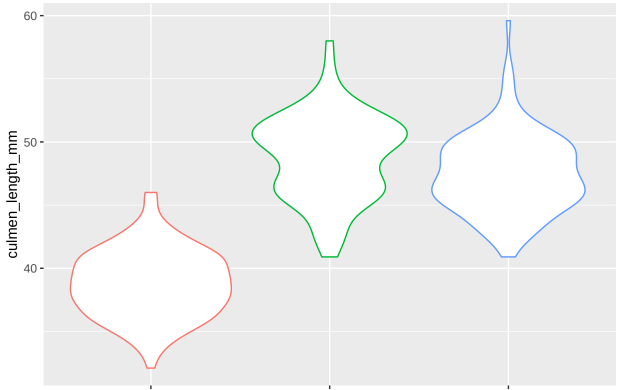
# MT3 Computer Assignment - Reproducible\_Figures\_R

#### 2023-12-04

 $\#\#\mathrm{Q1}$ ) Create a figure using the Palmer penguin dataset that is correct but badly communicates the data. Do not make a box plot.  $\#\#\mathrm{A}$ ) Figure inserted below:

```
library(ggplot2)
library(palmerpenguins)
library(janitor)
##
## Attaching package: 'janitor'
## The following objects are masked from 'package:stats':
##
##
       chisq.test, fisher.test
library(dplyr)
## Attaching package: 'dplyr'
## The following objects are masked from 'package:stats':
##
##
       filter, lag
  The following objects are masked from 'package:base':
##
##
       intersect, setdiff, setequal, union
## Warning: Removed 2 rows containing non-finite values ('stat_ydensity()').
```



Adelie Penguin (Pygoscelis ad laine)strap penguin (Pygoscelis antar lie anta

##B) Write about how your design choices mislead the reader about the underlying data. ##While technically this figure is not incorrect, this violin plot has some issues representing the data. The violin plot does not show individual data points and their distributions so it is difficult to observe the true nature of the data. It is therefore easy to misinterpret the data. Due to the natural complexity of the violin plot it can be inaccesible to those who are not familiar with this plot.

##Q2)Write a data analysis pipeline in your rmd. You should be aiming to write a clear explanation of the steps as well as clear code.

##Firstly you need to install the following packages:

```
install.packages(c("ggplot2", "palmerpenguins", "janitor", "dplyr"))
```

##You then need to load the packages now that they are installed:

```
library(ggplot2)
library(palmerpenguins)
library(janitor)
library(dplyr)
```

##Next you need to load in the data set

```
head(penguins)
```

```
## # A tibble: 6 x 8
## species island bill_length_mm bill_depth_mm flipper_length_mm body_mass_g
```

```
##
     <fct>
             <fct>
                                 <dbl>
                                                <dbl>
                                                                   <int>
                                                                               <int>
                                                 18.7
                                                                                3750
## 1 Adelie Torgersen
                                  39.1
                                                                     181
## 2 Adelie Torgersen
                                  39.5
                                                 17.4
                                                                     186
                                                                                3800
                                                                                3250
## 3 Adelie Torgersen
                                  40.3
                                                 18
                                                                     195
## 4 Adelie Torgersen
                                  NA
                                                 NA
                                                                     NA
                                                                                  NA
## 5 Adelie Torgersen
                                  36.7
                                                 19.3
                                                                     193
                                                                                3450
## 6 Adelie Torgersen
                                                 20.6
                                  39.3
                                                                     190
                                                                                3650
## # i 2 more variables: sex <fct>, year <int>
```

##Cleaning data by firstly looking at the column names

#### names (penguins)

```
## [1] "species" "island" "bill_length_mm"
## [4] "bill_depth_mm" "flipper_length_mm" "body_mass_g"
## [7] "sex" "year"
```

