Final Project

Ingrid J. Lu 260773949; Grace Ma 260761707; Xinbei Wan 260777034; Wanqi Wang

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Background

Your sister and her partner are expecting a child soon, and she just went to her obstetrician for her routine check. Because she in her second trimester, her obstetrician asks her to do a fetal cardiogram. The results will not get back to her until a week later. Your sister is a bit of a hypochondriac, so she is afraid that there is something wrong with her child.

So, her partner turns to you, a Masters student who is studying inferential statistics for health, hoping to find some results to calm your sister down.

Luckily, you have identified a publicly available dataset that includes thousands of fetal cardiogram results, and the classification of these babies' health status. You need to convince your sister that she and her child will be safe.

Note: the objective of this exercise is to consolidate all the important concepts covered in EPIB607. When answer each question, be sure to include any units and assumptions and define all parameters, when appropriate. The following questions are based on the publicly available dataset "Fetal Classification", please find all attribute information of the data from the link.

Use the following code to set-up your dataframe:

```
# Set-up
df_fh <- readr::read_csv(here::here("fetal_health.csv")) %>%
select(!starts_with("histogram"))
```

Question 1 Data Visualization and Summary Statistics

 \mathbf{a}

Is this data ready for you to work with? If no, transform it into a ready to use form, if yes, explain.

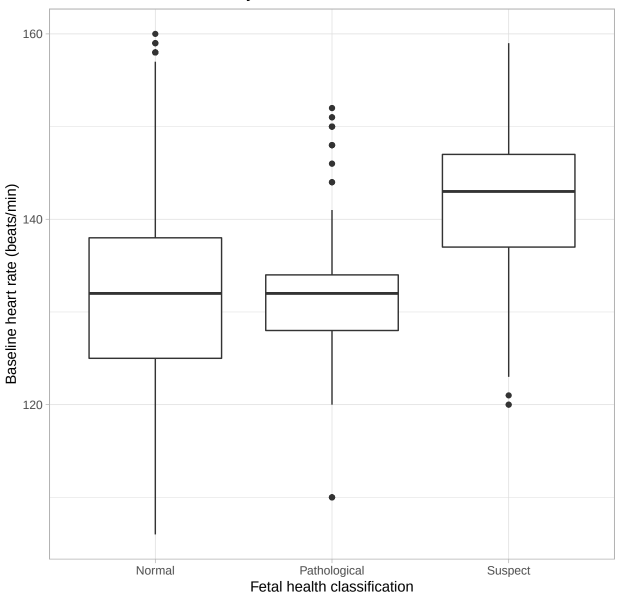
Solution: Yes, this data is ready to use because it is in a tidy format. Each column is a variable and each row is one observation. Each cell contains 1 value of the variable.

b)

Looking at the 3 different classification of fetal health status and each fetus' baseline heart rate, provide an appropriate graphic summarizing the distribution of each of baseline heart rates for each class. Be sure to provide the correct title and label for the plot.

Solution:

Baseline fetal heart rate by health classification



$\mathbf{c})$

Comment on the boxplot, what are the characteristics of each category? Solution:

d)

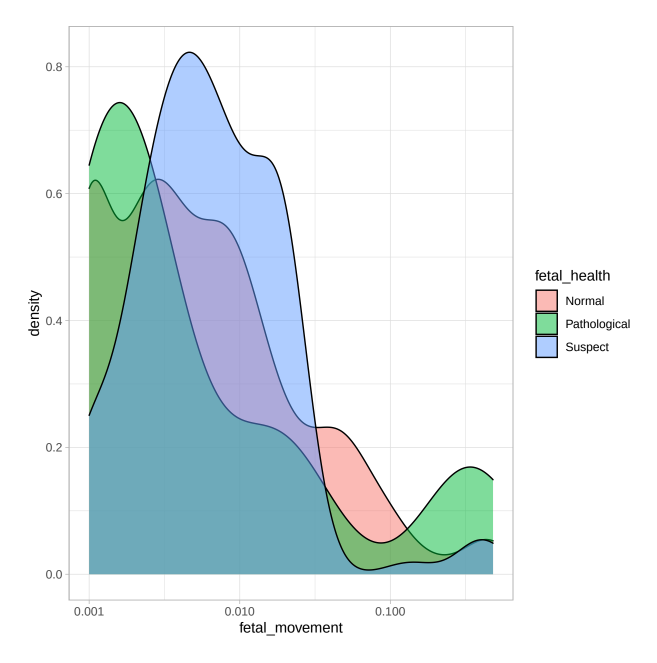
Your sister is concerned that her baby does not move as much, which could be a sign of an unhealthy pregnancy.

