TMA4268 - Project 1

```
# Libraries used throughout the exercise
library(ggplot2)
library(palmerpenguins) # Contains the data set 'penguins'.
library(ggfortify)
library(tidyverse)
library(GGally)
library(MASS)
library(class)
library(base)
library(ROSE)
```

Problem 1

a)

If we have that $y_0 = f(x_0) + \epsilon$, where ϵ is a random error independent of x with 0 mean. And x_0 is a unseen test observation we find,

$$E\left[\left(y_{0} - \hat{f}(x_{0})\right)^{2}\right] = E\left[\left(f(x_{0}) - \hat{f}(x_{0})\right)^{2}\right] + E\left[\epsilon^{2}\right] + 2E\left[\left(f(x_{0}) - \hat{f}(x_{0})\right) \cdot \epsilon\right]$$

$$= E\left[f(x_{0})^{2} - 2f(x_{0})\hat{f}(x_{0}) + \hat{f}(x_{0})^{2}\right] + E\left[\epsilon^{2}\right] + 2E\left[\left(f(x_{0}) - \hat{f}(x_{0})\right)\right] \cdot E[\epsilon]$$

$$= f(x_{0})^{2} - 2f(x_{0})E[\hat{f}(x_{0})] + E[\hat{f}(x_{0})^{2}] + Var\left[\epsilon\right]$$

$$= \left(f(x_{0}) - E[\hat{f}(x_{0})]\right)^{2} + Var[\hat{f}(x_{0})] + Var\left[\epsilon\right]$$

$$= \text{Bias}^{2} + \text{Variance} + \text{Irreducible error}$$

b)

The irreducible error is a term that can't be reduced by fitting the data well. The variance is a term describing how much uncertainty our statistical model \hat{f} , how much \hat{f} changes if we change the training data. The bias is the expected difference between the true model f and our statistical model \hat{f} .

c)

- i) True
- ii) False
- iii) True
- iv) False

d)

- i) False
- ii) False
- iii) True
- iv) False

e)

(iii) 0.76

$$\rho_{x_1, x_2} = \frac{\text{cov}(x_1, x_2)}{\sigma_{x_1} \sigma_{x_2}} = \frac{33}{\sqrt{50}\sqrt{38}} = 0.76$$

Problem 2

a)

One error made is excluding sex because of a low p-value, the p-value is the probability to observe a result equal or more extreme than the one we did, given that the null hypothesis $H_0: \beta = 0$, so a lower p-value means the result is more statistically significant.

Another error made is deciding whether to reject the null hypothesis that the species coefficient overall is actually zero based on the p-values of the coefficients. Since the null hypothesis in this case is H_0 : $\beta_{Chinstrap} = \beta_{Gentoo} = 0$ at the same time. We need to do a F-test.

A third error made is claiming that chinstrap penguins are the largest since they have the largest coefficient. Having a large coefficient only means that for a given set of parameters a chinstrap penguin is going to be heavier than another type of penguin. And when we include interaction terms which changes the the slopes, a large coefficient only guarantees it is larger for a range of parameters near zero. And by looking at body mass of the penguins in the data set we find that in fact gentoo penguins have the highest mean body mass.

