 1.6 1.9 1.4 1.6 1.4 1.5 1.4 4.7 4.5 4.9 4.0 ## [55] 4.6 4.5 4.7 3.3 4.6 3.9 3.5 4.2 4.0 4.7 3.6 4.4 4.5 4.1 4.5 3.9 4.8 4.0 ## [73] 4.9 4.7 4.3 4.4 4.8 5.0 4.5 3.5 3.8 3.7 3.9 5.1 4.5 4.5 4.7 4.4 4.1 4.0 ## [91] 4.4 4.6 4.0 3.3 4.2 4.2 4.2 4.3 3.0 4.1 6.0 5.1 5.9 5.6 5.8 6.6 4.5 6.3 ## [109] 5.8 6.1 5.1 5.3 5.5 5.0 5.1 5.3 5.5 6.7 6.9 5.0 5.7 4.9 6.7 4.9 5.7 6.0 ## [127] 4.8 4.9 5.6 5.8 6.1 6.4 5.6 5.1 5.6 6.1 5.6 5.5 4.8 5.4 5.6 5.1 5.1 5.9 ## [145] 5.7 5.2 5.0 5.2 5.4 5.1
```

Those are the basic functions you should know about R moving forward. Now we can explore more tools you can use to explore and analyze a set of data.

Exploring Your Data

Once we have our data, it is important to explore it before beginning analysis. First, you want to come up with the question that you will attempt to answer before analyzing your data. This is important since we want to remain objective throughout the data analysis process.

For this tutorial, we're interested in the following questions:

- 1. Are sepal and petal dimensions correlated within a single species?
- 2. Do sepal and petal dimensions differ between Iris species?

One way to begin to understand your dataset is to run the **str()** (structure) function. This outputs lots of information about the dataset. It will show you each column of the data, the class of the column (what type of data it is: a number (num), a word (or character: chr), etc), the number of observations in the column, and some of the data within each column. This is useful for ensuring that each column contains the data you expect. For example, one column in the dataset is sepal length, so I would expect to find in the output of str() that there is a column called sepal length (it is actually called sepal.length, but that is good), it should be a number, there should be 150 observations, and the data should actually be numbers.

In the chunk below I will run str() on the iris dataset to confirm my expectations and make sure that the data looks right.

str(IrisData)

```
## spec_tbl_df [150 x 5] (S3: spec_tbl_df/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 0.4 0.3 0.2 0.2 0.1 ...
                  : chr [1:150] "Setosa" "Setosa" "Setosa" "Setosa" ...
##
   $ varietv
   - attr(*, "spec")=
##
##
     .. cols(
          sepal.length = col_double(),
##
##
          sepal.width = col double(),
##
          petal.length = col double(),
##
         petal.width = col double(),
##
          variety = col character()
##
   - attr(*, "problems")=<externalptr>
```

One thing that stands out to me after running **str()** is the "variety" column. For starters, you may want to rename the column to something that's easier for you to understand, such as "species." You can do this using the **rename()** function as explained below. Once you've run that code, you can click on the dataset in your Environment and should see the column has been renamed.

If we want to compare variables between species of irises, this column will be very useful. To make those comparisons easier, I am going to tell R that this variable is categorical. We can group the data into those categories using the as.factor() function below.

```
IrisData$species <- as.factor(IrisData$species)</pre>
#This code tells R to replace the current character species column of IrisData with the new factor spec
str(IrisData)
## spec_tbl_df [150 x 5] (S3: spec_tbl_df/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 ...
##
                  : Factor w/ 3 levels "Setosa", "Versicolor", ...: 1 1 1 1 1 1 1 1 1 1 1 ...
##
   - attr(*, "spec")=
##
##
     .. cols(
##
          sepal.length = col_double(),
##
          sepal.width = col_double(),
          petal.length = col_double(),
##
          petal.width = col_double(),
##
##
          variety = col_character()
##
     ..)
   - attr(*, "problems")=<externalptr>
```

 $\#Running\ str()\ again\ shows\ that\ species\ is\ now\ a\ factor\ with\ 3\ levels\ which\ correspond\ to\ the\ 3\ different factor$

It can also be useful to look at the top and bottom of your dataset using the **head()** and **tail()** functions, which show the first 6 rows and last 6 rows respectively. This is useful to ensure that the data makes sense (the data is numeric if you were expecting that) and to make sure that the correct number of rows are present (for example, make sure that the last few rows are not empty). Or, if the data is time dependent, you can use these functions to make sure that the first times are in the first row and the last times are at the end.

#This chunk will output two functions, and it will separate them into two windows if you run the whole head(IrisData)