Use a density graph, show her the distribution of fetal movements according to different fetal health classifications. Interpret to her the meaning of the box plots.

Solution:

```
## 1.d

df_fh %>%
    ggplot(aes(x = fetal_movement, fill = fetal_health))+
    geom_density(alpha = 0.5) +
    scale_x_log10()
```

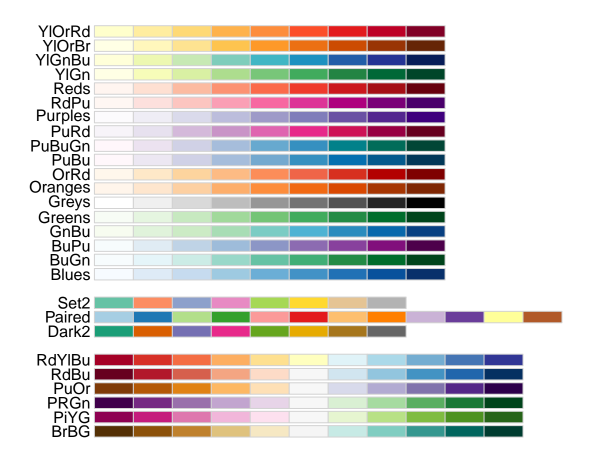


 $\mathbf{e})$

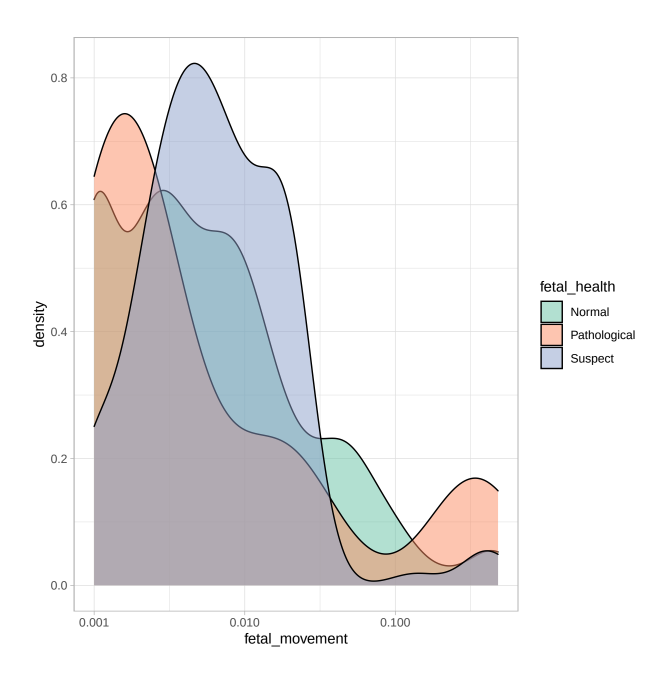
Your sister's partner tells you that he is colourblind, please fix the graph produced above to provide him with a colourblind-friendly graph.

Solution: We can use the "RColorBrewer" package and look for a colourblind-friendly palette. Then apply the package to our ggplot.

```
display.brewer.all(colorblindFriendly = TRUE)
```



```
df_fh %>%
  ggplot(aes(x = fetal_movement, fill = fetal_health))+
  geom_density(alpha = 0.5) +
  scale_x_log10() +
  scale_fill_brewer(palette="Set2")
```



Question 2

a)

Your sister thinks that more uterine contraction means that her fetus is unhealthy and she is more likely to give a pre-term birth.

Is her statement true? What is the mean uterine contraction for each class? Solution:

```
#---Question 2------
df_fh_n <- df_fh %>% filter(fetal_health == "Normal")
mean(df_fh_n$uterine_contractions)

## [1] 0.004780665

df_fh_s <- df_fh %>% filter(fetal_health == "Suspect")
mean(df_fh_s$uterine_contractions)

## [1] 0.002389831

df_fh_p <- df_fh %>% filter(fetal_health == "Pathological")
mean(df_fh_p$uterine_contractions)
```

[1] 0.003784091

Looking at the mean, this statement is not true. The Normal category has the highest number of mean uterine contractions. And the mean uterine contraction in the Pathological category is lower than the mean in the Normal category. The Suspect category has the lowest mean uterine contraction out of the three classes.

