## MLR Initial Report

#### Hannah Waddel

# Initial Report: Relationship between Birthweight and Several Variables

In this exercise we are going to perform a multiple linear regression to assess whether age, weight, smoking status, or hypertension status were significantly associated with birthweight.

We will begin by reading in the data. I have provided the data in a SAS dataset, "birthweight.sas7bdat".

### Importing the Data

The data are stored as a SAS dataset, "birthweight.sas7bdat". Before importing the data, make sure that your working directory is set to the folder where you have saved the dataset.

```
setwd("~/Work/BIOS 591P 2021/R Materials/3 MLR")
```

We need to get the data into a format which R can use. R does not have a built-in capability to read in a SAS dataset, but we can install packages which expand R's capabilities.

Packages are easy to install and make R an incredibly flexible programming language. People create R packages to implement new statistical methods or expand R's existing methods.

If a package is of publication quality and makes a non-trivial contribution to R's capabilities, it is stored on the Comprehensive R Archive Network (CRAN). If a package is on CRAN, you can install it using the <code>install.packages()</code> function.

The package we are installing today is called "sas7bdat". It allows us to read in SAS datasets, which are stored as ".sas7bdat" files on your computer.

To install the package, we are using the *install.packages()* function, which requires the name of the package we want as a string (text enclosed by " ").

Let's install sas7bdat now. Make sure that the package name is in quotation marks.

```
install.packages("sas7bdat")
```

```
## Installing package into 'C:/Users/hbwadde/Documents/R/win-library/4.0'
## (as 'lib' is unspecified)

## package 'sas7bdat' successfully unpacked and MD5 sums checked
##
## The downloaded binary packages are in
## C:\Users\hbwadde\AppData\Local\Temp\RtmpkrkkXa\downloaded_packages
```

R will tell you where it is putting the files for the package (default settings should be fine). Your installation was successful if you see "package 'sas7bdat' successfully unpacked and MD5 sums checked".

The sas7bdat package gives us an important function read.sas7bdat(). This works like the read.csv() function we have used, but for SAS datasets.

However, if we try to use read.sas7bdat() right now, we get an error.

```
bwt <- read.sas7bdat("birthweight.sas7bdat")</pre>
```

## Error in read.sas7bdat("birthweight.sas7bdat"): could not find function "read.sas7bdat"

R cannot find the function "read.sas7bdat" because while we have installed the sas7bdat package, we have not loaded it properly for R to use. After we have installed a package, we have to load it into R's library with the *library()* function before R can use the functions inside the package.

```
library(sas7bdat)
```

Notice that we don't need to use quotation marks around the name of the package when we are loading it with library().

Now that we have loaded the sas7bdat function, we can use the read.sas7bdat() function to read in the dataset. We assign the dataset the name "bwt".

```
bwt <- read.sas7bdat("birthweight.sas7bdat")</pre>
```

We can check that the dataset read in correctly with the following functions that do similar things as PROC CONTENTS.

```
#Check number of rows and columns
dim(bwt)
```

```
## [1] 189 7
```

```
#See the first few lines of the dataset head(bwt)
```

```
BWT AGE WT SMOKE HT RACE
        709
             28 120
                             0
## 1
                         1
## 2 10 1021
              29 130
                         0
                             0
                                  2
## 3 11 1135 34 187
                         0
                             1
                                  1
## 4 13 1330
              25 105
                          0
                                  2
## 5 15 1474
                         0
                            0
                                  2
              25
                  85
## 6 16 1588
             27 150
                            0
                                  2
```

```
#See that the columns have the right name names(bwt)
```

```
## [1] "ID" "BWT" "AGE" "WT" "SMOKE" "HT" "RACE"
```

## **Descriptive Statistics**

We use the mean(), sd() and quantile() functions to find descriptive statistics for the continuous/numeric variables.

```
mean(bwt$BWT)
## [1] 2944.714
sd(bwt$BWT)
## [1] 729.187
quantile(bwt$BWT)
    0% 25% 50% 75% 100%
##
  709 2414 2977 3475 5001
mean(bwt$AGE)
## [1] 23.2381
sd(bwt$AGE)
## [1] 5.298678
quantile(bwt$BWT)
    0% 25% 50% 75% 100%
##
## 709 2414 2977 3475 5001
mean(bwt$WT)
## [1] 129.8148
sd(bwt$WT)
## [1] 30.57938
quantile(bwt$WT)
##
     0% 25% 50% 75% 100%
##
     80 110 121 140 250
```

For the discrete/categorical variables, we use the table() function to get the frequency of outcomes.

```
table(bwt$SMOKE)
##
##
     0
          1
## 117
         72
table(bwt$HT)
##
##
     0
          1
## 177
        12
To get the percentages, we divide the output from table() by the number of observations (rows in the dataset),
then multiply by 100.
```

```
table(bwt$SMOKE)/nrow(bwt) * 100

##
## 0 1
## 61.90476 38.09524

table(bwt$HT)/nrow(bwt) * 100

##
## 0 1
## 93.650794 6.349206
```

