STAT8111 Assigment1

Phan Vinh Phu 45747989

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Question 1

pemax

1.0000

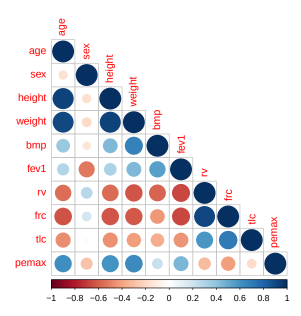
a.Examine first graphically and numerically correlation between variables, then comment:

Numerical correlation

```
num_cor <- round(cor(data), 4)</pre>
num_cor
##
                   sex height weight
                                         bmp
                                                fev1
                                                               frc
                                                                      tlc
            age
## age
         1.0000 -0.1671
                       0.9261 0.9059 0.3778
                                             0.2945 -0.5519 -0.6394 -0.4694
## sex
         -0.1671 1.0000 -0.1675 -0.1904 -0.1376 -0.5283 0.2714
                                                            0.1836
                       1.0000 0.9207
## height 0.9261 -0.1675
                                      0.4408
                                             0.3167 -0.5695 -0.6243 -0.4571
## weight 0.9059 -0.1904
                        0.9207
                               1.0000
                                      0.6725
                                             0.4488 -0.6215 -0.6173 -0.4185
         0.3778 -0.1376
                        0.4408
                               0.6725
                                      1.0000
                                             0.5455 -0.5824 -0.4344 -0.3649
## bmp
## fev1
         0.5455
                                              1.0000 -0.6659 -0.6651 -0.4430
## rv
        -0.5519 0.2714 -0.5695 -0.6215 -0.5824 -0.6659
                                                    1.0000 0.9106 0.5891
## frc
        -0.6394   0.1836   -0.6243   -0.6173   -0.4344   -0.6651
                                                    0.9106
                                                            1.0000
        -0.4694 0.0242 -0.4571 -0.4185 -0.3649 -0.4430
## tlc
                                                    0.5891
                                                            0.7044 1.0000
         ## pemax
##
          pemax
## age
         0.6135
         -0.2886
## sex
## height 0.5992
## weight 0.6352
## bmp
         0.2295
## fev1
         0.4534
## rv
        -0.3156
## frc
        -0.4172
## tlc
        -0.1816
```

Graphical correlation

```
corrplot(num_cor, type="lower")
```



Comment:

From the correlation plot, it can be seen that :

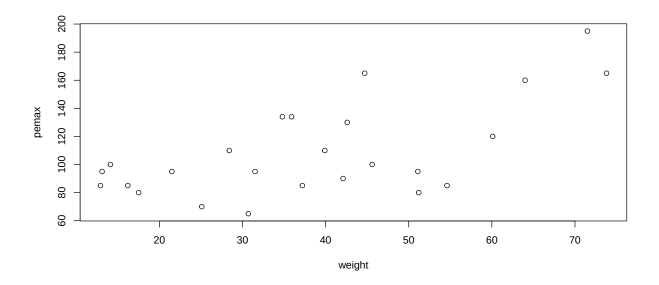
- There might be a strong positive linear relationship between age and height, weight and pemax. Beside that, age is negatively correlated with frc.
- height and weight are strongly correlated. In addition, weight has moderate negative correlation with rv, frc, tlc and positive correlation with bmp, pemax.
- Similarly, fev1 has moderate negative correlation with rv, frc, tlc.
- Lastly, rv highly positively correlated with frc.

b. the relationship between weight and pemax

In this part, the relationship between weight and pemax will be examined. Specifically, pemax is the dependent variable (Y) and weight is the independent variable (X).

Scatter Plot

```
plot(x = data$weight, y = data$pemax, xlab="weight", ylab = "pemax")
```



The graph illustrates that weight is positively related with pemax. The trend is that when weight increase, pemax will increase.