Nevertheless, further testing is required to confirm this observation.

b)

Since we have a small sample size for those who are suspected to be pathological and those who are determined to be pathological, what is one method that we can use to artificially create a pseudo-population and calculate the median of these two groups? State your assumptions.

Solution: Bootstrapping, the assumptions are simple random sampling and the samples are representative of the population.

```
B <- 1000
set.seed(3949)

R_s <- replicate(B, { # first argument, # of replicates
    df_fh_s %>%
        dplyr::slice_sample(n = nrow(df_fh_s), replace = T) %>%
        summarise(mean = mean(df_fh_s$uterine_contractions)) %>%
        pull(mean)
})

mean(R_s)
```

[1] 0.002389831

```
R_p <- replicate(B, { # first argument, # of replicates
    df_fh_p %>%
    dplyr::slice_sample(n = nrow(df_fh_p), replace = T) %>%
    summarise(mean = mean(df_fh_p$uterine_contractions)) %>%
```

```
\begin{array}{c} \text{pull(mean)} \\ \text{}) \\ \\ \text{mean(R_p)} \end{array}
```

[1] 0.003784091

c)

What is one weakness of using the bootstrapping method?

Solution: The bootstrapping method assumes that the sample is representative of the whole population. However, when we have a small sample size, that is not always the case. There could be sampling errors which can skew the distribution of the sample. And the bootstrapping method cannot correct this error as it can only sample within this small set of data. As the saying goes, "garbage in, garbage out". If the sample is not representative of the population, the results obtained from bootstrap will not be accurate and will result in an incorrect inference of the population parameter.

Question 3. p-value, power

For the purpose of this question only, we treat the 2126 individuals as **the entire target population of newborns**.

a)

Calculate the mean and standard deviation of the baseline fetal heart rate.

```
mean_hr <- mean(df_fh$`baseline value`)
mean_hr

## [1] 133.3039

sd_hr <- sqrt(var(df_fh$`baseline value`)*(length(df_fh$`baseline value`)-1)/length(df_fh$`baseline value`)
## [1] 9.83853</pre>
```

b)

Your sister claimed that, she read on a magazine, that the baseline fetal heart rates of fetuses with "suspect" health status are above average. Take a simple random sample of 10 fetuses with "suspect" health status, and measure their heart rate to obtain a sample mean of 141.68. Heart rates are scaled to be normally distributed. Does the sample provide evidence to reject null hypothesis? State your null and alternative hypothesis.

```
pnorm(q = 141.68, mean = 133.30, sd = 9.84/sqrt(10), lower.tail = FALSE)
## [1] 0.003539786
```

```
H_0: \mu = 133.30, H_A: \mu > 133.30
```

The p-value of one sided test is 0.0035. This sample provides evidence against the null hypothesis. The p-value tells us the probability of observing the sample size mean of 141.68 under the null hypothesis distribution is very unlikely.

c)

So your sister asks you now, what is the probability that you can detect the baseline fetal heart rates of fetuses with "suspect" health status are at least 8.38 heart beats higher than average, using a one-sided test and sample size 10 and a 0.05 level test?

```
H_0: \mu = 133.30, H_A: \mu > 141.30
```

```
# cutoff to reject the null
cutoff <- qnorm(p = 0.95, mean = 133.30, sd = 9.84/sqrt(10))
cutoff</pre>
```

[1] 138.4183

```
# probability of observing this cutoff or greater under the alternative
pnorm(q = cutoff, mean = 141.68, sd = 9.84/sqrt(10), lower.tail = FALSE)
```

[1] 0.8527324

d)

A sample size of 10 fetuses with "suspect" health status will have at least 85% power to detect a difference of 8.38 heart beats. Use a simulation based approach to reproduce the sample size calculation for the baseline fetal heart rates of fetuses with "suspect" health status and average.

```
set.seed(490)

power_distribution <- replicate(n = 1000, expr={
    sample.size <- 10

suspect <- rnorm(sample.size, mean = 141.68, sd = 9.83)
    SEM <- sd(suspect)/sqrt(sample.size)

pnorm(q = mean(suspect), mean = 133.30, sd = SEM, lower.tail = FALSE) < 0.05
})

prop.table(table(power_distribution))</pre>
```

```
## power_distribution
## FALSE TRUE
## 0.146 0.854
```

The percentage of samples that results in a p-value less than 0.05 is 85.4%, which shows the study is powered at 85% to detect the difference.

