# Final Project

#### Your Name Here

2023-11-29

## Final Details

#### Purpose

The goal of this final assignment is to assess your ability to integrate the many skills you have learned over the semester: filtering and summarizing data, creating new columns, choosing the appropriate data visualizations, and performing and interpreting the appropriate statistical tests.

#### Task

Write R code which produces the correct data, summaries, plots and analyses. Correctly interpret the results of these plots and analyses.

#### Criteria for Success

- Code chunks run without errors
- Code produces the correct result
  - Code that produces the correct answer will receive **full** credit
  - Code attempts with logical direction will receive **partial** credit
- Appropriate plot types are used to visualize the data
- Appropriate statistical tests are used to analyze the data
- Written answers address the questions in sufficient detail

## **Due Date**

Dec 11 at midnight MST

#### Final

For your final this semester, I am presenting you with 3 problem sets, totaling up to 60 points.

I'm expecting you to be able to filter and summarize the data in the ways you need, choose the appropriate visualization, choose the appropriate analysis, and correctly interpret the analysis for the question I've asked you.

It is important to note that I will not be giving you an answer key for this final project since a major part of your grade for the assignment is being able to choose the appropriate visualizations and analyses for the question and the data.

We are going to use the palmerpenguins R package and data set, which we've used many times before! You can learn more about it here. This is a real data set from a Long-Term Ecological Research (LTER) site in Antarctica.

## Set-Up

Be sure to run both of these code chunks before you begin! I've gone ahead and included the code to load the two packages you will need to successfully complete this project: the tidyverse and palmerpenguins. Be sure to run this code chunk!

```
library(tidyverse)
library(palmerpenguins)
```

**Important!** I've also included one more code chunk below. Be sure to run this code chunk, as well! It does two key things:

- first, it adds the penguins data frame to your environment, which I imagine you will find helpful
- second, it removes all rows that have any NA values, which will make completing this assignment a bit easier.

Once you've run this line of code, you should see the **penguins** data frame pop up in your environment with 333 observations (rows) and 8 variables (columns).

penguins <- penguins %>% drop\_na()



### Structure & Guidelines

Like the practice version, this final is structured as 3 different problem sets. For each problem set, I am presenting you with an initial question to guide your thinking and analysis.

#### Data

Assume that nothing carries over between problem sets.

Each problem set is stand-alone, meaning that you should always start with the penguins data frame at the beginning of each problem set. If you should use a data frame that you created *within* the problem set, I explicitly state so.

For example, in Problem Set 1, you should use the biscoe data frame that you create for the entire problem set; at the start of Problem Set 2, start over with the penguins data frame.

#### **Interpreting Statistical Results**

When I ask you to interpret statistical results, you should roughly follow these guidelines.

- the cut-off for our p-values is always 0.05
- report the p-value that we are focused on
- if there are multiple p-values of interest, report all of them
- state whether the p-value indicates a significant difference/relationship
- if applicable, state whether we should or should not reject the null hypothesis

#### **Plotting**

All plots should be made using ggplot2.

Your options for plot types to choose from are:

- multiple histogram plots
  - use transparency (alpha)
  - use position = "identity" with multiple groups to see the full distributions
- multiple density plots
  - use transparency (alpha) with multiple groups to see the full distributions
- box-and-whisker plot
  - add points on top of the box plot to show the distribution of the points
- scatter plot
  - add the linear model to every scatter plot

#### Note: All plots should have modified axis labels and legend labels.

In many cases, this might mean capitalizing the axis label or legend label. In other cases, you might want to put units in parentheses after the words (e.g., Body Mass (g)).

#### Generative AI Guidelines

While I do not recommend using generative AI to complete this assignment (throughout the semester, I've seen it cause more problems than not...), you are allowed to use it on the final.

However, you must do so according to the guidelines I've set out for all other assignments this semester.

You can find clear examples of these guidelines on D2L in Resources > Helpful Info > Assignment Reminders.

**CAUTION!** If you do use generative AI without following the course guidelines, you will receive a 0 for this final and you will not have the option to resubmit. Proceed wisely.

