GY7702: Coursework 1

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Introduction

This document has been created to increase the **reproducibility** of this coursework assignment, written in RMarkdown. To support the reproducibility of the document please refer to the *GitHub data repository* for the commits that document the development of this Coursework 1

Libraries

This coursework use the library tidyverse

```
library(tidyverse)
```

Also the library knitr

```
library(knitr)
```

Other libraries are also used for specific question for instance in question 2 the library **palmerpenguins** these specific libraries will be referred to within each question

Questions

Question 1: Question 1 deals with a vector of 25 numbers between 1 and 7, with each value representing answers to survey questions. Some values are missing. Vector was defined by the question:

```
# vector survey_responses contains 25 elements
survey_responses <- 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)
```

```
# specify the survey_response variable
survey_responses %>%
    # using na.omit removes NA values from the vector
na.omit() %>%
    # is. element will return logical output, if 7 or completely agree was answered in the
    # survey it will be returned as TRUE
    is.element(7) %>%
    # is. element will return logical output, if 1 or completely agree was answered in the
    # survey it will be returned as TRUE
    is.element(1)
```

Question 1.1:

```
## [1] FALSE FALSE
```

Question 1.2 Extract the indexes who at least somewhat agreed or more (values 5 to 7)

```
# Function which see if any value between 5 and 7 and return the index of each
# survey participant
which(survey_responses %in% 5:7)
```

```
## [1] 5 9 11 14 15 16 18 24
```

###Question 2: Question 2 looks data from Adélie, Chinstrap, and Gentoo penguins observed on islands in the Palmer Archipelago near Palmer Station, Antarctica. Palmerpenguins library can be found at Palmerpenguins GitHub Repositry

Question 2.1 Question 2.1 ask for the library (palmerpenguins) to be installed and loaded

```
library(tidyverse)
library(knitr)
# install.packages("palmerpenguins")
library(palmerpenguins)
```

Question 2.2 Question 2.2 creates a table showing *species, island, bill length and body mass* of the 10 Gentoo penguins in the penguins table with the highest body mass

```
# Starts from the entire palmerpenguins libraries
palmerpenguins::penguins %>%
    # Selects only the necessary columns
    dplyr::select(species, island, bill_length_mm, body_mass_g
) %>%
    # Retain only rows representing the Gentoo species
    dplyr::filter(species == "Gentoo"
) %>%
    # Sort by descending body mass in g
    dplyr::arrange(desc(body_mass_g))
```

```
## # A tibble: 124 x 4
##
      species island bill_length_mm body_mass_g
##
      <fct>
             <fct>
                              <dbl>
                                          <int>
## 1 Gentoo Biscoe
                               49.2
                                           6300
##
   2 Gentoo Biscoe
                               59.6
                                           6050
## 3 Gentoo Biscoe
                               51.1
                                           6000
## 4 Gentoo Biscoe
                               48.8
                                           6000
## 5 Gentoo
             Biscoe
                               45.2
                                           5950
##
                               49.8
                                           5950
  6 Gentoo Biscoe
## 7 Gentoo Biscoe
                               48.4
                                           5850
## 8 Gentoo Biscoe
                               49.3
                                           5850
## 9 Gentoo Biscoe
                               55.1
                                           5850
## 10 Gentoo Biscoe
                               49.5
                                           5800
## # ... with 114 more rows
```

Question 2.3 Question 2.3 creates a table with average bill length per island, ordered by average bill length

```
# Starts from the entire palmerpenguins libraries
palmerpenguins::penguins %>%
    # Selects only the necessary columns
    dplyr::select(bill_length_mm, island) %>%
    # Grouped by island
    dplyr::group_by(island) %>%
```

```
# Drops rows containing NAs in the bill_length_mm column
# otherwise the mean function will return NA
dplyr::filter(!is.na(bill_length_mm)) %>%
# Calculates the average of bill_length_mm
dplyr::summarise(average_bill_length = mean(bill_length_mm)) %>%
# Ordered by descending average_bill_length
dplyr::arrange(desc(average_bill_length)) %>%
# kable improves tibble format
knitr::kable()
```

