# Homework 2

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Link to the Github repository

Due: Tue, Feb 14, 2023 @ 11:59pm

Please read the instructions carefully before submitting your assignment.

- 1. This assignment requires you to only upload a PDF file on Canvas
- 2. Don't collapse any code cells before submitting.
- 3. Remember to make sure all your code output is rendered properly before uploading your submission.

Please add your name to the author information in the frontmatter before submitting your assignment

For this assignment, we will be using the Abalone dataset from the UCI Machine Learning Repository. The dataset consists of physical measurements of abalone (a type of marine snail) and includes information on the age, sex, and size of the abalone.

We will be using the following libraries:

```
library(readr)
library(tidyr)
```

```
Warning: package 'tidyr' was built under R version 4.2.2

library(ggplot2)
library(dplyr)

Warning: package 'dplyr' was built under R version 4.2.2

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

library(purrr)

Warning: package 'purrr' was built under R version 4.2.2

library(cowplot)

Warning: package 'cowplot' was built under R version 4.2.2
```

## Question 1



EDA using readr, tidyr and ggplot2

### 1.1 (5 points)

Load the "Abalone" dataset as a tibble called abalone using the URL provided below. The abalone\_col\_names variable contains a vector of the column names for this dataset (to be consistent with the R naming pattern). Make sure you read the dataset with the provided column names.

```
library(readr)
  url <- "http://archive.ics.uci.edu/ml/machine-learning-databases/abalone/abalone.data"</pre>
  abalone_col_names <- c(
    "sex",
    "length",
    "diameter",
    "height",
    "whole_weight",
    "shucked_weight",
    "viscera_weight",
    "shell_weight",
    "rings"
  )
  abalone <- read_csv(url, col_names = abalone_col_names)</pre>
`curl` package not installed, falling back to using `url()`
Rows: 4177 Columns: 9
-- Column specification ------
Delimiter: ","
chr (1): sex
dbl (8): length, diameter, height, whole_weight, shucked_weight, viscera_wei...
i Use `spec()` to retrieve the full column specification for this data.
i Specify the column types or set `show_col_types = FALSE` to quiet this message.
```

### 1.2 (5 points)

Remove missing values and NAs from the dataset and store the cleaned data in a tibble called df. How many rows were dropped?

#### abalone

```
# A tibble: 4,177 x 9
```

	sex	length	${\tt diameter}$	height	whole_weight	shucked_wei~1	visce~2	$\verb shell-3 $	rings
	<chr></chr>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>
1	M	0.455	0.365	0.095	0.514	0.224	0.101	0.15	15
2	M	0.35	0.265	0.09	0.226	0.0995	0.0485	0.07	7

```
3 F
          0.53
                    0.42
                            0.135
                                          0.677
                                                        0.256
                                                                 0.142
                                                                           0.21
                                                                                      9
4 M
          0.44
                    0.365
                            0.125
                                                        0.216
                                                                 0.114
                                                                           0.155
                                          0.516
                                                                                     10
                                                                           0.055
5 I
          0.33
                    0.255
                            0.08
                                          0.205
                                                        0.0895
                                                                 0.0395
                                                                                      7
6 I
          0.425
                    0.3
                            0.095
                                          0.352
                                                        0.141
                                                                 0.0775
                                                                           0.12
                                                                                      8
7 F
          0.53
                                                        0.237
                                                                 0.142
                                                                           0.33
                    0.415
                           0.15
                                          0.778
                                                                                     20
8 F
          0.545
                    0.425
                            0.125
                                          0.768
                                                        0.294
                                                                 0.150
                                                                           0.26
                                                                                     16
9 M
          0.475
                    0.37
                            0.125
                                          0.509
                                                        0.216
                                                                 0.112
                                                                           0.165
                                                                                      9
                            0.15
10 F
          0.55
                    0.44
                                          0.894
                                                        0.314
                                                                 0.151
                                                                           0.32
                                                                                     19
```