## Let's Begin!

- Have you run the 2 code chunks in the "Set-Up" section?
- Have you read through the "Structure & Guidelines" section?

If so, proceed! If not, scroll back up...

## Problem Set 1 (30 points total)

This problem set has 2 parts.

Part 1 (20 points)

Question: Is there a significant relationship between bill length and bill depth for penguins on Biscoe Island? For this problem set, we are going to use data from Biscoe island only.

1. Create a new data frame called biscoe that includes only penguins from Biscoe island. This new data frame should have 163 rows. (2 points)

```
biscoe <- penguins %>%
filter(island == "Biscoe")
```

You will want to use the biscoe data set for the rest of Problem Set #1.

This is a scenario where there is no independent and no dependent variable. Go ahead and **treat bill length** as the independent variable (x-axis) and bill depth as the dependent variable (y-axis).

- 2. Determine whether bill length and bill depth are continuous or categorical (2 points)
  - bill depth: dependentbill length: independent

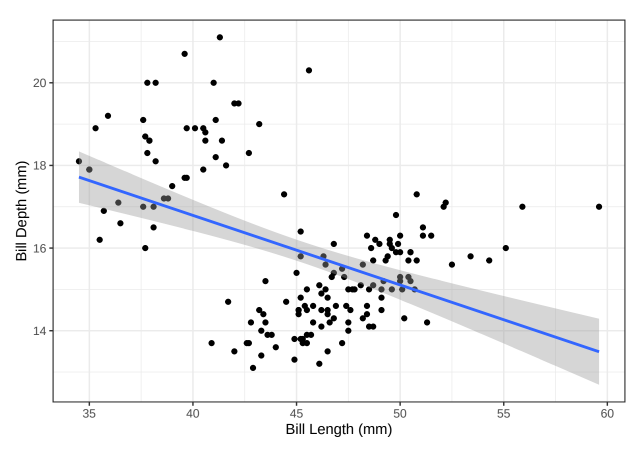
For now, ignore species. We will address species in Part 2 of the problem set.

3. Write the correct pair of statistical hypotheses for our question. (2 points)

Null: no relationship between bill length and bill depth
Alternative: there is a relationship between bill length and bill depth

- 4. Plot the relationship between bill length and bill depth using the appropriate plot type. (3 points)
  - Be sure to add a line of best fit
  - Ensure that the plot has clear labels on the axes (follow the Structure & Guidelines).
  - Remember, we are ignoring species for now.

## 'geom\_smooth()' using formula = 'y ~ x'



- 5. Describe the relationship that you see in the plot you just made. Is the relationship positive, negative, or nonexistent? How do you know? (2 points)
  - Answer: negative; downward slope, high values associated with low values on the other axis
- 6. Calculate the correlation coefficient, r. What does this value confirm for us about the relationship (positive, negative, no relationship)? (2 points)

```
r <- cor(biscoe$bill_length_mm, biscoe$bill_depth_mm)
r</pre>
```

## [1] -0.4446658

Answer: negative relationship

7. Calculate the  $r^2$  value. Remember, this number is typically expressed as a percent (x 100). What is this value telling us? (2 points)

```
r^2 * 100
```

## [1] 19.77277

Answer: About 20% variation explained

8. Let's see if there is a significant relationship between bill length and bill depth. Perform the correct statistical analysis (1 point) and interpret the results. (5 points total)

```
summary(lm(data = biscoe, bill_depth_mm ~ bill_length_mm))
```

```
##
## Call:
## lm(formula = bill_depth_mm ~ bill_length_mm, data = biscoe)
## Residuals:
##
      Min
               10 Median
                               3Q
                                      Max
## -3.2027 -1.2536 -0.1135 1.0735 4.5279
##
## Coefficients:
                 Estimate Std. Error t value Pr(>|t|)
##
                 23.52509
                             1.21614 19.344 < 2e-16 ***
## (Intercept)
## bill_length_mm -0.16835
                             0.02673 -6.299 2.74e-09 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 1.642 on 161 degrees of freedom
## Multiple R-squared: 0.1977, Adjusted R-squared: 0.1927
## F-statistic: 39.68 on 1 and 161 DF, p-value: 2.738e-09
```