```
## # A tibble: 6 x 5
     sepal.length sepal.width petal.length petal.width species
                                                     <dbl> <fct>
##
             <dbl>
                          <dbl>
                                        <dbl>
               5.1
                            3.5
                                          1.4
                                                       0.2 Setosa
## 1
## 2
               4.9
                            3
                                          1.4
                                                       0.2 Setosa
## 3
               4.7
                            3.2
                                          1.3
                                                       0.2 Setosa
                                                       0.2 Setosa
## 4
               4.6
                            3.1
                                          1.5
## 5
               5
                            3.6
                                          1.4
                                                       0.2 Setosa
## 6
               5.4
                            3.9
                                                       0.4 Setosa
                                          1.7
tail(IrisData)
```

```
## # A tibble: 6 x 5
## sepal.length sepal.width petal.length petal.width species
```

##		<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<fct></fct>
##	1	6.7	3.3	5.7	2.5	Virginica
##	2	6.7	3	5.2	2.3	Virginica
##	3	6.3	2.5	5	1.9	Virginica
##	4	6.5	3	5.2	2	Virginica
##	5	6.2	3.4	5.4	2.3	Virginica
##	6	5.9	3	5.1	1.8	Virginica

A very useful function for data exploration is **summary()**. This outputs the mean, median, min, max, and first and third quartiles of any numeric data, and the number of observations within our variables. This is an important step in any data exploration. This can be useful for validating your data against external sources. If external sources show that the average sepal length of irises is 100 cm long, but our sepal lengths range from 4.3 to 7.9 cm, something might be wrong. Luckily the average is closer to 5-7 cm, so our data does make sense.

summary(IrisData)