##Looking at the summary of data to see what is happening

### summary(penguins)

```
##
                           island
                                     bill_length_mm
                                                     bill_depth_mm
         species
##
                                            :32.10
    Adelie
             :152
                              :168
                                     Min.
                                                     Min.
                                                            :13.10
                    Biscoe
                                     1st Qu.:39.23
                                                     1st Qu.:15.60
##
    Chinstrap: 68
                    Dream
                              :124
                                     Median :44.45
                                                     Median :17.30
##
    Gentoo
            :124
                    Torgersen: 52
##
                                     Mean
                                            :43.92
                                                     Mean
                                                            :17.15
##
                                                     3rd Qu.:18.70
                                     3rd Qu.:48.50
##
                                            :59.60
                                     Max.
                                                     Max.
                                                             :21.50
##
                                     NA's
                                            :2
                                                     NA's
                                                             :2
##
   flipper_length_mm body_mass_g
                                          sex
                                                         year
##
  Min.
           :172.0
                      Min.
                              :2700
                                      female:165
                                                   Min.
                                                           :2007
##
   1st Qu.:190.0
                      1st Qu.:3550
                                      male :168
                                                   1st Qu.:2007
## Median :197.0
                      Median:4050
                                      NA's : 11
                                                   Median:2008
           :200.9
                                                           :2008
## Mean
                      Mean
                              :4202
                                                   Mean
##
    3rd Qu.:213.0
                      3rd Qu.:4750
                                                   3rd Qu.:2009
##
   Max.
           :231.0
                      Max.
                              :6300
                                                   Max.
                                                           :2009
##
  NA's
           :2
                      NA's
                              :2
```

##Realise that there are some N/A results and decide to remove them

```
penguins <- na.omit(penguins)</pre>
```

##Shortening of island names:

```
Clean_data <- penguins %>%
  mutate(island = substr(island,1,3))
```

##Also shortening species names which can be seen in this pipeline below ##Pipeline for this cleaning:

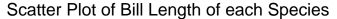
```
Clean_data <- penguins %>%
  na.omit() %>%
  mutate(island = substr(island,1,3)) %>%
  mutate(species = substr(species,1,3))
```

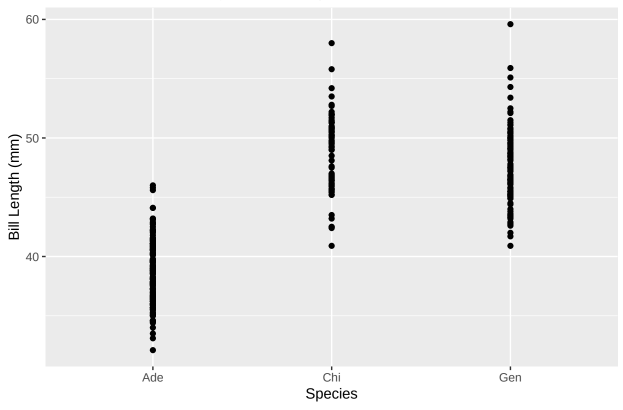
 $\#\#\#\mathrm{Observing}$  the cleaning made to the data

```
head(Clean_data)
```

```
## # A tibble: 6 x 8
     species island bill_length_mm bill_depth_mm flipper_length_mm body_mass_g
     <chr>
             <chr>
                              <dbl>
                                            <dbl>
                                                               <int>
                                                                           <int>
                               39.1
                                                                            3750
## 1 Ade
             Tor
                                             18.7
                                                                 181
## 2 Ade
             Tor
                               39.5
                                             17.4
                                                                 186
                                                                            3800
## 3 Ade
             Tor
                               40.3
                                             18
                                                                 195
                                                                            3250
## 4 Ade
                               36.7
                                                                 193
                                                                            3450
             Tor
                                             19.3
## 5 Ade
             Tor
                               39.3
                                             20.6
                                                                 190
                                                                            3650
## 6 Ade
             Tor
                               38.9
                                             17.8
                                                                 181
                                                                            3625
## # i 2 more variables: sex <fct>, year <int>
```

##Explanatory figure





## Null hypothesis: there is no significant difference in mean bill length between the three different species of penguin.

##Alternative hypothesis: there is signficant difference in mean bill length between atleast two of the three species.

##Run a statistical test. I decided to do an ANOVA test looking at the differences between mean bill length across the different species.

```
model <- lm(bill_length_mm ~ species, data = Clean_data)
anova_result <- anova(model)</pre>
```

##It is time to analyse the results of the anova...

```
print(anova_result)
```

##This anova has produced an incredibly small p value it almost practically 0. This tells us that there is strong evidence that the results are highly statistically significant. The null hypothesis is infact wrong and there are differences in mean bill length among species.