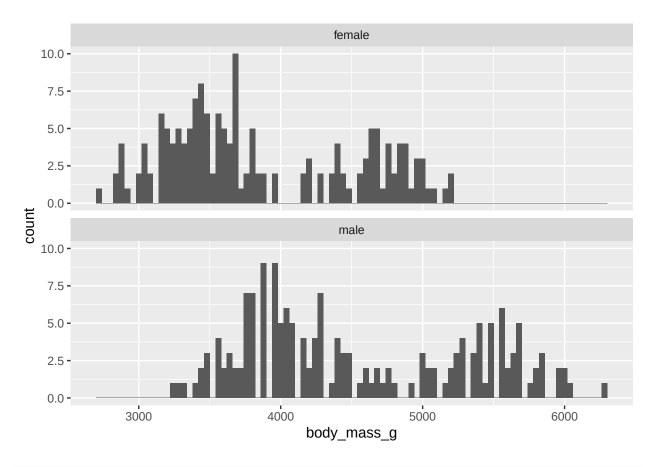
b)

```
data(penguins)

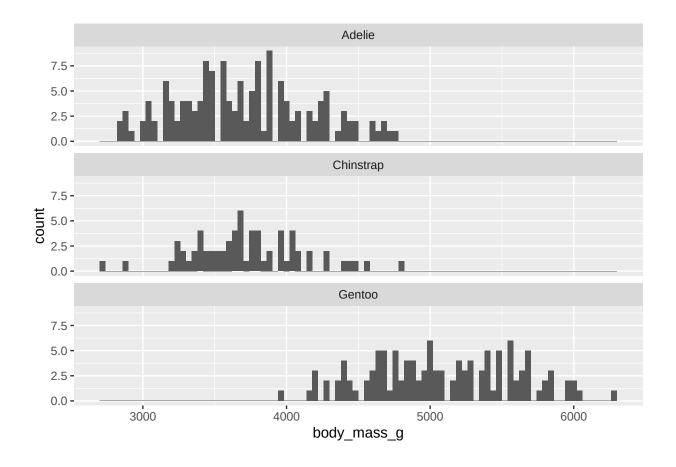
Penguins <- subset(penguins, select = -c(island, year))

Penguins <- na.omit(Penguins)

ggplot(data = Penguins) +
   geom_histogram(binwidth = 40, mapping = aes(x = body_mass_g)) +
   facet_wrap(~ sex, nrow = 2)</pre>
```



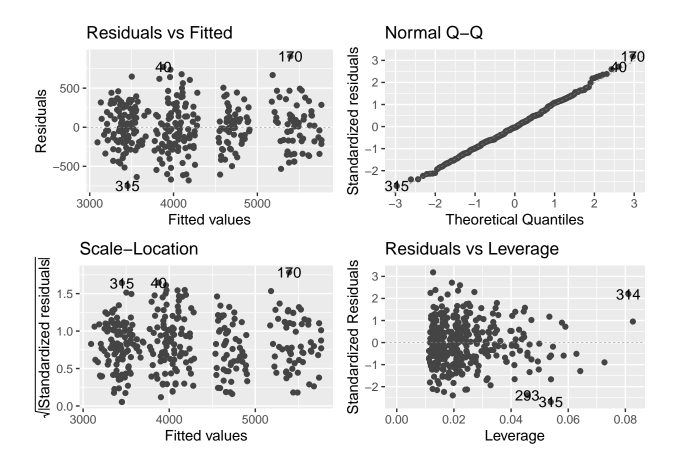
```
ggplot(data = Penguins) +
geom_histogram(binwidth = 40, mapping = aes(x = body_mass_g)) +
facet_wrap(~ species, nrow = 3)
```



c)

```
Penguins <- subset(penguins, select = -c(island, year))</pre>
penguin.model <- lm(body_mass_g ~ flipper_length_mm + sex + bill_depth_mm * species,</pre>
                    data = Penguins)
summary(penguin.model)
##
## Call:
## lm(formula = body_mass_g ~ flipper_length_mm + sex + bill_depth_mm *
##
       species, data = Penguins)
##
## Residuals:
##
      Min
              1Q Median
                                  Max
## -751.2 -183.8
                  -9.8 191.1 906.9
##
## Coefficients:
                                  Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                                                646.92 -2.066 0.039615 *
                                  -1336.58
                                                         5.971 6.17e-09 ***
## flipper_length_mm
                                      17.38
                                                  2.91
                                                       9.699 < 2e-16 ***
## sexmale
                                    432.90
                                                 44.63
## bill_depth_mm
                                     82.98
                                                 22.32 3.717 0.000237 ***
## speciesChinstrap
                                   1460.15
                                                680.39 2.146 0.032610 *
```

```
## speciesGentoo
                                     644.88
                                                542.57
                                                        1.189 0.235481
## bill_depth_mm:speciesChinstrap
                                     -83.53
                                                 37.01 -2.257 0.024666 *
                                      36.17
## bill_depth_mm:speciesGentoo
                                                 34.48
                                                         1.049 0.294955
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
##
## Residual standard error: 286.8 on 325 degrees of freedom
     (11 observations deleted due to missingness)
## Multiple R-squared: 0.8758, Adjusted R-squared: 0.8732
## F-statistic: 327.5 on 7 and 325 DF, p-value: < 2.2e-16
To test the null hypothesis H_0: \beta_{Chinstrap} = \beta_{Gentoo} = 0, we do a F-test.
anova(penguin.model)
## Analysis of Variance Table
##
## Response: body_mass_g
                          Df
                                Sum Sq
                                          Mean Sq
                                                    F value
                                                               Pr(>F)
## flipper_length_mm
                           1 164047703 164047703 1994.7424 < 2.2e-16 ***
                                          9416589 114.5013 < 2.2e-16 ***
## sex
                               9416589
## bill_depth_mm
                           1
                               3667377
                                          3667377
                                                    44.5936 1.051e-10 ***
                                                    64.8743 < 2.2e-16 ***
## species
                           2 10670525
                                          5335262
## bill_depth_mm:species
                           2
                                729458
                                           364729
                                                     4.4349
                                                              0.01258 *
## Residuals
                         325 26728014
                                            82240
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
We see that the p-value for species is very low, so we do not reject H_0
autoplot(penguin.model, smooth.colour = NA)
## Warning: Removed 333 row(s) containing missing values (geom_path).
## Removed 333 row(s) containing missing values (geom_path).
## Removed 333 row(s) containing missing values (geom_path).
```