# 2: viscera\_weight, 3: shell\_weight

```
df <- drop_na(abalone)
df</pre>
```

# A tibble: 4,177 x 9

	sex	length	${\tt diameter}$	height	whole_weight	shucked_wei~1	visce~2	shell~3	rings
	<chr></chr>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>
1	M	0.455	0.365	0.095	0.514	0.224	0.101	0.15	15
2	M	0.35	0.265	0.09	0.226	0.0995	0.0485	0.07	7
3	F	0.53	0.42	0.135	0.677	0.256	0.142	0.21	9
4	M	0.44	0.365	0.125	0.516	0.216	0.114	0.155	10
5	I	0.33	0.255	0.08	0.205	0.0895	0.0395	0.055	7
6	I	0.425	0.3	0.095	0.352	0.141	0.0775	0.12	8
7	F	0.53	0.415	0.15	0.778	0.237	0.142	0.33	20
8	F	0.545	0.425	0.125	0.768	0.294	0.150	0.26	16
9	M	0.475	0.37	0.125	0.509	0.216	0.112	0.165	9
10	F	0.55	0.44	0.15	0.894	0.314	0.151	0.32	19

<sup># ...</sup> with 4,167 more rows, and abbreviated variable names 1: shucked\_weight,

No rows were dropped from the dataset.

### 1.3 (5 points)

Plot histograms of all the quantitative variables in a **single plot** <sup>1</sup>

<sup># ...</sup> with 4,167 more rows, and abbreviated variable names 1: shucked\_weight,

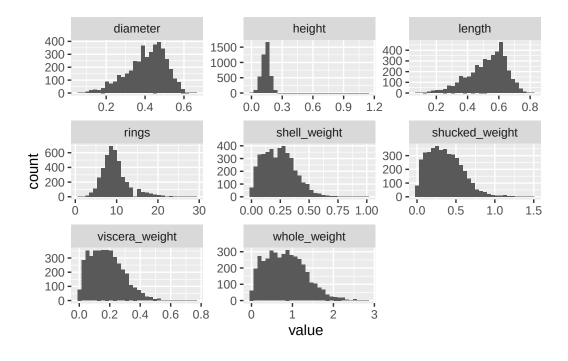
<sup># 2:</sup> viscera\_weight, 3: shell\_weight

 $<sup>^1</sup>$ You can use the facet\_wrap() function for this. Have a look at its documentation using the help console in R

```
new_df <-
    df %>%
    select(!sex) %>%
    # using gather function to gather all variables into one column
    gather(cols, value)

plt <-
    ggplot(new_df, aes(x = value)) +
    geom_histogram() +
    facet_wrap(~cols, scales="free")
plt</pre>
```

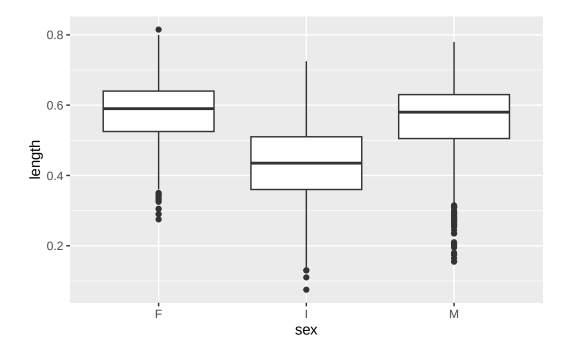
`stat\_bin()` using `bins = 30`. Pick better value with `binwidth`.



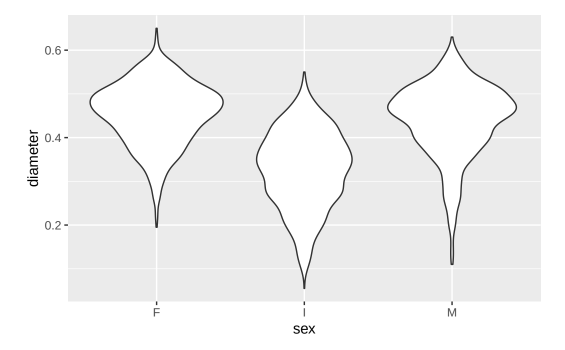
## 1.4 (5 points)

