Penguin Assignment

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Organising and cleaning data:

Setting working directory as "PenguinProjects" folder (please adjust to suit your working directory):

```
setwd("~/RWorkingDirectory/PenguinProjects")
```

Loading in libraries, data and functions:

```
source("functions/libraries.r")
source("functions/cleaning.r")
source("functions/plotting.r")
```

Creating a new folder for the raw data and saving the raw data:

```
#Creating a new data folder
dir.create("data")

#Creating a new raw data folder
dir.create("data/data_raw")

#Saving the raw data
write.csv(penguins_raw, "data/data_raw/penguins_raw_06-12-2022.csv")
```

Cleaning the data, creating a new folder for the clean data and saving the clean data:

```
#Using function from "cleaning.r" to clean the data
penguins_clean <- cleaning(penguins_raw)

#Creating a new clean data folder
dir.create("data/data_clean")

#Saving the clean data
write.csv(penguins_clean, "data/data_clean/penguins_clean_06-12-2022.csv")</pre>
```

Looking at the data:

```
head(penguins_clean)
```

Preparing data for analysis:

Removing rows without sex or body mass recorded and defining variables:

```
#Using function from "cleaning.r" to prepare data
penguins_mass <- remove_empty_body_mass(penguins_clean)

#Defining factors and numerical variables
penguins_mass$species <- as.factor(penguins_mass$species)
penguins_mass$sex <- as.factor(penguins_mass$sex)
penguins_mass$body_mass_g <- as.numeric(penguins_mass$body_mass_g)

#Saving mass data
write.csv(penguins_mass, "data/data_clean/penguins_mass_06-12-2022.csv")</pre>
```

Looking at the data:

```
head(penguins_mass, 3)
```

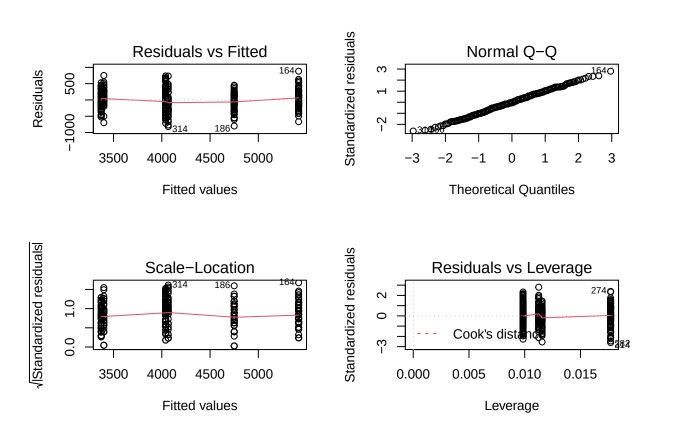
```
## # A tibble: 3 x 3
## species sex body_mass_g
## <fct> <fct> <fct> <dbl>
## 1 Adelie Penguin (Pygoscelis adeliae) MALE 3750
## 2 Adelie Penguin (Pygoscelis adeliae) FEMALE 3800
## 3 Adelie Penguin (Pygoscelis adeliae) FEMALE 3250
```

Analysing the data:

Q: To what extent does the species and sex of a penguin influence body mass?

Creating a model with no interaction and testing assumptions of statistical analysis:

```
#Creating the model
lmSpecies_Sex_Mass_No_Interaction <- aov(body_mass_g ~ species + sex, data = penguins_mass)
#Plotting to test assumptions
par(mfrow = c(2,2))
plot(lmSpecies_Sex_Mass_No_Interaction)</pre>
```



Overall, the data appears to meet assumptions of ANOVA.

Running ANOVA:

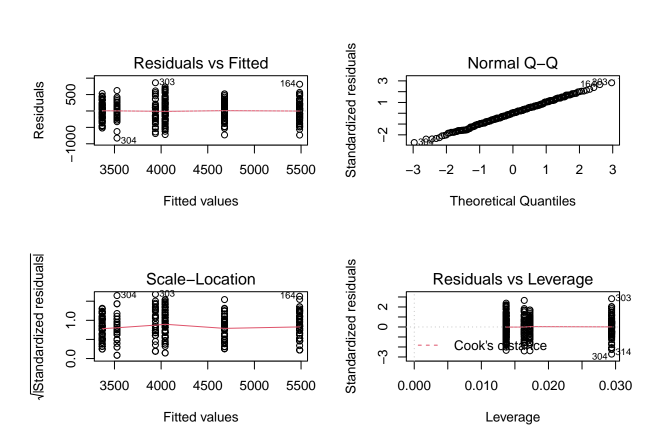
```
summary(lmSpecies_Sex_Mass_No_Interaction)
```

```
## Df Sum Sq Mean Sq F value Pr(>F)
## species    2 145190219 72595110          724.2 <2e-16 ***
## sex          1 37090262 37090262 370.0 <2e-16 ***
## Residuals 329 32979185 100241
## ---
## Signif. codes:    0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1</pre>
```

Both species and sex appear to have a significant effect on body mass.

Building a model that takes into account interaction between species and sex:

```
#Creating the model
lmSpecies_Sex_Mass_Interaction <- aov(body_mass_g ~ species * sex, data = penguins_mass)
#Plotting to test assumptions
par(mfrow = c(2,2))
plot(lmSpecies_Sex_Mass_Interaction)</pre>
```



Overall, the data appears to fit the assumptions of ANCOVA.

Running ANCOVA:

summary(lmSpecies_Sex_Mass_Interaction)

```
##
                Df
                      Sum Sq Mean Sq F value
                                                Pr(>F)
## species
                 2 145190219 72595110 758.358
                    37090262 37090262 387.460
                                               < 2e-16 ***
##
                 2
                     1676557
                               838278
                                        8.757 0.000197 ***
## species:sex
## Residuals
               327
                    31302628
                                95727
## Signif. codes:
                   0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
```

There is a significant effect of interaction between species and sex on body mass.