I decided to conduct a post-hoc test using the Tukey method as the results of my anova suggest there are significant differences and I want to see which species differ from eachother in particular.

```
library(agricolae)
# Conduct Tukey's HSD post-hoc test
tukey_result <- HSD.test(model, "species")</pre>
# Print the results
print(tukey_result)
## $statistics
##
      MSerror Df
                      Mean
##
     8.828839 330 43.99279 6.754143
##
## $parameters
##
      test name.t ntr StudentizedRange alpha
##
     Tukey species
                                3.329537 0.05
##
## $means
                                                                  Q50
                                                                         Q75
##
       bill_length_mm
                                           se Min Max
                                                            Q25
                            std
                                  r
             38.82397 2.662597 146 0.2459095 32.1 46.0 36.725 38.85 40.775
## Ade
## Chi
             48.83382 3.339256 68 0.3603275 40.9 58.0 46.350 49.55 51.075
## Gen
             47.56807 3.106116 119 0.2723819 40.9 59.6 45.350 47.40 49.600
##
## $comparison
## NULL
##
## $groups
##
       bill_length_mm groups
## Chi
             48.83382
## Gen
             47.56807
                            b
## Ade
             38.82397
##
## attr(,"class")
```

##The results of this test suggest Chinstrap has a significantly different mean bill length compared to Gentoo and Adelie. Gentoo and Adelie do not show a significant difference in mean bill length.

##In order to visualise the results of these statistical tests I chose to create a dot-plot:

## [1] "group"

```
library(ggplot2)

tukey_df <- as.data.frame(tukey_result$groups)</pre>
```

```
tukey_df$groups <- factor(tukey_df$groups, levels = c("a", "b", "c"), labels = c("Ade", "Gen", "Chi"))

ggplot(tukey_df, aes(x = groups, y = bill_length_mm, color = groups)) +
    geom_point(position = position_dodge(width = 0.8), size = 3) +
    labs(title = "Tukey's HSD Test",
        x = "Species",
        y = "Mean Bill Length (mm)",
        color = "Species") +
    theme_minimal()</pre>
```



## The results of my statistical tests as shown in the figure above clearly present that there is a significant difference in mean bill length between at least two of the three species. We can therefore reject our null hypothesis that states there is no significant differences in mean bill length between the three species.

Gen Species Chi

##Q3) Open Science:

Ade

- ##A) My github link: https://github.com/BioBabe2002/Reproducible-Figures-R
- ##B) Partner's github link: https://github.com/Elephant34/ReproducibleScience
- ##C) Reflect on your experience running their code. What elements of their code helped you to understand their data pipeline? Did it run and did you need to fix anything? What suggestions would you make for improving their code to make it more understandable or reproducible and why? If you needed to alter your partner's figure using their code, do you think that would be easy or difficult and why?

My partner's code was successful in allowing me to understand their data pipeline for a number of reasons. Firstly, their code was continously annotated. This let me know exactly what function a line of code had so that I could follow their process. Secondly, they handily make note of where figures can be found in their repo which was helpful to compare to when running the code myself. Their code ran very smoothly without need for alteration. I would argue that the removal of N/A values could be included in their "DataCleaning" block but this is a minimal comment. It would also be useful to contain session information indicating the versions of software used. It would also be easy to alter my partner's figure using their code as the steps appear to be very transparent and possible for me to integrate my own desired code.

##D) Reflect on your own code based on your experience with your partner's code and their review of yours.

##Their review was very plausible. I avoided added in code to install given packages but in hindsight I could have made note that these packages should be installed and even produced code to only run install.packages if the package is not already installed which would look hopefully something like this:

```
##firstly checking that the Agricolae package is installed:
if (!requireNamespace("Agricolae", quietly = TRUE)) {
    # then if it is not installed, it will install
    install.packages("Agricolae", dependencies = TRUE)
}
```

##My partner also comments that by splitting the pipeline into seperate functions this would improve readibility and I agree. By doing so it would make my code easier to follow along and reproduce. ##Writing code for other people has helped me to generate my clarity in my work. When coding for myself it is easy to make brief notes that only I would understand or lack thereof hoping I will know what it means when I return to my work. By outlining descriptions of my steps taken it ensures that my work is useful.