Problem 3

Now, the main idea is to classify the species of the penguins for a given body mass and flipper length. The dataset contains 3 type of penguin species, yet, classifying 3 penguin species is not straightforward. So, for simplicity, we define our goal to classify a penguin belonging to the species - Adelie, or not Adelie. This will give us two-class classification problem instead of three.

Since, we will use only three parameters from the dataset, it is wise to create a small dataset, and work with it. First, the variables should be converted to numeric (because knn function cannot handle the int class, and generates errors) and removes any missing observations.

Now, we need two datasets: the one to train the model, then to test the model. It is common to divide 70% of dataset as a training dataset, and the rest will be a test dataset.

```
set.seed(4268)
# 70% of the sample size for training set
```

```
training_set_size <- floor(0.7 * nrow(Penguins_reduced))
train_ind <- sample(seq_len(nrow(Penguins_reduced)), size = training_set_size)
train <- Penguins_reduced[train_ind, ]
test <- Penguins_reduced[-train_ind, ]</pre>
```

a)

i) Logistic regression

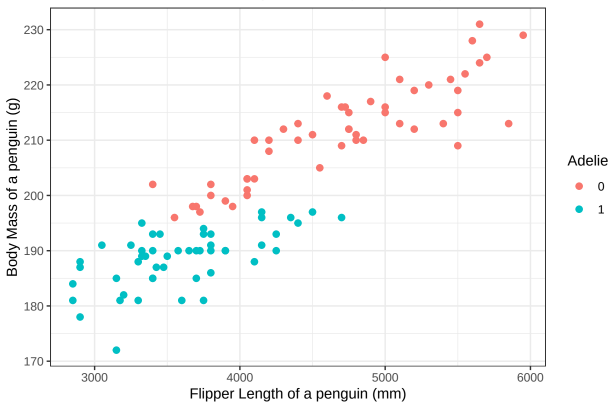
```
## Estimate Std. Error z value Pr(>|z|)
## (Intercept) 37.7618776 5.1761640773 7.295340 2.979055e-13
## body_mass_g 0.0007120 0.0004619996 1.541127 1.232859e-01
## flipper_length_mm -0.2055804 0.0324291723 -6.339367 2.307116e-10
```

Here, we train the model using train dataset, and the result is available above.

```
prob_test_lr <- predict(log_reg, newdata = test, type = "response")
pred_test_lr <- ifelse(prob_test_lr > 0.5, 1, 0)
```

predict() gives the probabilities of the species = Adelie (class = 1). Then, we apply a cutoff value of 0.5 to those probabilities, to get predicted classes.





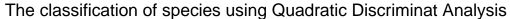
Then, we use test dataset for testing the model. The plot illustrates the testing result.

