Lab 4: Healthy Momma, Healthy Baby

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A Nice Introduction that Makes Us Sound Like Pros

According to the NIH, having a healthy pregancy is one of the best ways to promote a healthy birth and that getting early and regular prenatal care improves the chances of a healthy pregnancy.[1] According to Hack et all, while most low birth weight children will end up having normal outcomes, as a group they generally have more health issues than healthy weight babies[2].

Using data from the National Center for Health Statistics and from birth certificates, we will look at the impact of prenatal health care on health outcomes for newborn infants.

According to Montgomery, the Apgar scores are used as an evaluative measure to see if a newborn needs immediate attention. However, the using Apgar scores to attempt to predict long-term developmental outcomes of infants in not appropriates, so we will not be using Apgar scores in our outcome variable for newborn health. [3]

Therefore we will use birthweight as our outcome variable for our analysis based on historical research because of the limitations of our dataset.

Something about higher birthweight that talks about neural development big babies, big brains

Step 1: Read in the Data

```
load('/Users/nicholeh/student285/w203/w203_lab_4/bwght_w203.RData')
desc
```

```
##
      variable
                                            label
## 1
          mage
                             mother's age, years
## 2
         meduc
                           mother's educ, years
## 3
        monpre
                      month prenatal care began
## 4
         npvis total number of prenatal visits
          fage
## 5
                             father's age, years
## 6
         feduc
                            father's educ, years
## 7
         bwght
                             birth weight, grams
## 8
         omaps
                          one minute apgar score
## 9
                         five minute apgar score
         fmaps
## 10
          cigs
                          avg cigarettes per day
                             avg drinks per week
## 11
         drink
## 12
           lbw
                             =1 if bwght <= 2000
## 13
          vlbw
                             =1 if bwght <= 1500
## 14
                                 =1 if baby male
          male
## 15
         mwhte
                              =1 if mother white
## 16
         mblck
                              =1 if mother black
## 17
                           =1 if mother is other
          moth
## 18
         fwhte
                              =1 if father white
## 19
         fblck
                              =1 if father black
## 20
          foth
                           =1 if father is other
## 21
        lbwght
                                      log(bwght)
## 22
        magesq
                                           mage<sup>2</sup>
```

Step 2: Exploratory Data Analysis

First, get summary statistics on each element of the dataset:

nrow(data)