- What is the equation of the line of best fit? Use both variable names and values from the statistical analysis. (1 point)
- What is the p-value? (1 point)
- Is there a significant relationship? (1 point)
- What should we do with the null hypothesis? (1 point)

Answer: bill\_depth = -0.168\*bill\_length + 23.5; p = 2.74e-9, highly significant

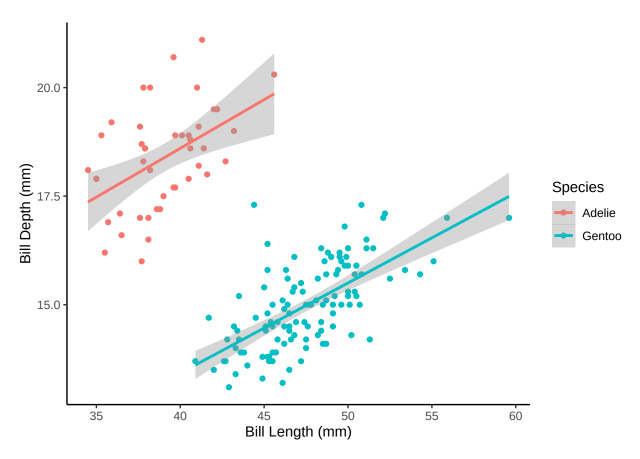
#### Part 2 (10 points)

When we look at the plot of the data, it looks like there might be two different groups in the data. Let's see what happens when we add in species to this analysis.

9. Make the color of the points and the linear models differ by species on Biscoe Island. Be sure to adjust all labels on the plot accordingly. (3 points)

```
ggplot(biscoe, aes(bill_length_mm, bill_depth_mm, color = species)) +
  geom_point() +
  geom_smooth(method = "lm") +
  labs(x = "Bill Length (mm)",
        y = "Bill Depth (mm)",
        color = "Species") +
  theme_classic()
```

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



10. Run the appropriate statistical test, adding species (and the interaction between the two independent variables) into the model. (2 points)

```
summary(lm(data = biscoe, bill_depth_mm ~ bill_length_mm * species))
```

```
##
## Call:
## lm(formula = bill_depth_mm ~ bill_length_mm * species, data = biscoe)
##
## Residuals:
##
        Min
                  1Q
                       Median
                                     3Q
                                             Max
   -2.08441 -0.64479 -0.02957
                               0.46954
                                         2.96109
##
##
## Coefficients:
##
                                 Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                                             2.02455
                                  9.62634
                                                       4.755 4.42e-06 ***
## bill_length_mm
                                 0.22435
                                             0.05184
                                                       4.328 2.66e-05 ***
## speciesGentoo
                                 -4.50539
                                             2.34915
                                                      -1.918
                                                                0.0569
                                             0.05755
                                                     -0.291
                                                                0.7715
## bill_length_mm:speciesGentoo -0.01674
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.8434 on 159 degrees of freedom
```

```
## Multiple R-squared: 0.791, Adjusted R-squared: 0.7871
## F-statistic: 200.6 on 3 and 159 DF, p-value: < 2.2e-16</pre>
```

11. Interpret the results of the test above. Is species significant? Is there a significant interaction between bill length and species? (1 points)

Answer: neither species nor interaction is significant

12. Based on the plots, the linear relationships, and the linear models, write 2-3 sentences discussing if and/or how adding species into our linear model changes our interpretation of the data. Did the type of relationship change? Did significance levels change? Do the two linear models tell us different things? (3 points)

Answer: negative to positive; models still tell us similar things in terms of significance, though

## Problem Set 2 (15 points)

Question: Is there a difference in the average flipper length between male and female Chinstrap penguins?

1. Our first step is to create a new data frame that includes only Chinstrap penguins. Call this new data frame chinstrap. (1 point)

```
chinstrap <- penguins %>%
  filter(species == "Chinstrap")
```

We will be using the chinstrap data frame for the rest of this problem set.