Comparing the two models:

anova(lmSpecies_Sex_Mass_No_Interaction,lmSpecies_Sex_Mass_Interaction)

```
## Analysis of Variance Table
##
## Model 1: body_mass_g ~ species + sex
## Model 2: body_mass_g ~ species * sex
## Res.Df RSS Df Sum of Sq F Pr(>F)
## 1 329 32979185
## 2 327 31302628 2 1676557 8.757 0.0001973 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Adding an interaction between species and sex significantly improves the model.

Running a Tukey HSD test:

The Tukey HSD test suggests significant differences between body mass of penguins grouped by species and sex

The comparisons between groups which are not significantly different are:

- 1) Chinstrap females and Adelie females
- 2) Chinstrap males and Adelie males

Producing figures to display the data:

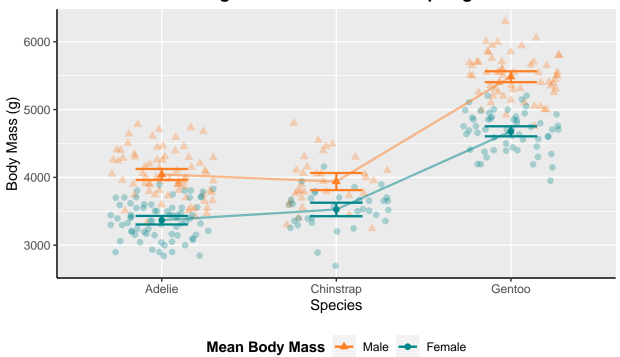
Creating a new folder to store figures in:

```
dir.create("figures")
```

Producing an interaction plot:

```
#Using function from "plotting.R" to produce graph
plot_mass_figure(penguins_mass)
```

Interaction Plot of Body Mass, Species and Sex of Penguins in the Palmer Archipelago:



Error bars represent 95% confidence intervals

Saving the figure as a PNG file:

```
#Using function from "plotting.R" to save figure as PNG save_mass_plot_png(penguins_mass, "figures/fig_01_06-12-2022.png", 20, 600, 1.4)
```

Saving the figure as an SVG file:

```
#Using function from "plotting.R" to save figure as SVG save_mass_plot_svg(penguins_mass, "figures/fig_01_06-12-2022.svg", 20, 1.4)
```

Session info:

sessionInfo()

```
## R version 4.0.5 (2021-03-31)
## Platform: x86_64-w64-mingw32/x64 (64-bit)
## Running under: Windows 10 x64 (build 22000)
## Matrix products: default
##
## locale:
## [1] LC_COLLATE=English_United Kingdom.1252
## [2] LC_CTYPE=English_United Kingdom.1252
## [3] LC_MONETARY=English_United Kingdom.1252
## [4] LC NUMERIC=C
## [5] LC_TIME=English_United Kingdom.1252
##
## attached base packages:
## [1] stats
                 graphics grDevices utils
                                                datasets methods
                                                                    base
## other attached packages:
                            ragg_1.2.2
## [1] svglite_2.1.0
                                                  dplyr 1.0.7
## [4] janitor_2.1.0
                                                  palmerpenguins_0.1.1
                            ggplot2_3.3.3
##
## loaded via a namespace (and not attached):
## [1] Rcpp_1.0.8.3
                            lubridate_1.8.0
                                                 lattice_0.20-41
## [4] deldir 1.0-6
                                                 assertthat 0.2.1
                            png 0.1-7
## [7] digest 0.6.27
                            utf8 1.2.1
                                                 R6 2.5.1
## [10] backports_1.4.1
                            evaluate_0.18
                                                 highr 0.9
## [13] pillar_1.6.0
                            rlang_0.4.11
                                                 data.table_1.14.2
## [16] rstudioapi_0.14
                            rpart_4.1-15
                                                 Matrix_1.4-1
## [19] checkmate_2.0.0
                            rmarkdown_2.18
                                                 labeling 0.4.2
## [22] textshaping_0.3.6
                            splines_4.0.5
                                                 stringr_1.4.0
## [25] foreign_0.8-81
                            htmlwidgets_1.5.4
                                                 munsell_0.5.0
## [28] compiler_4.0.5
                            xfun_0.22
                                                 pkgconfig_2.0.3
## [31] systemfonts_1.0.4
                            base64enc_0.1-3
                                                 htmltools_0.5.2
## [34] nnet_7.3-15
                            tidyselect_1.1.1
                                                 htmlTable_2.4.1
## [37] gridExtra_2.3
                                                 Hmisc_4.7-0
                            tibble_3.1.1
## [40] fansi_0.4.2
                            crayon_1.5.2
                                                 withr 2.5.0
## [43] grid_4.0.5
                            gtable_0.3.1
                                                 lifecycle_1.0.0
## [46] DBI_1.1.3
                            magrittr_2.0.1
                                                 scales_1.1.1
## [49] cli_3.0.1
                            stringi_1.7.6
                                                 farver_2.1.0
## [52] latticeExtra_0.6-30 snakecase_0.11.0
                                                 ellipsis_0.3.2
                                                 Formula 1.2-4
## [55] generics 0.1.3
                            vctrs_0.3.8
## [58] RColorBrewer_1.1-3 tools_4.0.5
                                                 interp 1.0-33
## [61] glue_1.4.2
                            purrr 0.3.4
                                                 jpeg_0.1-9
## [64] fastmap 1.1.0
                            survival_3.2-10
                                                 yaml_2.3.5
## [67] colorspace_2.0-0
                            cluster_2.1.1
                                                 knitr_1.33
```