#### Calculating and Testing Correlation

To calculate correlation between all the variables of the birthweight dataset, we can use the cor() function.

```
cor(bwt)
```

```
BWT
                                          AGE
                                                                 SMOKE
                                                                                  ΗT
##
                  ID
                                                        WT
## ID
          1.00000000
                      0.98117762
                                  0.09504326
                                               0.21565483 -0.16061294 -0.112021205
## BWT
          0.98117762
                      1.00000000
                                  0.09017566
                                               0.18572889 -0.15071982 -0.146062658
## AGE
          0.09504326
                      0.09017566
                                   1.00000000
                                               0.18007315 -0.06008380 -0.015837000
                                               1.00000000 -0.06132677 0.236360402
## WT
          0.21565483 0.18572889
                                  0.18007315
## SMOKE -0.16061294 -0.15071982 -0.06008380 -0.06132677 1.00000000 -0.025532146
## HT
         -0.11202120 \ -0.14606266 \ -0.01583700 \ \ 0.23636040 \ -0.02553215 \ \ 1.000000000
         -0.18297545 -0.17406839 -0.08501461 -0.04520165 -0.03449021 -0.006734069
## RACE
##
                 RACE
## ID
         -0.182975449
         -0.174068392
## BWT
## AGE
         -0.085014613
## WT
         -0.045201649
## SMOKE -0.034490210
## HT
         -0.006734069
## RACE
         1.000000000
```

This gives us every calculated correlation value between every variable in the dataset. We are not interested in some of these variables, so we can also give the cor() function just two variables.

```
cor(bwt$WT,bwt$BWT)

## [1] 0.1857289

cor(bwt$AGE,bwt$BWT)

## [1] 0.09017566
```

To test the correlation values and get confidence intervals for the true correlation, we use the cor.test() function.

```
cor.test(bwt$WT,bwt$BWT)

##

## Pearson's product-moment correlation

##

## data: bwt$WT and bwt$BWT

## t = 2.5848, df = 187, p-value = 0.01051

## alternative hypothesis: true correlation is not equal to 0

## 95 percent confidence interval:

## 0.04416951 0.31997686

## sample estimates:

## cor

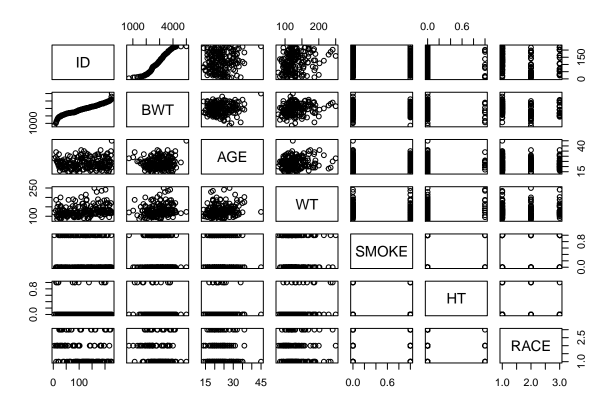
## 0.1857289
```

```
##
## Pearson's product-moment correlation
##
## data: bwt$AGE and bwt$BWT
## t = 1.2382, df = 187, p-value = 0.2172
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## -0.05323986  0.22994635
## sample estimates:
## cor
## 0.09017566
```

cor.test(bwt\$AGE,bwt\$BWT)

To get pairwise plots between each of the variables, we use the pairs() function.

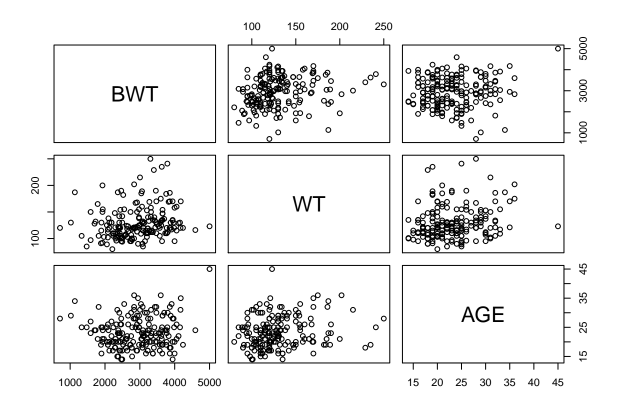
```
pairs(bwt)
```



However, this includes variables we are not interested in, like ID, and categorical variables. We can give pairs() the relevant columns of bwt and it will make the pairwise plots between each of them.