2. Let's summarize our data. Calculate one measure of central tendency and one measure of variability of the flipper length column for *each* sex: male and female. Save this dataframe as chinstrap\_summary. (2 points)

- 3. What if we wanted our summary data in centimeters instead of millimeters? We can acheive this in 2 steps.
  - a. First, create a function that will convert a number from millimeters to centimeters. There are 10 millimeters in a centimeter. (1 point)

```
mm_to_cm <- function(mm) {
  cm <- mm / 10
  return(cm)
}</pre>
```

b. Now, using the same code from question 3 above, add one line of code (in the correct location) that uses your newly created mm\_to\_cm function to create a new column in the dataframe. Make sure to edit the arguments in summary functions accordingly, as well. The result should be the same data frame that you produced in question 3 but with the summary values in cm instead of mm.(1 point)

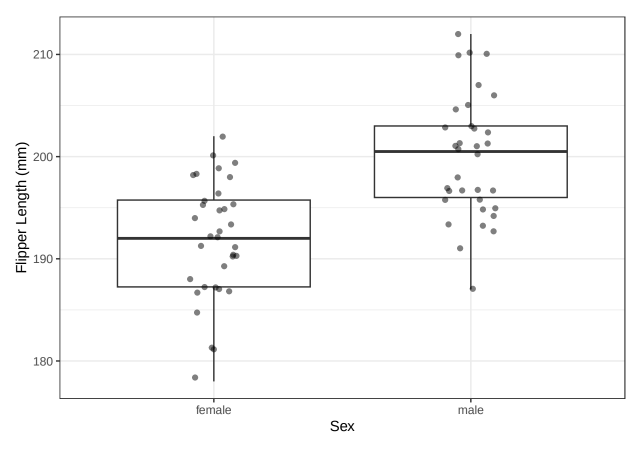
Ok, we have summarized the flipper length data for our two groups!

Let's get back to the chinstrap dataframe (not the summary dataframe), and keep working.

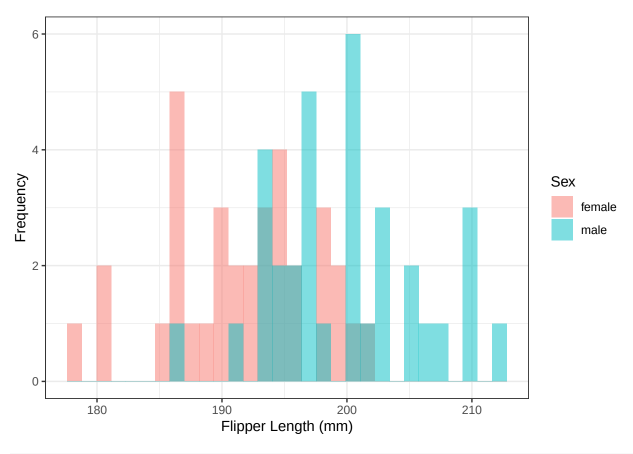
- 4. Determine which variable is *dependent* and which is *independent*. Also determine if each variable is *continuous* or *categorical*. (1 points)
- flipper length: dependent, continuous
- species: independent, categorical

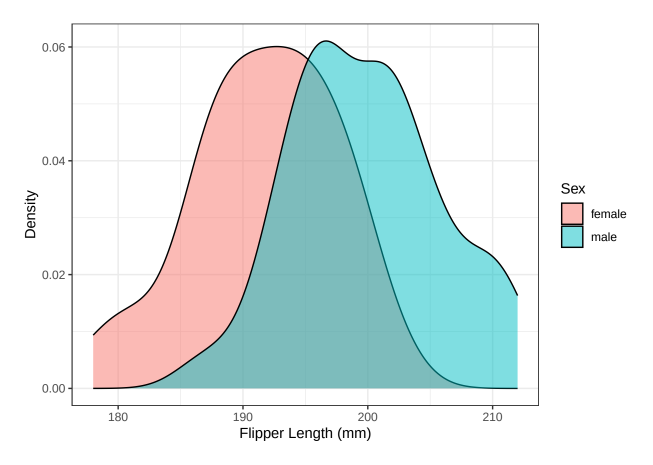
Let's plot the flipper length data for each sex.