Linear Model

```
model <- lm(pemax ~ weight, data = data)</pre>
summary(model)
##
## Call:
## lm(formula = pemax ~ weight, data = data)
##
## Residuals:
##
      Min
              1Q Median
                            3Q
                                  Max
##
   -44.30 -22.69
                   2.23
                         15.91
                                48.41
##
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) 63.5456
                           12.7016
                                     5.003 4.63e-05 ***
## weight
                 1.1867
                            0.3009
                                     3.944 0.000646 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 26.38 on 23 degrees of freedom
## Multiple R-squared: 0.4035, Adjusted R-squared: 0.3776
## F-statistic: 15.56 on 1 and 23 DF, p-value: 0.0006457
```

The model equation $pe\hat{m}ax = 63.5456 + 1.1867weight + \epsilon \ (\epsilon \sim N(0, \sigma^2))$

Model Fit

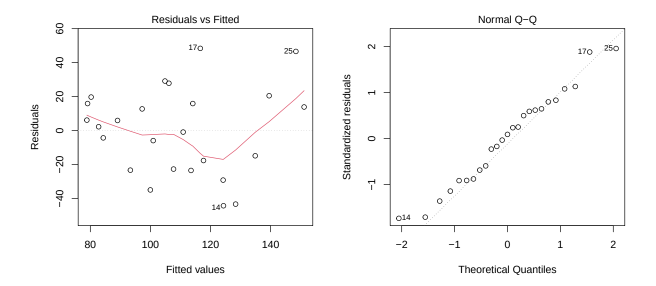
• The $R^2 = 0.4035$ which means 40.35% of variation in pemax can be explained by weight. This show that the model is not fit well.

Model interpretation

- According to the equation, for each unit increase in weight, the pemax will increase about 1.1867.
- weight is significant predictor since the p-value = 0.000646 (< 0.001).

```
par(mfrow = c(1,2))
plot(model, which = c(1,2))
```

Diagnostic Checking



- The standardized Residuals versus Fitted values plot appears to be a random scatter about zero, so the model is adequate. This graph also show some residuals which are 14, 17, 25 are low and high. This can be evidence of small amount of heteroscedasticity.
- the Normal Q-Q plot is approximately linear, so it can be said that the normality assumption holds.

c. Include in the previous model the sex variable

Model 2

```
model_2 <- lm(pemax ~ weight + sex, data = data)</pre>
summary(model_2)
##
## Call:
## lm(formula = pemax ~ weight + sex, data = data)
## Residuals:
##
       Min
                1Q Median
                                3Q
                                        Max
## -47.388 -16.850
                    0.073 13.168 43.748
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 70.9719
                           14.4644
                                     4.907 6.61e-05 ***
                 1.1248
                            0.3056
                                     3.681 0.00131 **
## weight
               -11.4776
                           10.7963 -1.063 0.29926
## sex
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 26.31 on 22 degrees of freedom
## Multiple R-squared: 0.4327, Adjusted R-squared: 0.3811
## F-statistic: 8.388 on 2 and 22 DF, p-value: 0.00196
The model 2 equation
pe\hat{m}ax = 70.9719 + 1.1248weight + (-11.4776)sex + \epsilon \quad (\epsilon \sim N(0, \sigma^2))
Model 3
model_3 <- lm(pemax ~ sex + weight, data = data)</pre>
summary(model 3)
##
## Call:
## lm(formula = pemax ~ sex + weight, data = data)
## Residuals:
##
       Min
                1Q Median
                                3Q
                                        Max
## -47.388 -16.850
                    0.073 13.168 43.748
##
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 70.9719
                           14.4644
                                     4.907 6.61e-05 ***
               -11.4776
                           10.7963
                                    -1.063 0.29926
## sex
                            0.3056
                                     3.681 0.00131 **
## weight
                 1.1248
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 26.31 on 22 degrees of freedom
## Multiple R-squared: 0.4327, Adjusted R-squared: 0.3811
## F-statistic: 8.388 on 2 and 22 DF, p-value: 0.00196
```

The model 3 equation

```
pe\hat{m}ax = 70.9719 + (-11.4776)sex + 1.1248weight + \epsilon \quad (\epsilon \sim N(0, \sigma^2))
```

Analyse the two proposed models

- Model_2 and model_3 are seem to be similar. The only difference is the order of weight and sex. The R^2 of both model are 0.4327 which mean 43.27% of variant in pemax can be explained by weight and sex. On the one hand, in both two model, sex is insignificant predictor with p-value = 0.29926. On the other hand, weight is still significant predictor.
- For one unit increase in weight, pemax will increase 1.1248. The coefficient of sex represents the difference in pemax between females and males, while weight is same. In this case, models indicate that, on average, females have a pemax that is lower by 11.4776 units compared to males.