Create a boxplot of length for each sex and create a violin-plot of of diameter for each sex. Are there any notable differences in the physical appearences of abalones based on your analysis here?

```
box_plt <-
  ggplot(df, aes(x=sex, y=length)) +
  geom_boxplot()
box_plt</pre>
```



```
violin_plt <-
  ggplot(df, aes(x=sex, y=diameter)) +
  geom_violin()
violin_plt</pre>
```

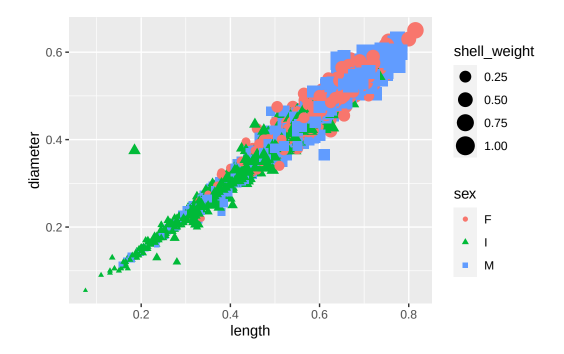


It would seem that the abalones of 'sex' "I" are significantly shorter in both 'diameter' and 'length' than those of 'sex' "M" and "F". In addition to this, the abalones of 'sex' "F" generally have a larger diameter than the other sexes.

## 1.5 (5 points)

Create a scatter plot of length and diameter, and modify the shape and color of the points based on the sex variable. Change the size of each point based on the shell\_wight value for each observation. Are there any notable anomalies in the dataset?

```
scatter_plt <-
ggplot(df, aes(x=length, y=diameter)) +
geom_point(aes(shape=sex, color=sex, size=shell_weight))
scatter_plt</pre>
```



The plot created above shows that as 'length' increases so too does 'diameter'. When it comes to any notable anomalies that are displayed by the plot, a few come to mind. For example, there is an outlier abalone of 'sex' "I" represented by a triangle that is far larger in 'diameter' than other abalones of proportional 'length'. In addition to this, it has become evident that abalones of 'sex' "M" have a much more diverse size range than those of 'sex' "F". In other words, male abalones can be much smaller than female abalones while still growing relatively large in line with the average size of females.

# 1.6 (5 points)

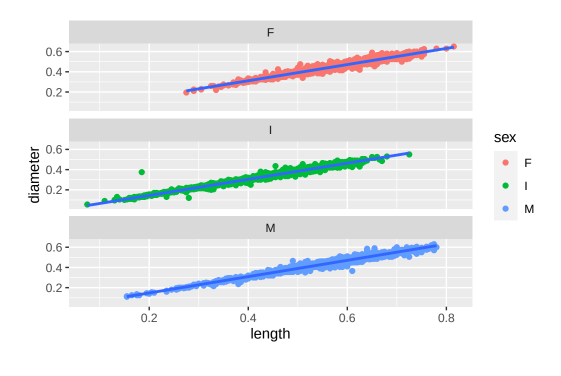
For each sex, create separate scatter plots of length and diameter. For each plot, also add a linear trendline to illustrate the relationship between the variables. Use the facet\_wrap() function in R for this, and ensure that the plots are vertically stacked **not** horizontally. You should end up with a plot that looks like this: <sup>2</sup>

```
abalone_plt <-
    ggplot(df, aes(x=length, y=diameter)) +
    geom_point(aes(color=sex)) +
    geom_smooth(method = "lm") +</pre>
```

 $<sup>^2</sup>$ Plot example for 1.6

```
facet_wrap(vars(sex), dir="v")
abalone_plt
```