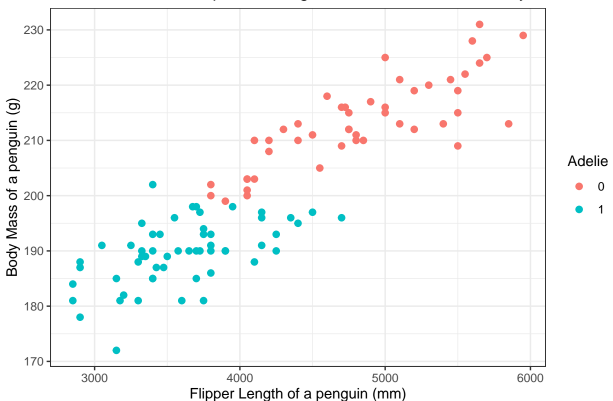
ii) Quadratic Discriminant Analysis We do the same for classification using Quadratic Discriminant Analysis

```
qda_reg = qda(adelie ~ body_mass_g + flipper_length_mm, data = train)
prob_test_qda <- predict(qda_reg, newdata = test, type = "response")$posterior
pred_test_qda <- predict(qda_reg, newdata = test, type = "response")$class</pre>
```

Because QDA identifies the probabilities of having species = Adelie (class = 1), and having species = not Adelie (class = 0), predict() function will give both probabilities and already cut-off value (0.5) applied predictions. So, \$posteriors will give the probabilities, \$class will give 0.5 cut-off value applied corresponding classes.

+ theme





Again, this plot illustrates the testing result.

```
knnMod = knn(train = train, test = test, cl = train$adelie, k = 25, prob = T)
knn_r <- data.frame(data.matrix(knnMod), attributes(knnMod)$prob)</pre>
```

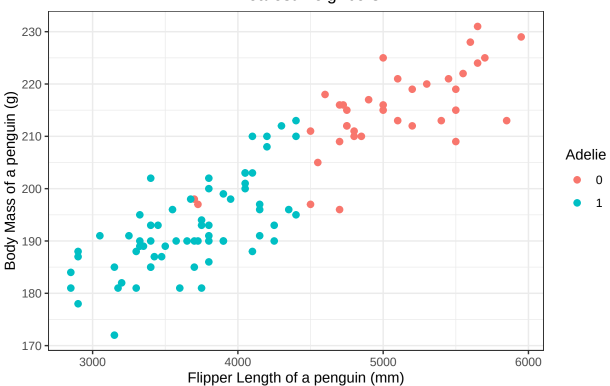
iii) K-nearest Neighbors After having the model, we put the result in matrix form for better usage.

```
probKNN = ifelse(knnMod == 0, 1 - attributes(knnMod)$prob, attributes(knnMod)$prob)
predKNN = ifelse(probKNN > 0.5, 1, 0)
```

This is the way to get the probabilities of species = Adelie (class = 1). Then, again we apply 0.5 cut-off value to get the predictions.

```
xlab("Flipper Length of a penguin (mm)") + ylab("Body Mass of a penguin (g)") +
labs(shape = "Adelie", color = "Adelie") + ggtitle("The classification of species using \n K-near
theme_bw() + theme(plot.title = element_text(hjust = 0.5))
gg_knn
```

The classification of species using K–nearest Neighbors



iv) Sensitivity & specificity For Logistic regression

```
##
     Estim. prob. of Y=1 Predicted class True class
               0.9616473
## 1
                                                     1
                                         1
## 2
               0.9028693
                                         1
                                                     1
               0.6321050
                                         1
## 3
                                                     1
## 4
               0.7859021
                                         1
                                                     1
               0.9324308
## 5
                                         1
                                                     1
## 6
               0.7688126
```

```
print("The confusion matrix is:")
## [1] "The confusion matrix is:"
conf_mat_log_r
##
            true
## predicted 0 1
           0 52 1
           1 8 42
##
test_error_rate_lr = (conf_mat_log_r[1, 2] + conf_mat_log_r[2, 1])/sum(conf_mat_log_r)
print(paste( "The test error for Logistic regression -", signif(test_error_rate_lr, digits = 4)))
## [1] "The test error for Logistic regression - 0.08738"
sensitivity_lr = conf_mat_log_r[2, 2]/(conf_mat_log_r[1, 2]+conf_mat_log_r[2, 2])
specificity_lr = conf_mat_log_r[1, 1]/(conf_mat_log_r[2, 1]+conf_mat_log_r[1, 1])
print(paste( "The sensitivity - ", signif(sensitivity_lr, digits=4)))
## [1] "The sensitivity - 0.9767"
print(paste( "The specificity - ", signif(specificity_lr, digits=4)))
## [1] "The specificity - 0.8667"
For Quadratic Discriminant Analysis
predictions_qda = data.frame(pred_test_qda, test[, "adelie"])
colnames(predictions qda) = c("Predicted class", "True class")
head(predictions_qda)
    Predicted class True class
## 1
                   1
## 2
                   1
                              1
## 3
                   1
                              1
## 4
                   1
                              1
## 5
                   1
                              1
## 6
                              1
conf_mat_qda = table(predicted = predictions_qda["Predicted class"][,1], true =
        predictions_qda["True class"][,1] )
print("The confusion matrix is:")
```