```
##
                                                         petal.width
     sepal.length
                       sepal.width
                                        petal.length
##
    Min.
            :4.300
                     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
##
            :5.843
                             :3.057
                                               :3.758
                                                                :1.199
    Mean
                     Mean
                                       Mean
                                                        Mean
##
    3rd Qu.:6.400
                     3rd Qu.:3.300
                                       3rd Qu.:5.100
                                                        3rd Qu.:1.800
##
    Max.
            :7.900
                     Max.
                             :4.400
                                       Max.
                                               :6.900
                                                        Max.
                                                                :2.500
##
          species
##
    Setosa
               :50
    Versicolor:50
##
##
    Virginica:50
##
##
##
```

Visualization with Graphs

Now that we've spent a substantial amount of time exploring and organizing our data, we're ready to start analyzing it! The first step here is to visualize our data since up until this point, all we've been looking at are numbers (which can be hard for identifying relationships and patterns between variables). When considering what graphs would be best for visualizing our data, we should consider our questions of interest and what types of variables we're dealing with. As a reminder, our two questions of interest are below:

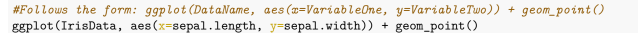
- 1. Are sepal and petal dimensions correlated within a single species?
- 2. Do sepal and petal dimensions differ between Iris species?

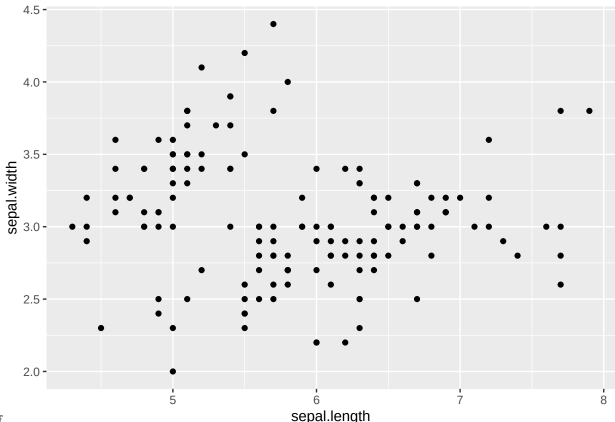
Are sepal and petal dimensions correlated within a single species?

Let's focus on question number one first. There are a couple different, more specific questions we can look at within this. Such as, the relationship between sepal length and sepal width, or petal length and petal width. Perhaps we can even compare sepal length and petal length, or sepal width and petal width. Wow! There are a ton of different relationships to look at. For our purposes, let's narrow in on two relationships:

- 1. Sepal length and sepal width
- 2. Sepal length and petal length

Since all of these variables are numerical, a scatterplot would best represent the data. The **ggplot** function is extremely useful for creating graphs. Below is the code for creating a basic scatterplot.



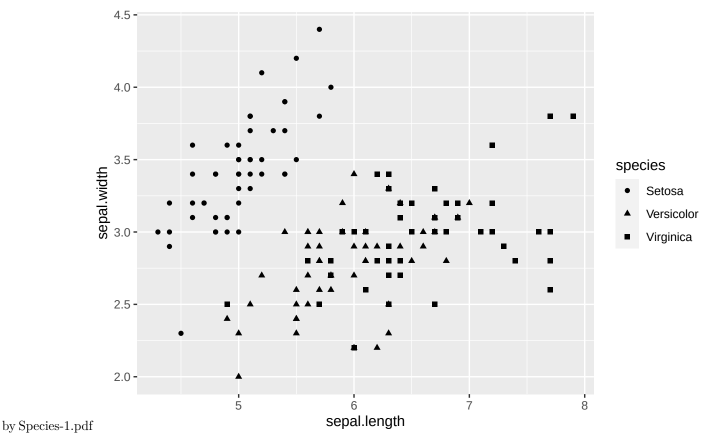


Scatterplot-1.pdf

#Learning check: Try running the code without + geom_point(). What happens?

In the output we see a basic scatterplot without any clear pattern. This could be because we're looking at data for all three species. Remember, our original question was about whether there were relationships between the variables within a *single* species. We can either filter the data so that we can plot one species at a time, or we can just color-code the points on the scatterplot by species so we can still look at all of the data (this is the method we'll use).

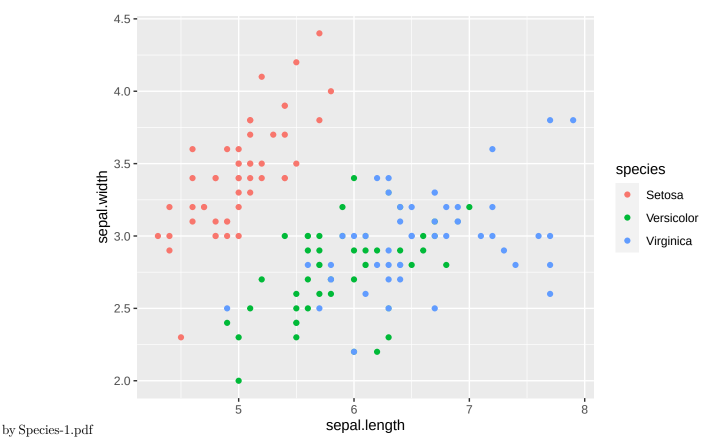
ggplot(IrisData, aes(x=sepal.length, y=sepal.width, shape=species)) + geom_point()



 $\#Adding\ shape=species\ in\ the\ code\ line\ tells\ R$ to group the points according to species and to differen

By separating the data points according to species we can start to see some patterns emerging between the variables within each species. However, it still takes a second to differentiate between Versicolor and Virginica since they occupy the same general space on the plot and you don't immediately notice the difference in shapes. To make it easier to distinguish between the two, we can use color to differentiate the species rather than shape.

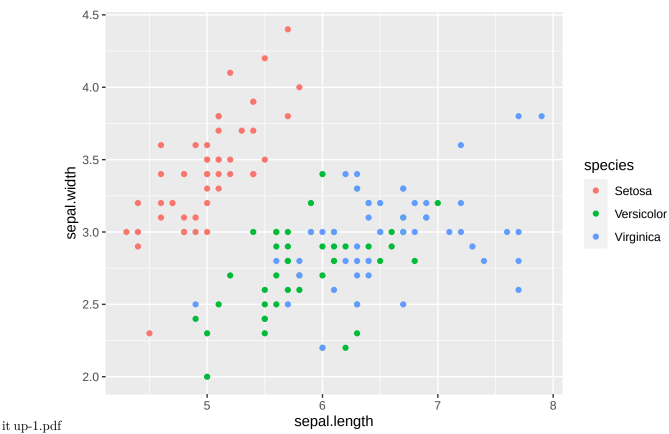
ggplot(IrisData, aes(x=sepal.length, y=sepal.width, color=species)) + geom_point()



 $\#Adding\ color=species\ in\ the\ code\ line\ tells\ R$ to differentiate the species by color as well as by shap

Based on this scatterplot, there seems to be a positive correlation between sepal length and sepal width within each species. Say you wanted to present this data in a formal report. You would likely want to clean it up a bit. Fortunately, **ggplot** has a *ton* of ways you can spruce up your graphs to look neater and more professional. Let's take a look at some of those tools.

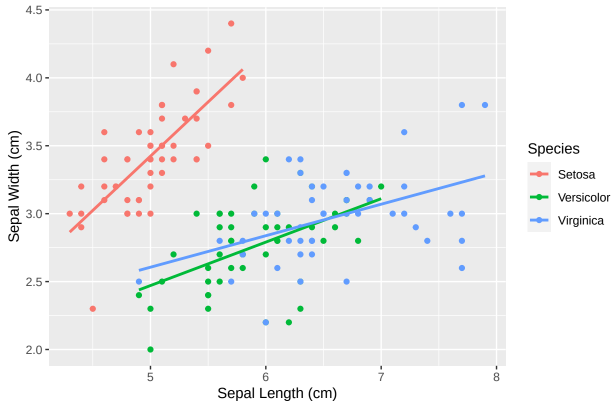
```
ggplot(IrisData, aes(x=sepal.length, y=sepal.width, color=species)) +
  geom_point()
```



#This code runs the same as the code we used previously, even though geom_point() is on a separate line
ggplot(IrisData, aes(x=sepal.length, y=sepal.width, color=species)) + #Basic scatterplot grouped by col
geom_point() + #Adds data points to plot
geom_smooth(method=lm, se=FALSE) + #Adds regression lines
labs(x = "Sepal Length (cm)", y = "Sepal Width (cm)") + #Renames axis labels
labs(col="Species") + #Renames legend
labs(title = "Sepal Width vs. Sepal Length in Three Iris Species") + #Gives the plot a title
theme(plot.title = element_text(hjust = 0.5)) #Centers the plot title

`geom_smooth()` using formula 'y ~ x'





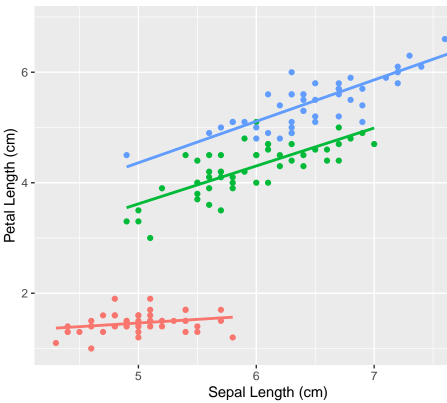
it up-2.pdf

This plot looks much nicer and is easier to interpret compared to our initial scatterplot. Now that we have the code all figured out, we can essentially copy and paste the same code but substitute in new variable to look at other relationships as well. Let's look at the relationship between