Recall the bracket notation [rows, columns] that we use to pull the columns of bwt that we want. We give it a list, created using c(), of the relevant column names as strings.

```
pairs(bwt[,c("BWT","WT","AGE")])
```



### Fitting Multiple Linear Regression Model

Our model is  $Y = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \beta_3 X_3 + \beta_4 X_4 + E$ 

Where

Y = birthweight (g)

 $X_1 = \text{mother's age (years)}$ 

 $X_2 = \text{mother's weight (lbs)}$ 

 $X_3 = \text{mother's smoking status } (1 = \text{smoked during pregnancy}, 0 = \text{did not smoke})$ 

 $X_4$  = mother's hypertension status (1 = history of hypertension, 0 = no history of hypertension)

 $E={\rm random~error,~assumed~Normal}(0,\,\sigma^2$  )

 $\beta_0 = \text{true y-intercept}$ 

 $\beta_i$  = true slope associated with  $X_i$ , adjusted for the other predictors.

We fit our model with the lm() function again. The arguments for the lm() function are a formula specifying our model and a dataset.

The formula uses the  $\sim$  operator again. The dependent variable (birthweight), goes on the left side of the  $\sim$ , and the independent variables go on the right side of the  $\sim$ . We are fitting a linear model where we add the effects of  $X_1$ ,  $X_2$ ,  $X_3$ , and  $X_4$ , so we add the variables age, weight, smoking status, and hypertension status on the right side of the formula. Remember that variable names in R are case-sensitive.

Make sure to save the results from lm() as an object.

```
bwt.lm <- lm(BWT ~ AGE + WT + SMOKE + HT,data=bwt)</pre>
```

Use the summary() function to see the estimates from lm().

#### summary(bwt.lm)

```
##
## Call:
## lm(formula = BWT ~ AGE + WT + SMOKE + HT, data = bwt)
## Residuals:
##
       Min
                  1Q
                       Median
                                    3Q
                                            Max
                        -0.28
  -2118.71 -436.99
                                        1855.62
##
                                518.01
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 2263.532
                           299.860
                                     7.549 1.97e-12 ***
## AGE
                  5.427
                             9.846
                                     0.551 0.58220
                                     2.953
## WT
                  5.184
                             1.755
                                           0.00355 **
## SMOKE
               -209.863
                           105.408
                                    -1.991 0.04797 *
## HT
               -597.684
                           215.769
                                    -2.770 0.00618 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 701.4 on 184 degrees of freedom
## Multiple R-squared: 0.09433,
                                    Adjusted R-squared:
## F-statistic: 4.791 on 4 and 184 DF, p-value: 0.001065
```

The overall F test is in the bottom row of the summary. The F-test p-value is 0.001065.

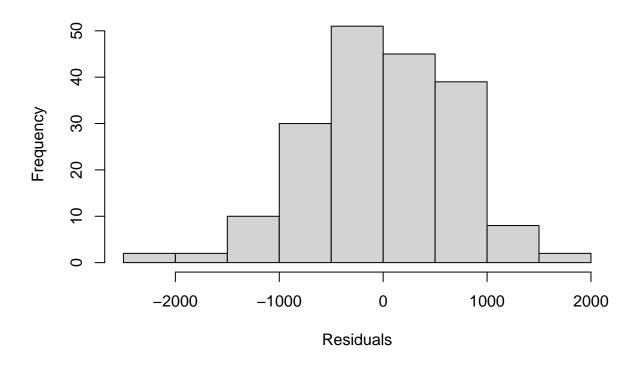
The adjusted R-squared value is just above the F statistic, and it is 0.07464.

The "Coefficients:" table has the variable name in the first column, then the estimated  $\hat{\beta}_i$ , the standard error, the test statistic value for the partial test, and the p-value for the partial tests.

The quantiles for the residuals are also in the summary, but we can get a histogram of the residuals by giving the hist() function the residuals of the bwt.lm linear model object, which we get with the residuals() function.

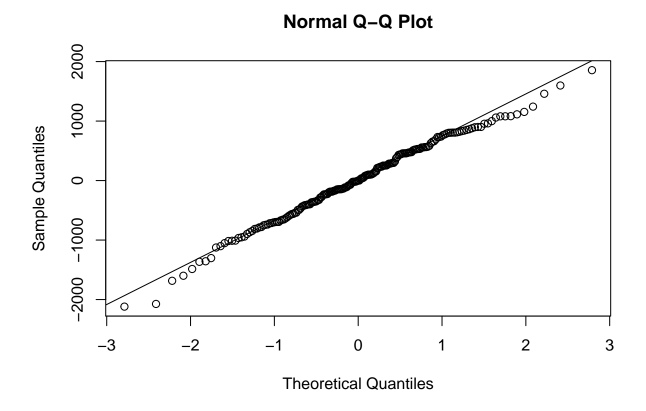
```
hist(residuals(bwt.lm), main="Histogram of Residuals", xlab="Residuals")
```

# **Histogram of Residuals**



We can also get the normal probability plot with the qqnorm() function, then the qqline() function to add the normal probability line to the plot.

```
qqnorm(residuals(bwt.lm))
qqline(residuals(bwt.lm))
```



You can export plots in Rstudio to put them in a report by clicking the "export" button just above the plot.