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



# Question 2

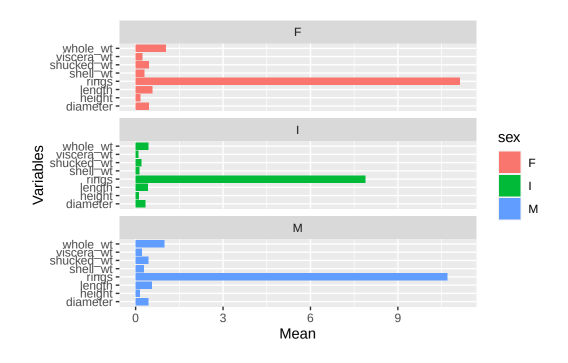


More advanced analyses using dplyr, purrrr and ggplot2

## 2.1 (10 points)

Filter the data to only include abalone with a length of at least 0.5 meters. Group the data by sex and calculate the mean of each variable for each group. Create a bar plot to visualize the mean values for each variable by sex.

```
glyph_ready_df <-</pre>
  df %>%
  group_by(sex) %>%
  summarize(across(everything(),mean)) %>%
  filter("length" >= 0.5) %>%
  # renaming just to make cleaner
  rename("shell_wt" = shell_weight,
         "shucked_wt" = shucked_weight,
         "viscera_wt" = viscera_weight,
         "whole_wt" = whole_weight) %>%
  # using pivot longer to put all variables into one column
  pivot_longer(!sex,
               names_to = "Variables",
               values_to = "Mean")
ggplot(glyph_ready_df, aes(x=Mean, y=Variables)) +
  geom_bar(stat="identity", aes(fill=sex)) +
  facet_wrap(vars(sex), dir="v")
```

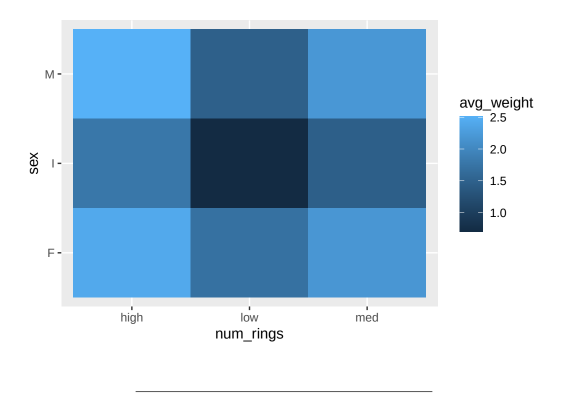


2.2 (15 points)

Implement the following in a **single command**:

- 1. Temporarily create a new variable called num\_rings which takes a value of:
- "low" if rings < 10
- "high" if rings > 20, and
- "med" otherwise
- Group df by this new variable and sex and compute avg\_weight as the average of the whole\_weight + shucked\_weight + viscera\_weight + shell\_weight for each combination of num\_rings and sex.
- 3. Use the geom\_tile() function to create a tile plot of num\_rings vs sex with the color indicating of each tile indicating the avg\_weight value.

`summarise()` has grouped output by 'num\_rings'. You can override using the `.groups` argument.



# 2.3 (5 points)

Make a table of the pairwise correlations between all the numeric variables rounded to 2 decimal points. Your final answer should look like this  $^3$ 

```
# creating new data frames with only numeric variables
df2 <- df %>%
    select(!sex)

# assigning variable of pairwise correlations
pair.cor <- round(cor(df2), 2)
pair.cor</pre>
```

	length	diameter	height	whole_weight	shucked_weight
length	1.00	0.99	0.83	0.93	0.90
diameter	0.99	1.00	0.83	0.93	0.89
height	0.83	0.83	1.00	0.82	0.77
whole_weight	0.93	0.93	0.82	1.00	0.97

 $<sup>^3</sup>$ Table for 2.3

shucked_weight	0.90	0.89	0.77	0.97	1.00
viscera_weight	0.90	0.90	0.80	0.97	0.93
shell_weight	0.90	0.91	0.82	0.96	0.88
rings	0.56	0.57	0.56	0.54	0.42
	viscera_	weight sh	ell_weight	rings	
length		0.90	0.90	0.56	
diameter		0.90	0.91	0.57	
height		0.80	0.82	0.56	
whole_weight		0.97	0.96	0.54	
shucked_weight		0.93	0.88	0.42	
viscera_weight		1.00	0.91	0.50	
shell_weight		0.91	1.00	0.63	
rings		0.50	0.63	1.00	