```
ggplot(IrisData, aes(x=sepal.length, y=petal.length, color=species)) +
  geom_point() +
  geom_smooth(method=lm, se=FALSE) +
  labs(x = "Sepal Length (cm)", y = "Petal Length (cm)") +
  labs(col="Species") +
  labs(title = "Petal Length vs. Sepal Length in Three Iris Species") +
  theme(plot.title = element_text(hjust = 0.5))
```

`geom_smooth()` using formula 'y ~ x'

Petal Length vs. Sepal Length in Three Iris Speci



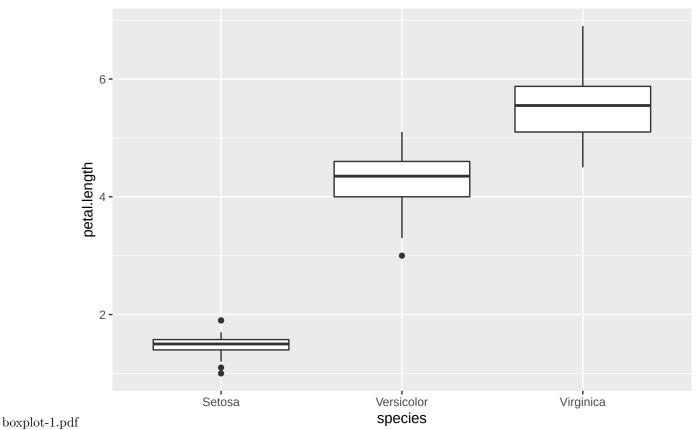
comparing Petal Length and Sepal Length-1.pdf

This scatterplot reveals that there also seems to be a positive correlation between sepal length and petal length within each species, though the strength of that relationship may differ between species. Feel free to use this scatterplot code to visualize relationships between other variables as well.

Do sepal and petal dimensions differ between Iris species?

Our next question of interest deals with comparing dimensions *between* species. Once again, there are a ton of different, more specific questions we can look at within this. For our purposes, let's focus on sepal length between species. Since species is a categorical variable, a boxplot would best represent the data. The code for a basic boxplot is below.

ggplot(IrisData, aes(x=species, y=petal.length)) + geom_boxplot()

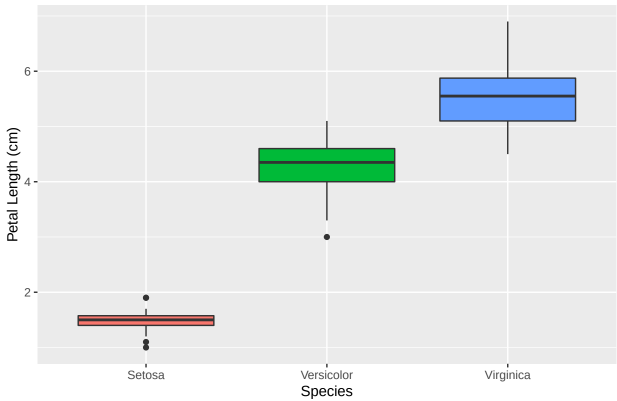


#Replacing geom_point() from our basic scatterplot code with geom_boxplot() tells R to make a boxplot.

There appears to be a difference in sepal length between species (note: determining whether or not that difference is significant is for later statistical testing). Just as we did before, let's spruce up our plot a bit.