[1] "The confusion matrix is:"

```
conf_mat_qda
##
            true
## predicted 0 1
          0 46 1
           1 14 42
test_error_rate_qda = (conf_mat_qda[1, 2] + conf_mat_qda[2, 1])/sum(conf_mat_qda)
print(paste( "The test error for Quadratic Discriminant Analysis -", signif(test_error_rate_qda, digits
## [1] "The test error for Quadratic Discriminant Analysis - 0.1456"
sensitivity_qda = conf_mat_qda[2, 2]/(conf_mat_qda[1, 2]+conf_mat_qda[2, 2])
specificity_qda = conf_mat_qda[1, 1]/(conf_mat_qda[2, 1]+conf_mat_qda[1, 1])
print(paste( "The sensitivity - ", signif(sensitivity_qda, digits=4)))
## [1] "The sensitivity - 0.9767"
print(paste( "The specificity - ", signif(specificity_qda, digits=4)))
## [1] "The specificity - 0.7667"
For K-nearest Neighbors
predictions knn = data.frame(knn r[, 2], knn r[, 1], test[, "adelie"])
colnames(predictions_knn) = c("Estim. prob. of Y=1", "Predicted class", "True class")
head(predictions knn)
    Estim. prob. of Y=1 Predicted class True class
##
## 1
              0.6000000
## 2
              0.6666667
                                       1
                                                  1
## 3
              0.6800000
                                       1
## 4
              0.6400000
                                       1
                                                  1
## 5
              0.6800000
                                       1
## 6
              0.6400000
                                                  1
                                       1
conf_mat_knn = table(predicted = predictions_knn["Predicted class"][,1], true =
        predictions_knn["True class"][,1] )
print("The confusion matrix is:")
## [1] "The confusion matrix is:"
conf_mat_knn
##
            true
## predicted 0 1
           0 35 2
##
##
           1 25 41
```

legend("bottomright", legend = c("Logistic Regression", "Quadradic Discriminant \nAnalysis", "K-nearest

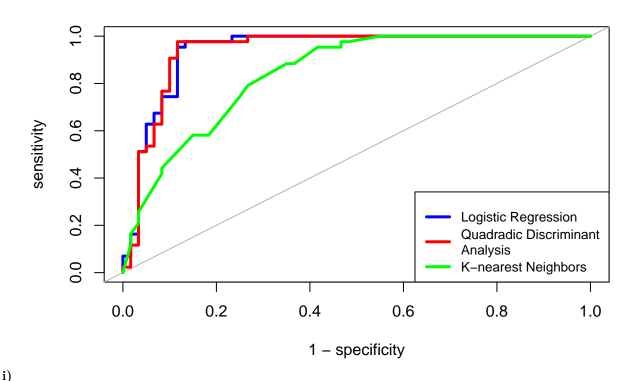
test_error_rate_knn = (conf_mat_knn[1, 2] + conf_mat_knn[2, 1])/sum(conf_mat_knn)

probKNN = ifelse(knnMod == 0, 1 - attributes(knnMod)\$prob, attributes(knnMod)\$prob)

knn_roc = roc.curve(test\$adelie, probKNN, col = "green", lwd=3, add = T)

col = c("blue", "red", "green"))

ROC curve



Areas Under Curves:

```
print(paste( "AUC of Logistic Regression ", signif(lr_roc$auc, digits = 4)))
```

[1] "AUC of Logistic Regression 0.9391"

```
print(paste( "AUC of Quadradic Discriminant Analysis ", signif(qda_roc$auc, digits = 4)))
```

[1] "AUC of Quadradic Discriminant Analysis 0.938"

```
print(paste( "AUC of K-nearest Neighbors ", signif(knn_roc$auc, digits = 4)))
```

[1] "AUC of K-nearest Neighbors 0.8417"

ii) ROC curve - Receiver operating characteristics curve displays the relation between False Positive rate and True Positive rate for all possible threshold values (0 to 1). Since True Positive rate is $\frac{True\ Positives}{All\ Positives}$, it is expressed as sensitivity, and False Positive rate as 1 - Specificity. If the model is ideal model, the sensitivity = 1, and specificity = 1, so, the curve will be on the top left. While a straight line will represent a model with random guesses on the outcome.