In conclusion, it appears that the order of variables (weight and sex) doesn't affect the results in this case. Model 2 and model 3 are better than the first model on question (b) but these two still are not good model.

Choose one model

In comparison of three models, model 2 and model 3 are appear to explain pemax better with higher R^2 and adjusted R^2 . Beside that, there are no collinearity issue between weight and sex. Therefore, I choose model 2.

d. Construct a statistical model for the response variables pemax based on the normal response distribution and the weight, bmp, fev1, rv,frc.

We are interested in predicting pemax (maximal expiratory pressure) by using weight, bmp, fev1, rv,frc. In this case, the stepwise backward selection will be used to find the best model. In stepwise backward selection, people first regress with all predictor variables in the model. Then, the predictors with the largest p-value in the t-test will be dropped. Next step is fitting the reduced model. This progess will be run iteratively until all variables in the model are significant.Let's start with the full model.

Full Model

```
full_model <- lm(pemax ~ weight + bmp + fev1 + rv + frc , data = data)
summary(full_model)</pre>
```

```
##
## Call:
## lm(formula = pemax ~ weight + bmp + fev1 + rv + frc, data = data)
## Residuals:
##
              10 Median
      Min
                             3Q
                                   Max
## -39.72 -12.17
                   4.83
                        15.29
                                 34.75
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 64.18640
                          54.73550
                                      1.173 0.255423
## weight
                           0.42529
                                      4.081 0.000637 ***
                1.73556
```

```
-1.35105
                          0.66763 -2.024 0.057303 .
## bmp
## fev1
                                    2.432 0.025078 *
               1.53087
                          0.62948
               0.13612
## rv
                          0.15668
                                    0.869 0.395787
                          0.31278 -0.079 0.937703
              -0.02477
## frc
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 23.34 on 19 degrees of freedom
## Multiple R-squared: 0.6142, Adjusted R-squared: 0.5127
## F-statistic: 6.05 on 5 and 19 DF, p-value: 0.001637
```

It can be seen that frc has the largest p-value which is 0.937703. Therefore, frc explains the least variation when added to the model. Now, frc will be dropped.

The reduced model without frc

```
reduced_model <- lm(pemax ~ weight + bmp + fev1 + rv , data = data)
summary(reduced_model)</pre>
```

```
##
## Call:
## lm(formula = pemax ~ weight + bmp + fev1 + rv, data = data)
## Residuals:
     Min
             1Q Median
                           3Q
                                 Max
## -39.77 -11.74
                 4.33 15.66
                               35.07
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 63.94669
                         53.27673
                                    1.200 0.244057
               1.74891
## weight
                          0.38063
                                    4.595 0.000175 ***
## bmp
              -1.37724
                          0.56534 -2.436 0.024322 *
                                    2.679 0.014410 *
## fev1
               1.54770
                          0.57761
               0.12572
                          0.08315
                                    1.512 0.146178
## rv
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 22.75 on 20 degrees of freedom
## Multiple R-squared: 0.6141, Adjusted R-squared: 0.5369
## F-statistic: 7.957 on 4 and 20 DF, p-value: 0.000523
```

Similarly, rv is the insignificant predictor and has the largest P-value which is 0.146178. Hence, I drop rv and fit the reduced model without frc and rv.

The reduced model without frc and rv

```
reduced_model_2 <- lm(pemax ~ weight + bmp + fev1 , data = data)
summary(reduced_model_2)</pre>
```

```
##
## Call:
## lm(formula = pemax ~ weight + bmp + fev1, data = data)
## Residuals:
##
      Min
               1Q Median
                               3Q
                                      Max
## -42.388 -13.496
                   3.991 14.856 40.373
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) 126.3336
                          34.7199
                                   3.639 0.001536 **
                                   4.216 0.000387 ***
                1.5365
                           0.3644
## weight
                           0.5793 -2.530 0.019486 *
## bmp
               -1.4654
                1.1086
                           0.5144
                                  2.155 0.042893 *
## fev1
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 23.44 on 21 degrees of freedom
## Multiple R-squared: 0.57, Adjusted R-squared: 0.5086
## F-statistic: 9.279 on 3 and 21 DF, p-value: 0.000418
```