Question 4.1

Suppose this data represents the **population of newborns in one hospital** and you take a simple random sample of 100 babies from the population.

a)

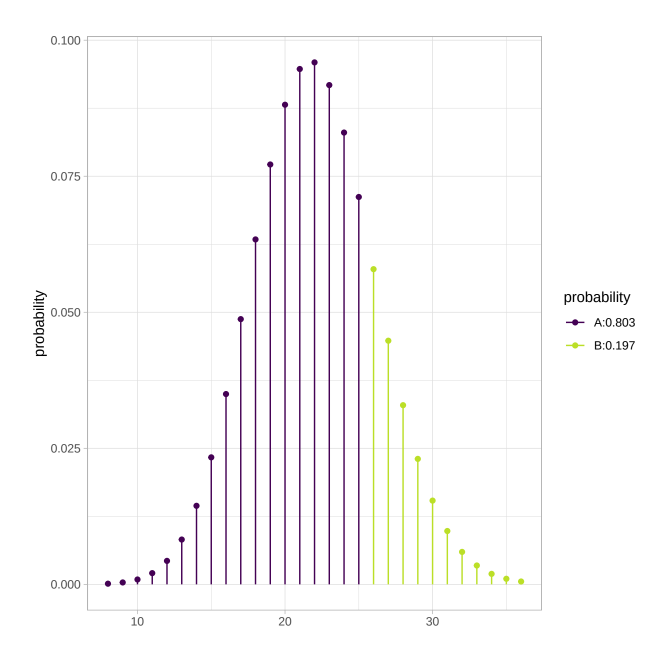
What is the probability that your sample contains more than 25 babies has abnormal health status (1 = health, 2 = suspect, 3 = Pathological)?

```
## ---- Question-5a

df_fh %>%
  select(fetal_health)%>%
  filter(fetal_health != 1)%>%
  nrow()
```

[1] 2126

```
#there are 471 abnormal in this population
#the probability of having abnormal is 471/2126 = 0.22
1 - mosaic::xpbinom(q = 25, size = 100, prob = 0.22)
```



[1] 0.1972269

The probability of having more than 30 abnormal is 0.197.

b)

Turns out that your sample actually contains 20 babies with abnormal health status. What is the 95% confidence interval of this proportion? Can you use a normal approximation for this sample? Why or why not?

```
## ---- Question-5b -----
mosaic::binom.test(x = 20, n = 100, ci.method = "Clopper-Pearson")
```

```
##
##
##
## data: 20 out of 100
## number of successes = 20, number of trials = 100, p-value = 1.116e-09
## alternative hypothesis: true probability of success is not equal to 0.5
## 95 percent confidence interval:
## 0.1266556 0.2918427
## sample estimates:
## probability of success
##
                      0.2
mosaic::binom.test(x = 20, n = 100, ci.method = "Wald")
   Exact binomial test (Wald CI)
##
##
## data: 20 out of 100
## number of successes = 20, number of trials = 100, p-value = 1.116e-09
## alternative hypothesis: true probability of success is not equal to 0.5
## 95 percent confidence interval:
## 0.1216014 0.2783986
## sample estimates:
## probability of success
##
                      0.2
```

The 95% CI using Clopper-Pearson method: [0.127,0.292] The 95% CI using exact method(ie: normal approximation): [0.122,0.278] The two methods give similar 95% CIs. Normal approximation can be used here since the sample size is large enough to generate a binomial distribution approximating the normal distribution and for CLT to kick in.

c)

##

Another sample taken have the same proportion of event but the sample size is now only 10 and the count of abnormal is 2. Calculate the 95% CI using this sample and compare it with the one you have in b). Describe their difference and the reason why.

```
## ---- Question-5c -----
mosaic::binom.test(x = 2, n = 10, ci.method = "Clopper-Pearson")
##
##
```

```
## data: 2 out of 10
## number of successes = 2, number of trials = 10, p-value = 0.1094
## alternative hypothesis: true probability of success is not equal to 0.5
## 95 percent confidence interval:
## 0.02521073 0.55609546
## sample estimates:
## probability of success
## 0.2
```

The 95% CI for this sample is: [0.0252, 0.556]. 95% CI in c) with sample size = 10 is wider than the 95% CI in b) with sample size = 100. The sample size is different so the standard error is different. The 95% CI is calculated using the formula:

$$\bar{y} - 1.96(\frac{\sigma}{\sqrt{n}}), \bar{y} + 1.96(\frac{\sigma}{\sqrt{n}})$$

If the sample size n is larger, the standard error(sigma/sqrt n) is smaller. The value 1.96*standard error is also smaller, and results in a narrower confident interval.