5. Choose an appropriate plot for the data. Be sure to adjust the x- and y-axis labels appropriately. (2 points)



## 'stat\_bin()' using 'bins = 30'. Pick better value with 'binwidth'.





6. Write the pair of statistical hypotheses for our question. (1 points)

Null: no difference in the mean flipper length between male and female chinstrap penguins Alternative: true difference in the mean flipper length between male and female chinstrap penguins

7. Perform the appropriate analysis to compare the flipper lengths of male and female chinstrap penguins. (2 points)

```
t.test(data = chinstrap, flipper_length_mm ~ sex)
```

```
##
##
   Welch Two Sample t-test
##
## data: flipper_length_mm by sex
## t = -5.7467, df = 65.905, p-value = 2.535e-07
## alternative hypothesis: true difference in means between group female and group male is not equal to
## 95 percent confidence interval:
   -11.017272 -5.335669
## sample estimates:
##
  mean in group female
                          mean in group male
##
               191.7353
                                    199.9118
```

- 8. Interpret the results of this test. (2 points)
  - is there a significant difference?
  - what does that significant difference mean?
  - should we reject the null hypothesis?

Answer: yes, there is a significant difference  $(p = 2.535 \times 10^{-7})$ ; reject null

9. Should we run pairwise comparisons? If no, explain why not. If yes, do so and interpret the results. (2 points)

Answer: no, t-test is comparing just one pair

## Problem Set 3 (15 points)

Question: Are there differences in the average bill length across the 3 islands in the data set: Dream, Biscoe, and Torgersen? (Ignore species) Let's start by summarizing the bill length data.

1. Calculate at least one measure of central tendency and one measure of variability for the bill length for *each* island. (2 points)

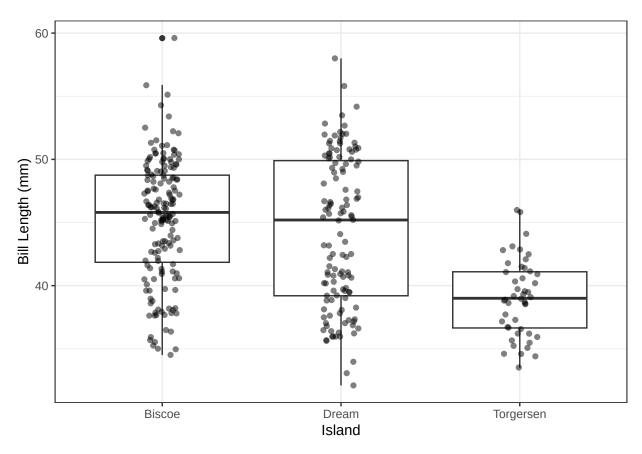
```
## # A tibble: 3 x 4
##
     island
               min_bill max_bill mean_bill
##
     <fct>
                            <dbl>
                                       <dbl>
                   <dbl>
## 1 Biscoe
                    34.5
                             59.6
                                        45.2
## 2 Dream
                    32.1
                             58
                                        44.2
## 3 Torgersen
                    33.5
                              46
                                        39.0
```

- 2. Which of our variables would be considered *independent* and which one *dependent*? Also determine whether each is *continuous* or *categorical*. (2 points)
  - island: independent, categorical
  - bill length: dependent, continuous

Now that we have an idea numerically of the differences between the islands, let's plot the differences.