[1] 1832

summary(data)

```
##
                          meduc
         mage
                                           monpre
                                                            npvis
##
    Min.
            :16.00
                     Min.
                             : 3.00
                                              :0.000
                                                        Min.
                                                                : 0.00
##
    1st Qu.:26.00
                     1st Qu.:12.00
                                       1st Qu.:1.000
                                                        1st Qu.:10.00
##
    Median :29.00
                     Median :13.00
                                       Median :2.000
                                                        Median :12.00
##
    Mean
            :29.56
                                                                :11.62
                     Mean
                             :13.72
                                       Mean
                                              :2.122
                                                        Mean
##
    3rd Qu.:33.00
                     3rd Qu.:16.00
                                       3rd Qu.:2.000
                                                        3rd Qu.:13.00
            :44.00
##
    Max.
                     Max.
                             :17.00
                                       Max.
                                              :9.000
                                                        Max.
                                                                :40.00
##
                     NA's
                             :30
                                       NA's
                                              :5
                                                        NA's
                                                                :68
##
                          feduc
                                           bwght
         fage
                                                           omaps
##
    Min.
            :18.00
                     Min.
                             : 3.00
                                       Min.
                                               : 360
                                                       Min.
                                                               : 0.000
                     1st Qu.:12.00
                                                       1st Qu.: 8.000
##
    1st Qu.:28.00
                                       1st Qu.:3076
                     Median :14.00
    Median :31.00
                                       Median:3425
                                                       Median: 9.000
##
##
    Mean
            :31.92
                     Mean
                             :13.92
                                       Mean
                                               :3401
                                                       Mean
                                                               : 8.386
    3rd Qu.:35.00
                     3rd Qu.:16.00
                                       3rd Qu.:3770
                                                       3rd Qu.: 9.000
##
            :64.00
                             :17.00
                                               :5204
                                                               :10.000
    Max.
                     Max.
                                       Max.
                                                       Max.
##
    NA's
            :6
                     NA's
                             :47
                                                       NA's
                                                               :3
##
        fmaps
                            cigs
                                             drink
                                                                 lbw
##
    Min.
           : 2.000
                              : 0.000
                                         Min.
                                                 :0.0000
                                                           Min.
                                                                   :0.00000
                      Min.
##
    1st Qu.: 9.000
                      1st Qu.: 0.000
                                         1st Qu.:0.0000
                                                           1st Qu.:0.00000
##
    Median : 9.000
                      Median : 0.000
                                                           Median :0.00000
                                         Median :0.0000
##
    Mean
           : 9.004
                      Mean
                              : 1.089
                                         Mean
                                                 :0.0198
                                                           Mean
                                                                   :0.01638
##
    3rd Qu.: 9.000
                                         3rd Qu.:0.0000
                      3rd Qu.: 0.000
                                                           3rd Qu.:0.00000
##
    Max.
            :10.000
                      Max.
                              :40.000
                                         Max.
                                                 :8.0000
                                                           Max.
                                                                   :1.00000
            :3
                              :110
##
    NA's
                      NA's
                                         NA's
                                                 :115
##
         vlbw
                              male
                                               mwhte
                                                                  mblck
##
    Min.
            :0.000000
                        Min.
                                :0.0000
                                           Min.
                                                   :0.0000
                                                              Min.
                                                                     :0.0000
    1st Qu.:0.000000
                         1st Qu.:0.0000
##
                                           1st Qu.:1.0000
                                                              1st Qu.:0.0000
##
                         Median :1.0000
                                                              Median :0.0000
    Median :0.000000
                                           Median :1.0000
    Mean
            :0.007096
                         Mean
                                :0.5136
                                           Mean
                                                   :0.8865
                                                              Mean
                                                                     :0.0595
##
    3rd Qu.:0.000000
                        3rd Qu.:1.0000
                                           3rd Qu.:1.0000
                                                              3rd Qu.:0.0000
##
    Max.
            :1.000000
                        Max.
                                :1.0000
                                           Max.
                                                   :1.0000
                                                              Max.
                                                                     :1.0000
##
##
         moth
                            fwhte
                                              fblck
                                                                   foth
##
    Min.
            :0.00000
                        Min.
                               :0.0000
                                          Min.
                                                  :0.00000
                                                              Min.
                                                                     :0.00000
##
    1st Qu.:0.00000
                        1st Qu.:1.0000
                                          1st Qu.:0.00000
                                                              1st Qu.:0.00000
##
    Median :0.00000
                        Median :1.0000
                                          Median :0.00000
                                                              Median :0.00000
##
    Mean
            :0.05404
                        Mean
                               :0.8897
                                          Mean
                                                  :0.05841
                                                                     :0.05186
                                                              Mean
##
    3rd Qu.:0.00000
                        3rd Qu.:1.0000
                                          3rd Qu.:0.00000
                                                              3rd Qu.:0.00000
            :1.00000
                               :1.0000
##
                                                  :1.00000
    Max.
                       Max.
                                          Max.
                                                              Max.
                                                                     :1.00000
##
##
        lbwght
                          magesq
                                           npvissq
##
    Min.
            :5.886
                     Min.
                             : 256.0
                                        Min.
                                              :
                                                    0.0
    1st Qu.:8.031
                     1st Qu.: 676.0
                                        1st Qu.: 100.0
```

```
## Median :8.139 Median : 841.0 Median : 144.0

## Mean :8.114 Mean : 896.4 Mean : 148.6

## 3rd Qu.:8.235 3rd Qu.:1089.0 3rd Qu.: 169.0

## Max. :8.557 Max. :1936.0 Max. :1600.0

## NA's :68
```