```
ggplot(IrisData, aes(x=species, y=petal.length, fill=species)) + #Addition of fill=species adds color
geom_boxplot() + #Specifies boxplot
labs(x = "Species", y = "Petal Length (cm)") + #Renames axes
labs(title = "Petal Length for Three Iris Species") + #Gives plot a title
theme(plot.title = element_text(hjust = 0.5)) + #Centers title
theme(legend.position = "none") #Removes legend
```

Petal Length for Three Iris Species

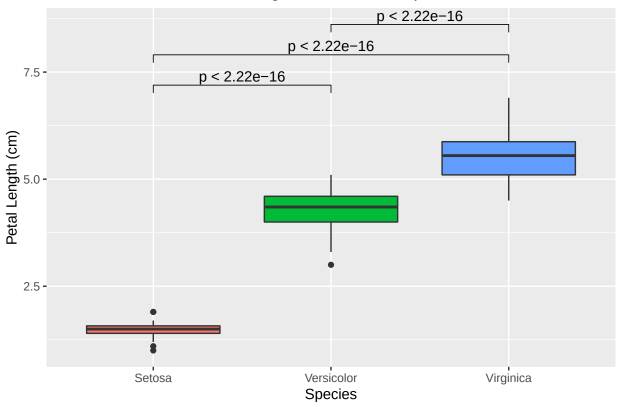


Boxplot-1.pdf

Once you get *really* good with R, you'll be able to make plots like the one below, where you include the significance bars for pairwise comparisons between each of the species. Even though we're getting ahead of ourselves here, feel free to play around with the code below to make plots for other variables and look at the p-values for pairwise comparisons between species.

```
compare_means(petal.length ~ species, data = IrisData)
## # A tibble: 3 x 8
##
                  group1
                             group2
                                                   p.adj p.format p.signif method
     .y.
##
     <chr>>
                  <chr>
                             <chr>
                                                   <dbl> <chr>
                                                                  <chr>
                                                                            <chr>
                                           <dbl>
                             Versicolor 5.65e-18 1.7e-17 <2e-16
## 1 petal.length Setosa
                                                                            Wilcoxon
## 2 petal.length Setosa
                             Virginica 5.67e-18 1.7e-17 <2e-16
                                                                  ****
                                                                            Wilcoxon
                                                                            Wilcoxon
## 3 petal.length Versicolor Virginica 9.13e-17 9.1e-17 <2e-16
my_comparisons <- list(c("Setosa", "Versicolor"), c("Setosa", "Virginica"), c("Versicolor", "Virginica"
ggplot(IrisData, aes(x=species, y=petal.length, fill=species)) + #Addition of fill=species adds color
  geom_boxplot() +
  labs(x = "Species", y = "Petal Length (cm)") + #Renames axes
  labs(title = "Petal Length for Three Iris Species") + #Gives plot a title
  theme(plot.title = element_text(hjust = 0.5)) + #Centers title
  theme(legend.position = "none") + #Removes legend
  stat_compare_means(comparisons = my_comparisons) #Adds p-values for pairwise comparisons
```

Petal Length for Three Iris Species



Hypothesis Testing with Statistical Tests

Up until this point we've been looking at the data with some questions in mind about relationships in the data. For statistical tests, we need a specific question or hypothesis to test. Our hypothesis is that petal length differs significantly between Iris species. We will test this hypothesis below.

Requirements for Tests

Now we can move on to analyzing our data more critically with statistical testing for specific hypotheses. When using statistical tests, always be sure to check the assumptions made by the particular test you plan to use. This is important because many common statistical tests, including the two-sample t-test we will use here, assume that the datasets being compared are normally distributed. While much biological data is normally distributed, this is not always true, so be sure to explore and understand the behavior of your data before performing tests.

The **shapiro.test** function allows us to check if our data is normally distributed. If the data is normally distributed, the p-value returned by the shapiro test should be greater than 0.05. If there are multiple groups/populations within a data set (for example, multiple species of iris), it's important to test for a normal distribution within each of those populations separately. Looking at the distribution as a whole may lead us to think the data is not normally distributed when it actually is within each population. This is actually the case for us, as you can see with the output below.

shapiro.test(IrisData\$petal.length) #Not normally distributed as a whole (p = 7.412e-10 < 0.05)