In order to check the overall performance of the model, AUC (Area Under Curve) values are compared, meaning that if AUC value is high, that means the curve is closer to the ideal situation.

The comparison of the models we have: Logistic regression, Quadratic Discriminant Analysis, K-nearest Neighbors

When we look at the ROC Curves, we clearly see that the model that performs worse than others is K-nearest Neighbors. This is proved by AUC area. (AUC area of KNN = 0.8417).

The performances of Logistic regression and Quadratic Discriminant Analysis are almost the same. Yet, there is a slight difference in AUC areas. Based on AUC areas, Logistic regression performs a bit better than Quadratic Discriminant Analysis (AUC area of LogReg = 0.9391, AUC area of QDA = 0.938). However, the overall conclusion is Logistic Regression and Quadratic Discriminant Analysis are similarly good models for interpreting the data.

iii) If the task is to create an interpretable model, the model that has to be chosen is Logistic Regression. The reason is Logistic Regression outputs a table that contains all the necessary information to analyze the relation between two variable. The features like Intercept, p-value, t-value have key roles in interpretation.

c)

```
\begin{split} odds &= \frac{p_i}{1 - p_i} = \frac{P(Y_i = 1 | X = x)}{P(Y_i = 0 | X = x)} = exp(\beta_0) \cdot exp(\beta_1 x_1) \cdot exp(\beta_2 x_2) \\ odds \; ratio &= \frac{odds[Y_i = 1 | X = x_1 + 1000]}{odds[Y_i = 1 | X = x_1]} = \frac{exp(\beta_0) \cdot exp(\beta_1 (x_1 + 1000)) \cdot exp(\beta_2 x_2)}{exp(\beta_0) \cdot exp(\beta_1 x_1) \cdot exp(\beta_2 x_2)} = exp(\beta_1 * 1000) \end{split}
```

We find β_1 from coefficient table of Logistic regression

```
summary(log_reg)$coef
```

```
## (Intercept) 37.7618776 5.1761640773 7.295340 2.979055e-13 ## body_mass_g 0.0007120 0.0004619996 1.541127 1.232859e-01 ## flipper_length_mm -0.2055804 0.0324291723 -6.339367 2.307116e-10 So, \beta_1 = 0.000712 print(paste( "The odds will be multiplied by ", signif(exp(coef(log_reg)[2]*1000), digits = 4))) ## [1] "The odds will be multiplied by 2.038"
```

d)

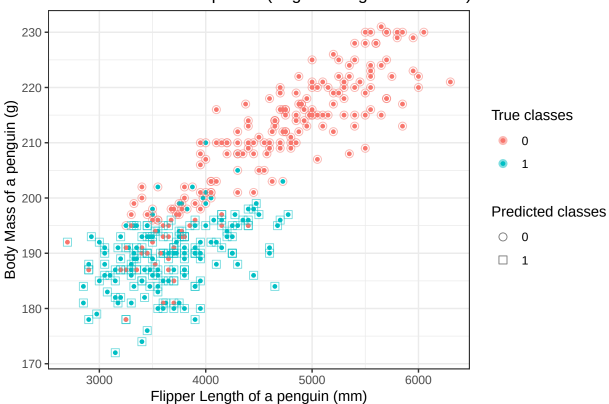
```
prob_whole_lr <- predict(log_reg, newdata = Penguins_reduced, type = "response")

pred_whole_lr <- ifelse(prob_whole_lr > 0.5, 1, 0)

pred_whole_lr_sh = pred_whole_lr

whole_lr_df = bind_rows(mutate(Penguins_reduced, pred_whole_lr, pred_whole_lr_sh))
whole_lr_df$pred_whole_lr = as.factor(whole_lr_df$pred_whole_lr)
whole_lr_df$pred_whole_lr_sh = as.factor(whole_lr_df$pred_whole_lr_sh)
Penguins_reduced$adelie = as.factor(Penguins_reduced$adelie)
```