Question 4.2

You continue to work with the simple random sample. This time you take 100 babies from the a different hospital as the sample from the population.

a)

Your sample contains 30 babies with abnormal health status. What is the rate and 95% CI of health abnormality? Interpret your result.

```
## ---- Question-6a -----
stats::poisson.test(x = 30, T = 100)

##
## Exact Poisson test
##
## data: 30 time base: 100
## number of events = 30, time base = 100, p-value = 4.154e-16
## alternative hypothesis: true event rate is not equal to 1
## 95 percent confidence interval:
## 0.2024087 0.4282687
## sample estimates:
## event rate
```

Rate: 0.3~95% CI: [0.20,0.43] There is a 95% chance that the confidence interval [0.20,0.43] captures the true rate.

b)

##

0.3

According to the data of the population with 2126 babies, 471 babies have abnormal health status. The expectation of the baby having abnormal health status of the original hospital is only 0.22. Does your sample suggest that the babies coming from two hospitals have significantly different rate of abnormal health? Calculate the 95% CI for the rate ratio both by hand and using a one-step canned function.

```
## ---- Question-6a ------
stats::poisson.test(x = 30, T = 100)
```

```
##
## Exact Poisson test
##
## data: 30 time base: 100
## number of events = 30, time base = 100, p-value = 4.154e-16
## alternative hypothesis: true event rate is not equal to 1
## 95 percent confidence interval:
## 0.2024087 0.4282687
## sample estimates:
## event rate
## 0.3
```

0.2024087/0.22

[1] 0.9200395

0.4282687/0.22

```
## [1] 1.946676
```

```
stats::poisson.test(x = 30, T = 471/2126*100)
```

```
##
## Exact Poisson test
##
## data: 30 time base: 471/2126 * 100
## number of events = 30, time base = 22.154, p-value = 0.1092
## alternative hypothesis: true event rate is not equal to 1
## 95 percent confidence interval:
## 0.9136327 1.9331192
## sample estimates:
## event rate
## 1.35414
```

95% CI by hand: [0.92.1.95] 95% CI by canned function: [0.91,0.933] According to the Poisson test and by hand calculation, two 95% CI for the rate ratio both contains the null value 1. The 95% CI and the p-value suggests that the difference between two hospitals is not statistically significant.

Question 5

a)

Your sister's partner said, their obstetrician told them that the Fetal Heart Rate could be a reflection of a lower value of short term variability. Can you conduct a linear regression to test it? Is it significant?

Solution:

```
reg1 <- lm(mean_value_of_long_term_variability ~ `baseline value`, data = df_fh)
summary(reg1)</pre>
```

```
##
## Call:
## lm(formula = mean_value_of_long_term_variability ~ 'baseline value',
       data = df_fh)
##
##
## Residuals:
     Min
              10 Median
                            3Q
                                  Max
## -8.615 -3.526 -0.756 2.647 42.525
##
## Coefficients:
                    Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                    10.63425
                                1.65791
                                          6.414 1.74e-10 ***
## 'baseline value' -0.01835
                                0.01240 - 1.480
                                                   0.139
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Residual standard error: 5.627 on 2124 degrees of freedom
## Multiple R-squared: 0.00103,
                                    Adjusted R-squared:
## F-statistic: 2.19 on 1 and 2124 DF, p-value: 0.1391
```

According to the linear regression, there is no significant association between baseline heart rate and value of short term variability.

b)

If you want to use a logistic regression model to use **Number of fetal movements per second** to the predict the classification of fetal health outcome. Describe how will you process these three variables to fit a logistic regression.