### 2.4 (10 points)

Use the map2() function from the purr package to create a scatter plot for each quantitative variable against the number of rings variable. Color the points based on the sex of each abalone. You can use the cowplot::plot\_grid() function to finally make the following grid of plots.

```
# creating data frame with all quantitative variables

df_quant <-
    df %>%
    select(!sex)

# creating data frame with only rings variable

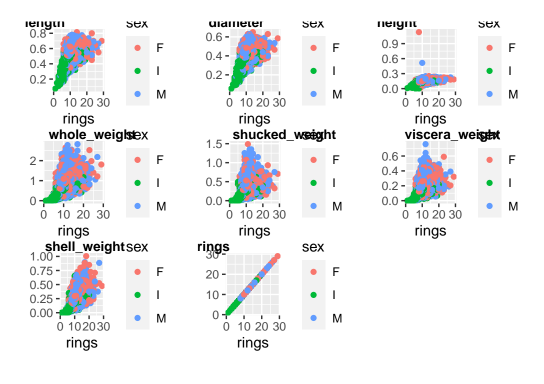
df_y <-
    df %>%
    select(rings)

# using 'map2()' function to make a plot of every quantitative variable vs rings

plot1 <- map2(df_quant, df_y, ~ ggplot(df) +
    geom_point(aes(x=rings, y=.x, color=sex)) +
    labs(x = "rings", y = " "))

# using 'plot_grid()' to put all plots into one grid

cowplot::plot_grid(plotlist = plot1, labels = colnames(df_quant), ncol = 3, label_size = 9</pre>
```



# Question 3



## 3.1 (10 points)

Perform a simple linear regression with diameter as the covariate and height as the response. Interpret the model coefficients and their significance values.

```
# assigning 'x' and 'y'
x = df$diameter
y = df$height

# creating the model and summarizing
model <- lm(y ~ x)</pre>
```

#### summary(model)

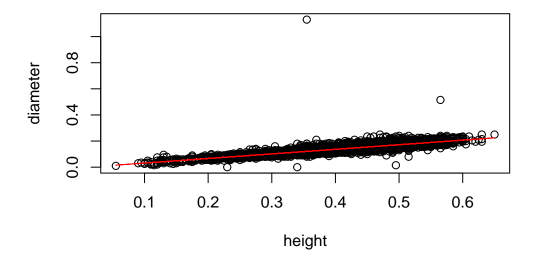
```
Call:
lm(formula = y \sim x)
Residuals:
     Min
               1Q
                    Median
                                 3Q
                                          Max
-0.15513 -0.01053 -0.00147 0.00852 1.00906
Coefficients:
             Estimate Std. Error t value Pr(>|t|)
                                  -2.515
(Intercept) -0.003803
                        0.001512
                                            0.0119 *
             0.351376
                        0.003602 97.544
                                            <2e-16 ***
X
                0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Signif. codes:
Residual standard error: 0.0231 on 4175 degrees of freedom
Multiple R-squared: 0.695, Adjusted R-squared:
F-statistic: 9515 on 1 and 4175 DF, p-value: < 2.2e-16
```

The model's  $\beta_0$  coefficient represents the intercept and has a value of -0.003803. This means that the y-intercept of the simple linear regression model is in the negatives. The model's  $\beta_1$  coefficient represents the slope and has a value of 0.351376. The p-value is very statistically significant therefore we reject the null hypothesis and accept the alternative hypothesis. The R-squared value, on the other hand, is not very large and is not significant.

### 3.2 (10 points)

Make a scatterplot of height vs diameter and plot the regression line in color="red". You can use the base plot() function in R for this. Is the linear model an appropriate fit for this relationship? Explain.

```
# creating scatterplot using 'plot()' function
plot(x, y, xlab="height", ylab="diameter")
lines(x, fitted(lm(y ~ x)), col="red")
```



Yes this linear model is an appropriate fit for this relationship because it successfully minimizes the distance from each point to the trend line. In addition to this, the p-value is statistically significant, as mentioned above.