```
##
## Shapiro-Wilk normality test
##
## data: IrisData$petal.length
```

```
## W = 0.87627, p-value = 7.412e-10
shapiro.test(IrisData$petal.length[IrisData$species=="Setosa"]) #Normally distributed (p = 0.05481 > 0.
##
   Shapiro-Wilk normality test
##
##
## data: IrisData$petal.length[IrisData$species == "Setosa"]
## W = 0.95498, p-value = 0.05481
shapiro.test(IrisData$petal.length[IrisData$species=="Versicolor"]) #Normally distributed (p = 0.1585 >
##
##
   Shapiro-Wilk normality test
##
## data: IrisData$petal.length[IrisData$species == "Versicolor"]
## W = 0.966, p-value = 0.1585
shapiro.test(IrisData$petal.length[IrisData$species=="Virginica"]) #Normally distributed (p = 0.1098 >
##
##
   Shapiro-Wilk normality test
##
## data: IrisData$petal.length[IrisData$species == "Virginica"]
## W = 0.96219, p-value = 0.1098
```

If the data you are using is not normally distributed, test to see if a logarithmic transformation would make the distribution normal. If so, you should use the log-transformed data. If not, continue on with the original, non-transformed data but note in your report that any statistical test results should be interpreted with caution, given that the base requirements for accurate use of the test have not been met.

Since the data for petal length is normally distributed within each species, a log transformation is not necessary in our case. We will proceed with the non-transformed data. However, here is an example of how you would perform a log transformation if you needed to:

```
#log transform the data and assign it to a variable
logpetal.length <- log(IrisData$petal.length)
#test for normality of this newly transformed data
shapiro.test(logpetal.length)

##
## Shapiro-Wilk normality test
##
## data: logpetal.length
## W = 0.81692, p-value = 2.038e-12</pre>
```

Note that each statistical test which assumes normality also has a sample size requirement which can allow for them to produce accurate results using non-normal data. For a two-sample t-test or a one-way ANOVA test, each group is required to have a sample size of at least 15 observations to overcome the need for normality. If your samples do fit this requirement, you can proceed without worrying about normality. This cutoff is different for each type of test, so always be sure to check before you disregard any requirements.

ANOVA and Two-sample T-tests In order to analyze whether petal length differs significantly between species, we can perform an ANOVA. We do this with the aov() function. It's best to assign the function to a variable that you can then run a summary of to see the output from the ANOVA test.

```
anova <- aov(petal.length ~ species, data = IrisData)
summary(anova)</pre>
```

```
## Df Sum Sq Mean Sq F value Pr(>F)
## species 2 437.1 218.55 1180 <2e-16 ***
## Residuals 147 27.2 0.19
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

The ANOVA test returned a p-value < 2e-16 which is much less than 0.05 (the standard significance level). If you recall from statistics, this gives us evidence that petal length is significantly different for at least one of the groups (species), but we don't know which. In order to further analyze petal length between species, we can conduct pairwise t-tests to see which species' petal lengths significantly differ from one another. In order to conduct pairwise comparisons, we need to filter the data into three data subsets, one for each pairwise comparison. We can do this using the **filter()** function and assign each filtered dataset to a new dataset.

```
#filter() uses the form: filter(DataName, VariableName=Value). This indicates that we want to pull all
SetosaVersicolor <- filter(IrisData, species!="Virginica")</pre>
SetosaVirginica <- filter(IrisData, species!="Versicolor")</pre>
VersicolorVirginica <- filter(IrisData, species!="Setosa")</pre>
t.test(petal.length ~ species, data = SetosaVersicolor) #Significant (p < 2.2e-16)
##
##
    Welch Two Sample t-test
## data: petal.length by species
## t = -39.493, df = 62.14, p-value < 2.2e-16
## alternative hypothesis: true difference in means between group Setosa and group Versicolor is not eq
## 95 percent confidence interval:
##
   -2.939618 -2.656382
## sample estimates:
##
       mean in group Setosa mean in group Versicolor
##
                      1.462
                                                4.260
t.test(petal.length ~ species, data = SetosaVirginica) #Significant (p <2.2e-16)
##
##
   Welch Two Sample t-test
##
## data: petal.length by species
## t = -49.986, df = 58.609, p-value < 2.2e-16
## alternative hypothesis: true difference in means between group Setosa and group Virginica is not equ
## 95 percent confidence interval:
  -4.253749 -3.926251
## sample estimates:
##
      mean in group Setosa mean in group Virginica
                     1.462
                                              5.552
t.test(petal.length ~ species, \frac{data}{data} = VersicolorVirginica) #Significant (p < 2.2e-16)
##
##
   Welch Two Sample t-test
## data: petal.length by species
## t = -12.604, df = 95.57, p-value < 2.2e-16
## alternative hypothesis: true difference in means between group Versicolor and group Virginica is not
## 95 percent confidence interval:
## -1.49549 -1.08851
```

sample estimates:

```
## mean in group Versicolor mean in group Virginica
## 4.260 5.552
```