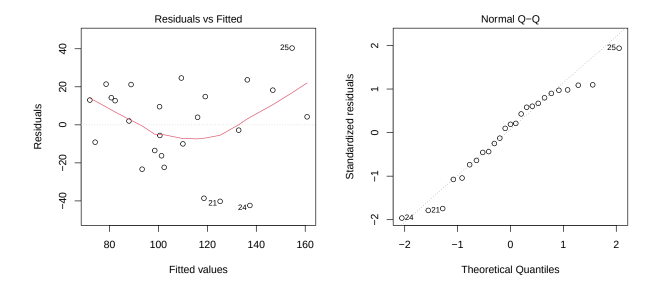
At this stage, all predictors are significant, therefore, selection process stops here. Nevertheless, there is moderate multi-collinearity since bmp, weight, fev1 are positively correlated.

```
# Calculate the correlation matrix
correlation_matrix <- cor(data[, c("weight", "bmp", "fev1")])
# Print the correlation matrix
print(correlation_matrix)</pre>
```

```
## weight bmp fev1
## weight 1.0000000 0.6725463 0.4488393
## bmp 0.6725463 1.0000000 0.5455204
## fev1 0.4488393 0.5455204 1.0000000
```

Diagnostic Checking

```
par(mfrow = c(1,2))
plot(reduced_model_2, which = 1:2)
```



- The Residuals vs Fitted plot look like random scatter around 0. Therefore, there is no obvious pattern in any of the residual plots so it appears the linearity and constant variance assumptions of the multiple linear model are justified.
- The quantile plot of residuals look approximately linear, suggesting the normality assumption for residuals is appropriate.

The final model equation

$$pe\hat{m}ax = 126.3336 + 1.5365weight + (-1.4654)bmp + 1.1086fev1 + \epsilon \quad (\epsilon \sim N(0, \sigma^2))$$

Interpretaion

From above equation,

- For every unit increase in weight (bmp and fev1 stay unchanged), the average pemax is expected to increase by 1.5365 units.
- For every unit increase in bmp (holding weight and fev1 constant), the average pemax is expected to decrease by 1.4654 units.
- For one extra unit of fev1, the average pemax is expected to increase by 1.1086 units.
- The R-squared value of 0.57 indicates that approximately 57% of the variability in the pemax can be explained by the predictor variables (weight, bmp, and fev1) included in the model.

Question 2

a. Show that the Inverse Gaussian distribution is a member of the exponential family

The Inverse Gaussian distribution is defined as

$$f(x;\mu,\gamma) = \sqrt{\frac{\gamma}{2\pi x^3}} \exp\left(-\frac{\gamma(x-\mu)^2}{2\mu^2 x}\right)$$

where

$$\mu > 0$$
 and $\gamma > 0$

$$\begin{split} f(x;\mu,\gamma) &= \exp\left(\frac{1}{2}log(\frac{\gamma}{2\pi x^3}) - \frac{\gamma}{2\mu^2}\frac{(x-\mu)^2}{x}\right) \\ f(x;\mu,\gamma) &= \exp\left(\frac{1}{2}log(\frac{\gamma}{2\pi x^3}) - \frac{\gamma}{2\mu^2}\frac{(x^2-2x\mu+\mu^2)}{x}\right) \\ f(x;\mu,\gamma) &= \exp\left(\frac{1}{2}log(\frac{\gamma}{2\pi x^3}) - \frac{\gamma x}{2\mu^2} + \frac{x}{\mu} + \frac{\gamma}{2x}\right) \\ f(x;\mu,\gamma) &= \exp\left(-\frac{\gamma x}{2\mu^2} + \frac{x}{\mu} + \frac{1}{2}log(\frac{\gamma}{2\pi x^3}) + \frac{\gamma}{2x}\right) \end{split}$$

The simpler form of the pdf:

$$f(y;\theta,\phi) = \exp\left(\frac{y\theta - b(\theta)}{a(\phi)} + c(y,\phi)\right)$$

in this case :

$$\frac{y\theta-b(\theta)}{a(\phi)}+c(y,\phi)=-\frac{\gamma x}{2\mu^2}+\frac{x}{\mu}+\frac{1}{2}log(\frac{\gamma}{2\pi x^3})+\frac{\gamma}{2x}$$

with:

$$\phi = \gamma, \ a = \frac{1}{\phi}, \ \theta = -\frac{1}{2\mu^2}, \ b = -\sqrt{-2\theta}, c = \frac{1}{2}log(\frac{\theta}{2\pi x^3}) + \frac{\theta}{2x}$$

.