HE3021 Tutorial 1 Attempt

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```
knitr::opts_chunk$set(echo = TRUE)
library("tidyverse")
library("foreign")
library("ggplot2")
library("lmtest")
library("sandwich")
```

Initial reading of raw data. Packages used are tidyverse, foreign, ggplot2, sandwich and lmtest.

```
dat <- read.dta(file = fil)
dat <- mutate(dat, npvis_squared = npvis**2)

model <- lm(bwght ~ npvis + npvis_squared + cigs + male, dat)</pre>
```

(a)

```
mean_med <- summary(dat[, "bwght"])
mean_bwght <- mean_med[4]
print(mean_bwght)

## Mean
## 3401.122</pre>
```

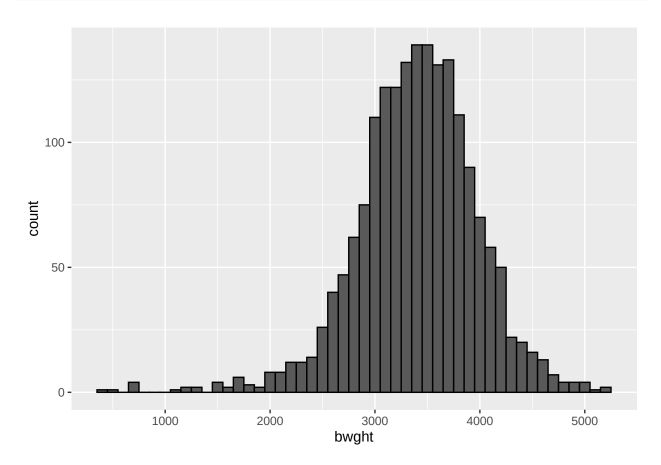
(b)

```
med_bwght <- mean_med[3]
print(med_bwght)
## Median</pre>
```

(c)

3425

```
plot_bwght <- ggplot(dat, aes(x = bwght)) +
    geom_histogram(binwidth = 100, color = "black")
print(plot_bwght)</pre>
```



(d)

```
beta_3 <- model$coefficients["cigs"]
print(beta_3)</pre>
```

cigs ## -9.901333

(e)

 $H_0: \beta_3 = -10, H_a: \beta_3 \neq -10, \alpha = 0.05,$ 2-tailed 1-sample t-test.

```
t_test_2tail_variable_hypo <- function(
   hypothesis, significance_lvl, df, sample_value, standard_error){
   confidence_interval <- hypothesis +c(-1, 1)*qt(p = 1-significance_lvl/2,</pre>
```

```
df = df)*standard_error
if (between(sample_value, confidence_interval[1], confidence_interval[2])){
    return(FALSE)
} else {return(TRUE)}
}

test_beta_3 <- t_test_2tail_variable_hypo(
    -10, 0.05, 1651, beta_3, 3.3330) # TRUE means insignificantly different from -10
confidence_interval <- -10 + c(-1, 1)*qt(p = 0.975, df = 1651)*3.3330
print(confidence_interval)</pre>
```

[1] -16.537352 -3.462648

Since $\beta_3 \in [-16.54, -3.46]$, then we have insufficient evidence to reject $H_0 = -10$.

(f)

 $H_0: \beta_4=100, \, H_a: \beta_4 \neq 100, \, \alpha=0.05, \, \text{2-tailed 2-sample Welsh t-test.}$

```
test_beta_4 <- t_test_2tail_variable_hypo(
    100, 0.05, 1651, beta_3, 27.9757) # TRUE means insignificantly different from +100
confidence_interval <- 100 + c(-1, 1)*qt(p = 0.975, df = 1651)*27.9757
print(confidence_interval)</pre>
```

[1] 45.12841 154.87159

Since $+100 \in [45.1, 154.9]$, then we have insufficient evidence to reject $H_0 = +100$.

(g)

```
beta_1 <- model$coefficients["npvis"]
beta_2 <- model$coefficients["npvis_squared"]</pre>
```

The partial effect is $32.8npvis + (-0.669)npvis^2$.

(h)

```
0 = \frac{\partial bwght}{\partial npvis} = 32.8 + (-0.669)npvis * 2, npvis = \frac{32.8}{2*(0.669)}
```

```
turning_pt <- beta_1/(2*-beta_2)
print(turning_pt)</pre>
```

```
## npvis
## 24.50519
```

npvis = 24.50519 Diminishing marginal returns of additional pre-natal visits since there is not much more the doctor can check.

(i)

 $H_0: 0 = \beta_1 = \beta_2 = \beta_3 = \beta_4$, $H_a: otherwise$, F-test of squared residuals against independent variables.

```
hetero_test <- model$model[, 2:5]
hetero_test <- mutate(hetero_test, residuals = model$residuals)
hetero_test <- mutate(hetero_test, residuals = residuals**2)
hetero_test_model <- lm(residuals ~ npvis + npvis_squared + cigs + male, hetero_test)
hetero_test_p_value <- 4.476*(10**(-5)) # heteroscedastic
```

Since $4.476 * (10^{-5}) < 0.05$, then we have sufficient evidence to reject H_0 , and we cannot have the homoscedascity assumption. To solve it, we can use robust standard errors as sample size n = 1656 > 120. The hypothesis tests for β_3 and β_4 would then be like so:

```
robust_t <- coeftest(model, vcov = vcovHC(model, type = "HCO"))
print(robust_t)</pre>
```

```
##
## t test of coefficients:
##
##
                   Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                 3097.59754
                              99.79408 31.0399 < 2.2e-16 ***
## npvis
                   32.80329
                              12.76511 2.5698 0.010264 *
## npvis_squared
                   -0.66931
                               0.39520 -1.6936 0.090525 .
## cigs
                  -9.90133
                               3.28212 -3.0167 0.002594 **
## male
                  81.31364
                              27.82588 2.9222 0.003523 **
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
test_beta_3_robust <- t_test_2tail_variable_hypo(</pre>
    -10, 0.05, 1651, beta_3, 3.28212) # TRUE means insignificantly different from -10
confidence_interval <-10 + c(-1, 1)*qt(p = 0.975, df = 1651)*3.28212
print(confidence_interval)
```

```
## [1] -16.437556 -3.562444
```

```
test_beta_4 <- t_test_2tail_variable_hypo(
    100, 0.05, 1651, beta_3, 27.82588) # TRUE means insignificantly different from +100
confidence_interval <- 100 + c(-1, 1)*qt(p = 0.975, df = 1651)*27.82588
print(confidence_interval)</pre>
```

[1] 45.42227 154.57773

Since $-10 \in [-16.437556, -3.562444]$, then we cannot reject $H_0: \beta_3 = -10$. Since $+100 \in [45.42227, 154.57773]$, then we cannot reject $H_0: \beta_4 = 100$.

(j)

No, I do not think we can have the zero conditional mean, because income can be correlated with pre-natal visits since poorer people can come from locations where clinics are inaccessible. Income itself can influence mother's nutrition, which can influence baby weight.