average_bill_length
45.25749
44.16774
38.95098

Question 2.4 Question 2.4 creates a table showing the *minimum*, *median and maximum* proportion between *bill length and bill depth by species*

```
# Starts from the entire palmerpenguins libraries
palmerpenguins::penguins %>%
  # Selects only the necessary columns
  dplyr::select(species, bill_length_mm, bill_depth_mm) %>%
  # Grouped by species
  dplyr::group_by(species) %>%
  # Drops rows containing NAs in the bill_length_mm column
  # otherwise the mean function will return NA
  dplyr::filter(!is.na(bill_length_mm))%>%
  # Drops rows containing NAs in the bill_depth_mm column
  # otherwise the mean function will return NA
  dplyr::filter(!is.na(bill_depth_mm)) %>%
  # Calculates the bill length to bill depth ratio
  dplyr::summarise(Proportion=
                     (bill_length_mm/bill_depth_mm)) %>%
   # using summariase again the minimum, median and maximum for each species can be calculated
  dplyr::summarise(min(Proportion),
                   median(Proportion),
                   max(Proportion)) %>%
  # Using the function kable formats the table
  knitr::kable()
```

species	$\min(\text{Proportion})$	median(Proportion)	$\max(\text{Proportion})$
Adelie Chinstrap	$1.639810 \\ 2.350516$	$\begin{array}{c} 2.136842 \\ 2.661577 \end{array}$	2.450000 3.258427
Gentoo	2.566474	3.166667	3.612676

Question 3:

Question 3 looks at a topical data set of new and cumulative **COVD19** cases in the UK between March 1st and October 17th 2020. **COVID19** data is sourced from the HM Government Coronavirus in the UK

Question 3.1 Question 3.1 asks for the data covid19 cases to be loaded

```
#using readr (part of tidyverse)
library(readr)
# reads the .CSV file with the correct directory
# Imports covid19_cases_20200301_20201017.csv and assings to a new variable
#covid_data
covid_data <-readr::read_csv("covid19_cases_20200301_20201017.csv")</pre>
```

Question 3.2 This question asked for an area specific table to be generated. Here Brentwood (Essex) COVID19 is presented. Brentwood Borough Council (COVID19 response and information)[https://www.brentwood.gov.uk/index.php?cid=2937]

```
# create a complete table containing a row for each day and area, replace Na
# with the value available for the previous date
# Resulting table will be stored in the new variable
# brentwood_complete_covide_data
Brentwood_complete_covid_data <-covid_data %>%
  #selects extracts wanted columns
  dplyr::select(specimen_date, area_name, newCasesBySpecimenDate,
                cumCasesBySpecimenDate)%>%
  # group by specimen_data & area_name leads to each area_name having a row
  # per specimen data
  dplyr::group_by(specimen_date, area_name) %>%
  # tidyr :: fill replace NA values with the values from the previous row
  # default direction is down
  tidyr::fill(newCasesBySpecimenDate, cumCasesBySpecimenDate) %>%
  # replace_na replaces any remaining NA with 0
  tidyr::replace_na()%>%
  # dplyr :: filter subsets the area name to Bentwood
  dplyr::filter(area_name == "Brentwood")%>%
  # converting to a data frame as initially when trying to drop area_name using
  # dplyr::select (-area_number) got an error message 'adding missing grouping
  # variables error: area_number so I converted it to a data frame
  data.frame()%>%
  # then drop area_name using select
  dplyr::select(-area_name) %>%
  # Then converted back to a tibble
  as tibble()
```

Question 3.3

load library lubridate

```
library("lubridate")

# start from copy Brentwood_day_before
Brentwood_day_before <- Brentwood_complete_covid_data %>%

# mutate specimen_date using lubridate

# format of ymd is specified then - 1 day

# as.character converts date into character string

dplyr::mutate(day_before = as.character(ymd(specimen_date - 1 ))) %>%

# drop specimen_date and cumCasesBySpecimenDate columns

dplyr::select(-specimen_date, -cumCasesBySpecimenDate) %>%
```

```
# Rename newCasesBySpecimenDate to newCases_day_before
dplyr::rename(newCases_day_before = newCasesBySpecimenDate)
# join Brentwood_day_before with Brentwood_complete_covid_data
# where specimen_date is equal to day_before using inner_join
```

Question 3.4 ###Question 4

Importing population per local authority and COVID19 case data

```
# import lad19_population data and assigned to a new variable LAD_pop
LAD_pop<- readr::read_csv("lad19_population.csv")
# import COVID19 case data and assigned to a new variable LAD_covid_cases
LAD_covid_cases <- readr::read_csv("covid19_cases_20200301_20201017.csv")</pre>
```

Carrying out table join for LAD_pop and LAD_covid_cases

```
# need to rename LAD_pop column lad19_area_name to area_name to match
# LAD_covid_cases table
covid_cases_lad_pop <- dplyr::rename(LAD_pop,area_name = lad19_area_name) %>%
# table join between LAD_pop and LAD_covid_cases
dplyr::left_join(LAD_covid_cases, LAD_pop, by = "area_name")
```

###Description and Interpretation