Problem 4

a)

Answers for the questions are as follows: (i) True; (ii) False; (iii) True; (iv) False;

b)

Heart disease (chd) probability for a non-smoking male with systolic blood pressure (sbp) 150 is 0.10096. The R code for the problem solution is provided below:

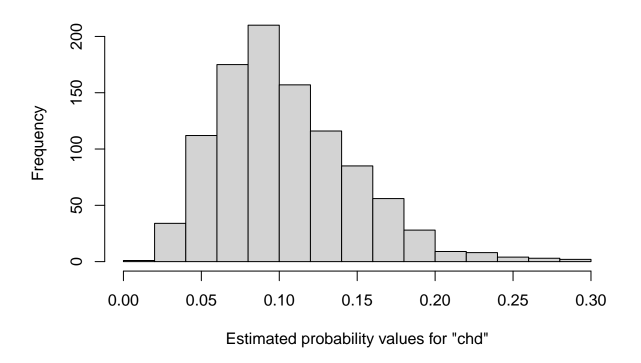
```
#Accessing the data
id<-"1chRpybM5cJn4Eow3-_xwDKPKyddL9M2N"
d.chd<-read.csv(sprintf("https://docs.google.com/uc?id=%s&export=download",id))
#Creating a predicting model
glm_fit = glm(chd~sex+sbp+smoking, data = d.chd, family = binomial)
#Predicting the disease probability
temp_d <- data.frame(sex=1, sbp=150, smoking=0)
predict(glm_fit,temp_d,type = 'response')

##     1
##     0.10096</pre>
c)
```

The R code generating solution for the first problem can be found below. As one can see most of the prediction values are between 0.05 and 0.15.

```
#Create function of interest
func_interest <- function(used_data){
   glm_fit = glm(chd~sex+sbp+smoking, data = used_data, family = binomial)
   predict(glm_fit,data.frame(sex=1, sbp=150, smoking=0),type = 'response')
}
#Implement bootstrap for 1000 iterations and save the results in A
B = 1000
A = replicate(B,0)
for (i in 1:B){
   A[i]=func_interest(d.chd[sample(1:nrow(d.chd), 500, replace = T),])
}
hist(A, main = 'Estimated probability for "chd"', xlab = 'Estimated probability values for "chd"')</pre>
```

Estimated probability for "chd"



The standard error derived from predictions above equals to 0.00134.

One can find the R code generated for solving the problem here.

```
#Calculating standard error
std_error_func <- function(x) sd(x) / sqrt(length(x))
Std_error <- std_error_func(A)
print(Std_error)</pre>
```

[1] 0.001387751

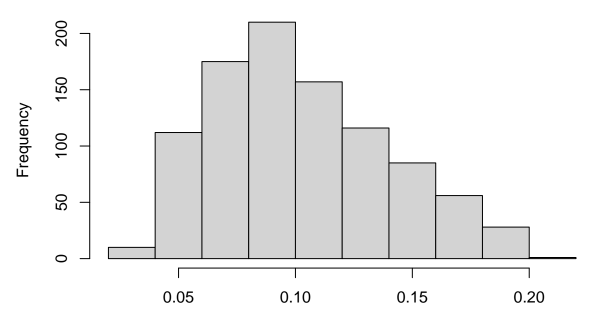
The 95 % quantile interval for the bootstrap samples lies between 0.03915905 and 0.20153356. One can see how they distributed in the histogram below.

```
#Deriving quantiles and sorting bootstrap results within them
q = quantile(A,probs = c(0.025, 0.975))

After_q <-A[A<q[2]]
After_q <-After_q[After_q>q[1]]

hist(After_q, main = 'Quantile of estimated probability of "chd"', xlab = 'Quatiled estimated probability
```

Quantile of estimated probability of "chd"



Quatiled estimated probability values of "chd"

The expected probability value is 0.1036039. In my opinion the range of plausib- le values are from 0.06 to 0.12. The R code for solving this problem is here

```
#Calculating values probabilities and expected probability value

EP = 0
for(i in 1:length(After_q)){
   EP = EP+After_q[i]*((1/950))
}
print(EP)
```

[1] 0.1025979

d)

The answers for the questions in problem d) are as follows. (i) True, (ii) False, (iii) True, (iv) False.