The results from the pairwise comparisons indicate that the mean petal length for the three species are each significantly different from one another. We actually see the same p-values returned by the pairwise t-tests displayed on the box-plot with p-values. This makes sense, since the p-values on the boxplot resulted from pairwise comparisons. Overall, we have evidence to support the hypothesis that petal length differs significantly between Iris species.

Linear Model

We can also use a linear model as a statistical test to look at relationships in the data. We already used a linear model to some degree when we created our scatterplot and included a regression line. However, now we're going to look at a linear model more quantitatively rather than qualitatively. The lm() function allows us to build a linear model with our data.

```
#The lm() function takes the form: lm(DependentVariable ~ IndependentVariable, data=DataName)
lmSpecies <- lm(petal.length ~ species, data=IrisData)
summary(lmSpecies)
```

```
##
## Call:
## lm(formula = petal.length ~ species, data = IrisData)
## Residuals:
##
              1Q Median
     Min
                            3Q
                                  Max
## -1.260 -0.258 0.038 0.240
                                1.348
##
## Coefficients:
                     Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                      1.46200
                                 0.06086
                                           24.02
                                                   <2e-16 ***
                      2.79800
                                 0.08607
                                           32.51
## speciesVersicolor
                                                   <2e-16 ***
## speciesVirginica
                      4.09000
                                 0.08607
                                           47.52
                                                   <2e-16 ***
##
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.4303 on 147 degrees of freedom
## Multiple R-squared: 0.9414, Adjusted R-squared: 0.9406
## F-statistic: 1180 on 2 and 147 DF, p-value: < 2.2e-16
```

The p-value returned with the linear model is < 2.2e-16. This shows us that petal length is dependent on/explained by species. This is what we expected based on earlier statistical tests. For the sake of this tutorial, let's explore another linear model that looks at petal length sepal width.

```
lmSepalWidth <- lm(petal.length ~ sepal.width, data=IrisData)
summary(lmSepalWidth)</pre>
```

```
##
## Call:
## lm(formula = petal.length ~ sepal.width, data = IrisData)
##
## Residuals:
##
                1Q Median
                                 30
       Min
                                        Max
## -3.7721 -1.4164 0.1719 1.2094
                                    4.2307
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
```

```
## (Intercept) 9.0632 0.9289 9.757 < 2e-16 ***
## sepal.width -1.7352 0.3008 -5.768 4.51e-08 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 1.6 on 148 degrees of freedom
## Multiple R-squared: 0.1836, Adjusted R-squared: 0.178
## F-statistic: 33.28 on 1 and 148 DF, p-value: 4.513e-08</pre>
```

This linear model returned a p-value of 4.513e-8. While this is still a very significant p-value, we see that our model which used species rather than sepal width was a better model. We can represent the difference in how well these two modes fit the data with an ANOVA that compares the two models.

```
anova(lmSepalWidth, lmSpecies, test='F')
```

```
## Analysis of Variance Table
##
## Model 1: petal.length ~ sepal.width
## Model 2: petal.length ~ species
                                         Pr(>F)
##
     Res.Df
              RSS Df Sum of Sq
## 1
        148 379.09
## 2
           27.22
                        351.87 1900.1 < 2.2e-16 ***
        147
                   1
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

The ANOVA returned a p-value < 2.2e-16. This is interpreted as having sufficient evidence that the second model in the code line (species model) is a better fit for the data than the first model in the code line (sepal width model). So while both models may be a good fit for the data, the model with species is a better fit overall. We can also consider whether a linear model that combines *both* variables would be an even better fit than a model that only considers species.