3. Choose an appropriate plot. Ensure that you follow the plotting guidelines in the Structure & Guidelines section above! (2 points)

```
ggplot(penguins, aes(island, bill_length_mm)) + # adding color is optional
geom_boxplot() +
geom_jitter(width = 0.1, alpha = 0.5) +
labs(x = "Island",
    y = "Bill Length (mm)") + # if adding color, must change color label
theme_bw() # can choose any theme but must include
```



```
# OR

ggplot(penguins, aes(bill_length_mm, fill = island)) +

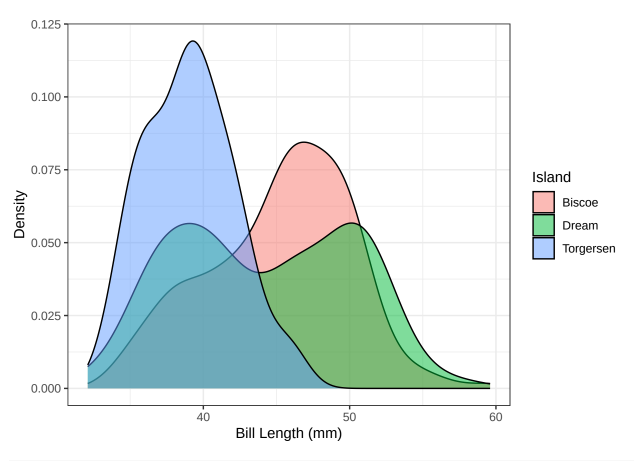
geom_density(alpha = 0.5) +

labs(x = "Bill Length (mm)",

y = "Density",

fill = "Island") +

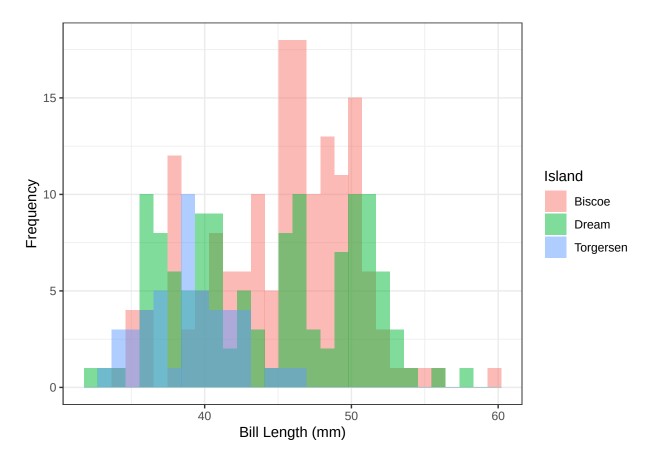
theme_bw()
```



```
# OR

ggplot(penguins, aes(bill_length_mm, fill = island)) +
  geom_histogram(alpha = 0.5, position = "identity") +
  labs(x = "Bill Length (mm)",
        y = "Frequency",
        fill = "Island") +
  theme_bw()
```

## 'stat\_bin()' using 'bins = 30'. Pick better value with 'binwidth'.



4. Write the correct pair of statistical hypotheses for our question. (2 points)

Null: no difference in the mean bill length between penguins on different islands
Alternative: true difference in the mean bill length between penguins on different islands

5. Run the appropriate statistical analysis for our question. (2 points)

```
aov_model <- aov(data = penguins, bill_length_mm ~ island)
summary(aov_model)</pre>
```

```
## Df Sum Sq Mean Sq F value Pr(>F)
## island 2 1417 708.6 27.47 9.21e-12 ***
## Residuals 330 8512 25.8
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
```

- 6. Interpret the results of this test. (3 points)
  - what is the p-value?
  - is there a significant difference?
  - should we reject the null hypothesis?

Answer: yes, p = 9.21e-12, which is smaller than 0.05, reject null. penguins on different islands have significantly different beak lengths

7. Should we run pairwise comparisons? If no, explain why not. If yes, do so and interpret the results. (2 points)

Answer: yes; significant differences between all pairs except Dream-Biscoe because p > 0.05

## TukeyHSD(aov\_model)

```
Tukey multiple comparisons of means
##
##
       95% family-wise confidence level
##
## Fit: aov(formula = bill_length_mm ~ island, data = penguins)
##
## $island
##
                         diff
                                    lwr
                                               upr
                                                       p adj
## Dream-Biscoe
                    -1.026515 -2.454611 0.4015809 0.2096455
## Torgersen-Biscoe -6.210168 -8.189815 -4.2305220 0.0000000
## Torgersen-Dream -5.183653 -7.234077 -3.1332293 0.0000000
```

## The End!

Great work, and thanks for a wonderful semester!