GY7702 Assignment

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The libraries that were used in this assignment:

```
library(tidyverse)
library(palmerpenguins)
library(dplyr)
library(knitr)
library(readr)
library(tinytex)
```

Question 1

A vector of 25 numbers between 1 and 7 are listed below. These values represent answers to a survey question

NA 3 4 4 5 2 4 NA 6 3 5 4 0 5 7 5 NA 5 2 4 NA 3 3 5 NA

- 1 = Completely disagree
- 2 = Disagree
- 3 = Somehow disagree
- 4 = Neither agree or disagree
- 5 = Somehow agree
- 6 = Agree
- 7 = Completely agree
- NA = missing value

Question 1.1

Write the code necessary to check whether all participants to the survey either completely disagree or completely agree, once the missing values are excluded

```
# First, let's create the vector.
num <- c(NA, 3, 4, 4, 5, 2, 4, NA, 6, 3, 5, 4, 0, 5, 7, 5, NA, 5, 2, 4, NA, 3, 3, 5, NA)
# Then, remove missing values.
num <- na.omit(num)
# And finally check whether all students either completely agree (= 7).
students_agree_complete <- all(num == 7)</pre>
```

```
# Or, completely disagree (=1).
students_disagree_complete <- all(num == 1)</pre>
```

Question 1.2

Write the code necessary to extract the indexes related to the participants in the survey who at least somehow agree or more.

In the vector *num*, which students selected values between 5 and 7.

```
students_agree <- which(num %in% c(5:7))
```

Question 2

Question 2.2

Write the code necessary to create a table showing species, island, bill length and body mass of the 10 Gentoo penguins in the penguins table with the highest body mass

The library palmerpenguins was created by Dr. Kristen Gorman and holds 2 data sets. For this assignment, we will work with penguins, a table containing 8 measured variables from 344 penguins!

Some of the penguins in *penguins* hold NA values for certain variables. Due to the analysis that we are going to do, these need to be removed

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

As this task requires multiple steps, it will be more efficient to start a pipe. Lets send the *na omitted* table, *pen* down the pipe.

It is important to note that the functions in the pipe operator do not require the first argument, *pen* because it is being 'fed down' the pipe. For more information on pipes, click here

```
pen %>%

# First, let's select all the variables that the question asks for
select(species, island, bill_length_mm, bill_depth_mm, body_mass_g)%>%

# and select all the 'Gentoo' penguins.
filter(species == "Gentoo")%>%

# Order body mass in ascending order
arrange(-body_mass_g)%>%

# Return the 10 heaviest Gentoo penguins
slice_head(n = 10)%>%
kable()
```

species	island	bill_length_mm	bill_depth_mm	body_mass_g
Gentoo	Biscoe	49.2	15.2	6300
Gentoo	Biscoe	59.6	17.0	6050
Gentoo	Biscoe	51.1	16.3	6000

	:-11	1.:11 1	1:11 141	1
species	island	bill_length_mm	biii_deptii_mm	body_mass_g
Gentoo	Biscoe	48.8	16.2	6000
Gentoo	Biscoe	45.2	16.4	5950
Gentoo	Biscoe	49.8	15.9	5950
Gentoo	Biscoe	48.4	14.6	5850
Gentoo	Biscoe	49.3	15.7	5850
Gentoo	Biscoe	55.1	16.0	5850
Gentoo	Biscoe	49.5	16.2	5800

Question 2.3

write the code necessary to create a table showing the average bill length per island, ordered by average bill length.

As this question states $per\ island$, we will use $group_by$ to perform an average bill length for each island. We will also see mean() to determine average. This is a function built into R.

After the data set is grouped by island, average bill length will be calculated. The result will be appended to pen in a column called avg_bill_length using the mutate function.

```
# Select the appropriate attributes
select(species, island, bill_length_mm, bill_depth_mm,)%>%

# Order by bill_length_mm. By default arrange sorts in ascending order.
arrange(bill_length_mm)%>%

# Use group_by to execute the average for each island
group_by(island)%>%

# Append the average bill_length_mm to the table pen
mutate(avg_bill_length_mm = mean(bill_length_mm))%>%

# Display 3 example average_bill_lengths for each island
slice_head(n = 3)%>%
kable()
```

species	island	$bill_length_mm$	$bill_depth_mm$	avg_bill_length_mm
Adelie	Biscoe	34.5	18.1	45.24847
Adelie	Biscoe	35.0	17.9	45.24847
Adelie	Biscoe	35.0	17.9	45.24847
Adelie	Dream	32.1	15.5	44.22195
Adelie	Dream	33.1	16.1	44.22195
Adelie	Dream	34.0	17.1	44.22195
Adelie	Torgersen	33.5	19.0	39.03830
Adelie	Torgersen	34.4	18.4	39.03830
Adelie	Torgersen	34.6	21.1	39.03830

Question 2.4

Write the code necessary to create a table showing the minimum, median and maximum proportion between bill length and bill depth by species.

The proportion between bill length and bill depth will be calculated as bill_length_mm / bill_depth_mm. This will return a unit-less measure which will be called bl_bd

R also contains in built functions for obtaining the minimum, median and maximum values from a vector. So by example, on line [] lets take the minimum $bill_length_mm$ and minimum $bill_depth_mm$ and divide them to get the unit-less measurement min_bl_bd .

```
# Select the necessary attributes
select(species, island, bill_length_mm, bill_depth_mm,)%>%

# Execute the proportions for each species
group_by(species)%>%

# Append the proportions to the table pen, using mutate
mutate(min_bl_bd = min(bill_length_mm) / min(bill_depth_mm))%>%
mutate(med_bl_bd = median(bill_length_mm) / median(bill_depth_mm))%>%
mutate(max_bl_bd = max(bill_length_mm) / max(bill_depth_mm))%>%

# Drop bill_length_mm and bill_depth_mm
subset(select = -c(bill_length_mm, bill_depth_mm))%>%

slice_head(n = 3)%>%
kable()
```

species	island	min_bl_bd	med_bl_bd	max_bl_bd
Adelie	Torgersen	2.070968	2.111413	2.139535
Adelie	Torgersen	2.070968	2.111413	2.139535
Adelie	Torgersen	2.070968	2.111413	2.139535
Chinstrap	Dream	2.493902	2.685637	2.788461
Chinstrap	Dream	2.493902	2.685637	2.788461
Chinstrap	Dream	2.493902	2.685637	2.788461
Gentoo	Biscoe	3.122137	3.160000	3.445087
Gentoo	Biscoe	3.122137	3.160000	3.445087
Gentoo	Biscoe	3.122137	3.160000	3.445087