```
lmSpeciesSepalWidth <- lm(petal.length ~ species + sepal.width, data=IrisData)
summary(lmSpeciesSepalWidth)</pre>
```

```
##
## Call:
## lm(formula = petal.length ~ species + sepal.width, data = IrisData)
##
## Residuals:
##
        Min
                  1Q
                       Median
                                     30
                                             Max
##
   -1.13073 -0.22613 -0.01516
                               0.23307
##
## Coefficients:
##
                     Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                     -0.17920
                                 0.33753
                                           -0.531
                                                     0.596
                                           30.420
                                                   < 2e-16 ***
## speciesVersicolor
                      3.11303
                                  0.10234
## speciesVirginica
                      4.30736
                                  0.09130
                                           47.179 < 2e-16 ***
                                            4.932 2.19e-06 ***
## sepal.width
                      0.47876
                                 0.09707
##
                  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
## Residual standard error: 0.3998 on 146 degrees of freedom
## Multiple R-squared: 0.9497, Adjusted R-squared: 0.9487
## F-statistic: 919.7 on 3 and 146 DF, p-value: < 2.2e-16
anova(lmSpecies, lmSpeciesSepalWidth, test='F')
```

```
## Analysis of Variance Table
##
## Model 1: petal.length ~ species
## Model 2: petal.length ~ species + sepal.width
## Res.Df RSS Df Sum of Sq F Pr(>F)
## 1 147 27.223
## 2 146 23.335 1 3.8879 24.326 2.186e-06 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

The ANOVA returned a p-value of 2.186e-6. This is interpreted as having sufficient evidence that the combined model is a better fit than the model that only considers species. This makes sense since our earlier analyses revealed that all of the variables in this dataset seemed to be closely related with one another.

Congratulations! You just got through the **R** for **Dummies** (**BIOL208 Edition**) tutorial, and that's no easy task. The world of coding in R still has so much for you to discover, but you have the skills now to get started with it. If this tutorial has left you craving for more data analysis and R knowledge, check out the additional tools below. Happy coding!

Additional Tools

Explanation of p-values

A p-value represents the probability that the null hypothesis is true given the outcome observed. For most purposes, a p-value of 0.05 or lower is considered statistically significant. However, staunchly relying on this value comes with some concerns because, although it is commonly used and accepted, it is essentially an arbitrary cutoff. Thus, be sure to understand what this cutoff means so you can accurately express the strength of each result.

There are two main ways in which you can unwittingly abuse p-values. The first is known as p-hacking and occurs when researchers end up with results which are not statistically significant but then continue to find new ways to manipulate the data until they eventually find something significant. The second is through what Stuart Ritchie calls "HARKing" or "Hypothesising After the Results are Known," in which researchers come up with new hypotheses after they find that their original hypothesis was insignificant. This is problematic because it is intellectually dishonest and hides insignificant results in the interest of only reporting those which are significant (often called the File Drawer Effect). Remember, a p-value of 0.05 means that there is a 5% chance of the observations being as they are if the null hypothesis is true. Thus, if you test 20 variables, it is statistically likely that at least one of them will show significance simply by chance.

R Markdown Formatting Tricks (& Knitting)

R markdown has a ton of useful syntax tricks for formatting text in the knitted version of your document. These include:

italics

bold

superscript¹⁰ strikethrough

Large Header

Medium-Larger Header

Medium Header

Small-Medium Header

Small header

- Bulleted list
 - Sub-item in list
- 1. First list item
- 2. Second list item
- 3. Third list item

To test out some of these tricks yourself, you'll have to knit your document to see what it looks like outside of R studio. My favorite file type to knit to is a pdf, because these are easy to read, save, and send. You can set the file type at the top of your document and then hit the knit button (refer to figure 3). It's important to note that in order to knit, you can't have any errors in your code. If you have any non-functional code, the file won't be able to knit. You also need to ensure that all your code chunks are named with a unique name in order to knit. If you try knitting and it doesn't work, check that you've met both of those requirements and try again.

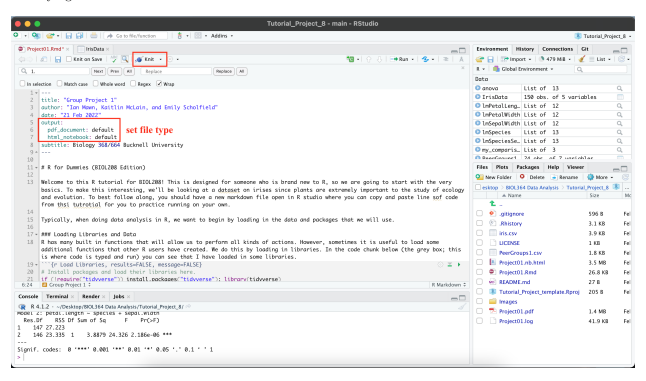


Figure 3: Knitting

Acknowledgements

1. Ritchie, Stuart. Science Fictions (2020) pp. 98–99

Group Member Contributions

- 1. Ian: Loading Libraries and Data, Using R, Exploring Your Data
- 2. Kaitlin: Tutorial screenshots, Additions to Using R, Visualization with Graphs, shapiro tests, ANOVA, t-tests, Linear Models, Markdown Formatting, GitHub

3. Emily: Log Transformation, p-value Discussion, README file