## 3.3 (10 points)

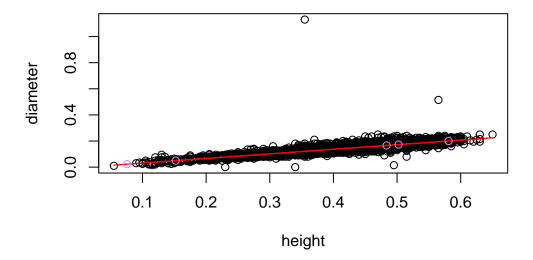
Suppose we have collected observations for "new" abalones with new\_diameter values given below. What is the expected value of their height based on your model above? Plot these new observations along with your predictions in your plot from earlier using color="violet"

```
new_diameters <- c(
    0.15218946,
    0.48361548,
    0.58095513,
    0.07603687,
    0.50234599,
    0.83462092,
    0.95681938,
    0.92906875,
    0.94245437,
```

```
0.01209518
)

# creating new variables
new_x <- data.frame(x = new_diameters)
new_y <- predict(model, new_x)

# plotting graph and adding predicted points
plot(x, y, xlab="height", ylab="diameter")
lines(x, fitted(lm(y ~ x)), col="red")
points(new_x %>% unlist(), new_y, col="violet")
```



The expected value of the "new" abalones height based on the model above is as follows:

```
new_y
```

1 2 3 4 5 6 0.0496723682 0.1661276096 0.2003304536 0.0229141546 0.1727090665 0.2894625947 7 8 9 10 0.3324002348 0.3226493217 0.3273527111 0.0004465615

# **Appendix**

```
i Session Information
Print your R session information using the following command
  sessionInfo()
R version 4.2.1 (2022-06-23 ucrt)
Platform: x86_64-w64-mingw32/x64 (64-bit)
Running under: Windows 10 x64 (build 22000)
Matrix products: default
locale:
[1] LC_COLLATE=English_United States.utf8
[2] LC_CTYPE=English_United States.utf8
[3] LC_MONETARY=English_United States.utf8
[4] LC_NUMERIC=C
[5] LC_TIME=English_United States.utf8
attached base packages:
[1] stats
              graphics grDevices datasets utils
                                                       methods
                                                                 base
other attached packages:
[1] cowplot_1.1.1 purrr_1.0.1
                                dplyr_1.1.0
                                               ggplot2_3.4.1 tidyr_1.3.0
[6] readr_2.1.4
loaded via a namespace (and not attached):
 [1] pillar_1.8.1
                      compiler_4.2.1
                                       tools_4.2.1
                                                         bit_4.0.5
 [5] digest_0.6.31
                      lattice_0.20-45 nlme_3.1-157
                                                         jsonlite_1.8.4
 [9] evaluate_0.20
                      lifecycle_1.0.3 tibble_3.1.8
                                                         gtable_0.3.1
[13] mgcv_1.8-40
                      pkgconfig_2.0.3 rlang_1.0.6
                                                         Matrix_1.4-1
[17] cli_3.6.0
                      parallel_4.2.1
                                       yaml_2.3.7
                                                         xfun_0.37
[21] fastmap_1.1.0
                      withr_2.5.0
                                       knitr_1.42
                                                         generics_0.1.3
[25] vctrs_0.5.2
                      hms_1.1.2
                                       bit64_4.0.5
                                                         grid_4.2.1
[29] tidyselect_1.2.0 glue_1.6.2
                                       R6_2.5.1
                                                         fansi_1.0.4
[33] vroom_1.6.1
                      rmarkdown_2.20
                                       farver_2.1.1
                                                         tzdb_0.3.0
```

```
[37] magrittr_2.0.3 splines_4.2.1 scales_1.2.1 ellipsis_0.3.2 [41] htmltools_0.5.4 colorspace_2.1-0 renv_0.16.0-53 labeling_0.4.2 [45] utf8_1.2.3 munsell_0.5.0 crayon_1.5.2
```