Solution:

- 1. In a logistic regression, the outcome variable is binary variable. In this example, the fetal health outcome will be classified as Pathological, and non-pathological (including normal and suspect).
- 2. We will find cut points for the Number of fetal movements per second based on previous literature and hypothesis.

c)

Suppose the normal number of fetal movements per second is within 0.01. Fit the logistic regression model and provide a 95% CI.

```
df_fh$f_mov <- ifelse(df_fh$fetal_movement>0.01, 1, 0)
df_fh$f_patho <- ifelse(df_fh$fetal_health=="Pathological", 1 , 0)</pre>
reg2 <- glm(f_patho~ f_mov, family = binomial (link = "logit"), data = df_fh)
summary(reg2)
##
## Call:
## glm(formula = f_patho ~ f_mov, family = binomial(link = "logit"),
##
       data = df fh)
##
## Deviance Residuals:
##
       Min
                 1Q
                      Median
                                   3Q
                                            Max
## -0.4975 -0.4048 -0.4048 -0.4048
                                         2.2550
##
## Coefficients:
##
               Estimate Std. Error z value Pr(>|z|)
## (Intercept) -2.46056
                           0.08535 -28.829
                                              <2e-16 ***
                                              0.0507 .
## f_mov
                0.43339
                           0.22181
                                     1.954
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 1214.0 on 2125 degrees of freedom
## Residual deviance: 1210.5 on 2124
                                       degrees of freedom
## AIC: 1214.5
##
## Number of Fisher Scoring iterations: 5
exp(confint(reg2))
                    2.5 %
                             97.5 %
## (Intercept) 0.07194141 0.1005501
## f_mov
               0.98016928 2.3459592
```

Question 6

a)

You saw on another paper, that the pathological diagnosis is dependent on fetal movements and abnormal long term variability, provide a regression equation for this model. Remember to define all parameters.

For the purpose of this question, code the presence and absence of abnormal long term variability as 0,1, using the percentage of time with abnormal long term variability. (0% of variability means no abnormal variability, any number higher than 0% suggests there is long term variability)

Solution:

Regression equation:

$$\frac{\pi}{1-\pi} = \frac{\pi_0}{1-\pi_0} * \theta_1^{mov} * \theta_2^{var} \begin{cases} mov = 1 & \text{if movement is equal to or higher than } 0.01 \\ mov = 0 & \text{if movement is lower than } 0.01 \\ var = 1 & \text{if there is abnormal variability} \\ var = 0 & \text{if there is no abnormal variability} \end{cases}$$

$$log(\frac{\pi}{1-\pi}) = log(\frac{\pi_0}{1-\pi_0}) + mov * log(\theta_1) + var * log(\theta_2)$$

 π is the probability of receiving a pathological diagnosis

 π_0 is the probability of receiving a pathological diagnosis, when fetal movement is lower than 0.01 and no abnormal long-term variability

 θ_1 is the odds ratio of receiving a pathological diagnosis, when fetal movement is equal to or higher than 0.01, adjusting for long-term variability.

 θ_2 is the odds ratio of receiving a pathological diagnosis, when there is abnormal long-term variability, adjusting for fetal movement.

mov and var are the risk parameters.

b)

Fit the regression equation, and compare the intercept value, is it different than the fitted value in Question 5.c? Why?

Solution:

```
df_fh$ab_var <- ifelse(
    df_fh$percentage_of_time_with_abnormal_long_term_variability == 0, 0, 1)

reg3 <- glm(f_patho ~ f_mov + ab_var, family = binomial (link = "logit"), data = df_fh)
summary(reg3)</pre>
```

```
##
## Call:
## glm(formula = f_patho ~ f_mov + ab_var, family = binomial(link = "logit"),
##
       data = df_fh)
##
## Deviance Residuals:
##
      Min
                1Q
                     Median
                                           Max
  -0.5081 -0.4182 -0.4182 -0.3866
                                        2.2938
##
##
## Coefficients:
              Estimate Std. Error z value Pr(>|z|)
                            0.1074 -22.284
## (Intercept) -2.3926
                                             <2e-16 ***
                0.4106
                            0.2230
                                   1.841
                                             0.0656 .
## f mov
## ab var
                -0.1635
                            0.1632 - 1.002
                                             0.3165
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
  (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 1214.0 on 2125 degrees of freedom
##
```

```
## Residual deviance: 1209.5 on 2123 degrees of freedom
## AIC: 1215.5
##
## Number of Fisher Scoring iterations: 5
```

You decided to draw a ROC curve and see how well your model works out. Use the pROC package to draw the curve, and comment on what you see.

Solution:

c)