Response Variables

The bwght, lbwght, omaps and fmaps variables are related to the health of the baby.

```
The bwght, bwght, omaps and imaps variables are related to the health of the baby.

The first thing to check is if these variables are collinar. We will omit bwghts as that is a function of lbwghts.

library(ggplot2)
cor(data$maps, data$fmaps, use = "complete.obs")

## [1] 0.5575238

cor(data$lbwght, data$fmaps, use = "complete.obs")

## [1] 0.2710456

p <- ggplot(data, aes(omaps, lbwght)) + geom_point(size = 0.25) + geom_smooth(method = "lm", se = FALSE) + geom_point(aes(colour = fmaps)) + ggtitle("Scatterplot of log(weight) against One Minute APGAR test,\n with 5 minute APGAR test heatmap")

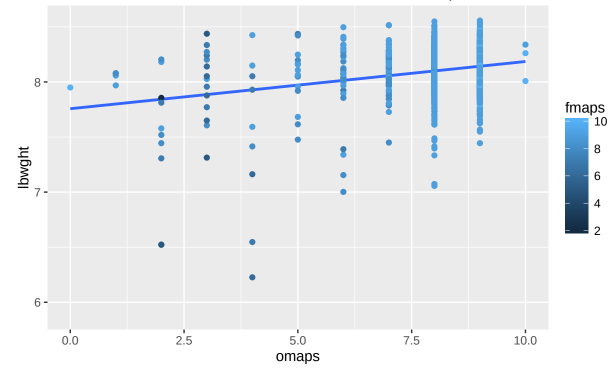
p

## Warning: Removed 3 rows containing non-finite values (stat_smooth).

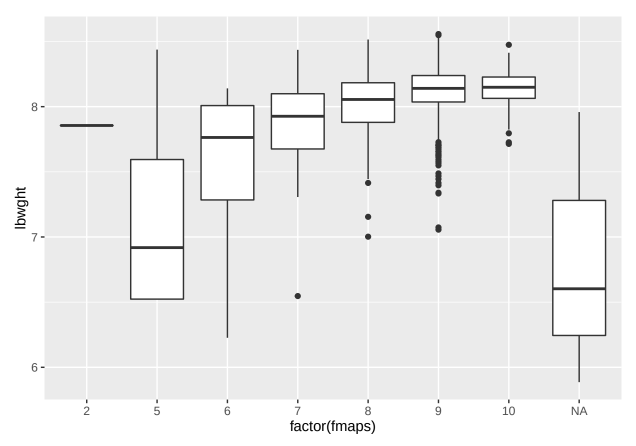
## Warning: Removed 3 rows containing missing values (geom_point).
```

Scatterplot of log(weight) against One Minute APGAR test,





p <- ggplot(data, aes(factor(fmaps), lbwght)) + geom_boxplot()</pre>



Look at the extreme fmaps case

data[data\$fmaps< 4,]</pre>

##		mage	meduc	monpre	npvis	fage	feduc	bwght	oma	ps	fmaps	cigs	drink	lbw
##	NA	NA	NA	NA	NA	NA	NA	NA		NA	NA	NA	NA	NA
##	837	32	12	2	10	40	16	2580		2	2	0	0	0
##	NA.1	NA	NA	NA	NA	NA	NA	NA		NA	NA	NA	NA	NA
##	NA.2	NA	NA	NA	NA	NA	NA	NA		NA	NA	NA	NA	NA
##		vlbw	male i	mwhte ml	olck m	oth f	whte f	blck f	oth	1	bwght	mages	q npv:	issq
##	NA	NA	NA	NA	NA	NA	NA	NA	NA		NA	N.	A	NA
##	837	0	1	1	0	0	1	0	0	7.8	355545	102	4	100
##	NA.1	NA	NA	NA	NA	NA	NA	NA	NA		NA	N.	A	NA
##	NA.2	NA	NA	NA	NA	NA	NA	NA	NA		NA	N.	A	NA

Looking at the data, we can be reasonably assured that the response variables are related, but not collinear. It may be best to make a combined variable of fmaps and omaps such as mapscombined = fmaps + omaps. The difference would not make much sense compared to the sum; 10 - 10 and 2 - 2 are both zero, after all.

Regressors

The variables monpre and npvis are related to the prenatal care given during pregnancy. Let us review them for collinearity:

```
cor(data$npvis, data$monpre, use = "complete.obs")
```

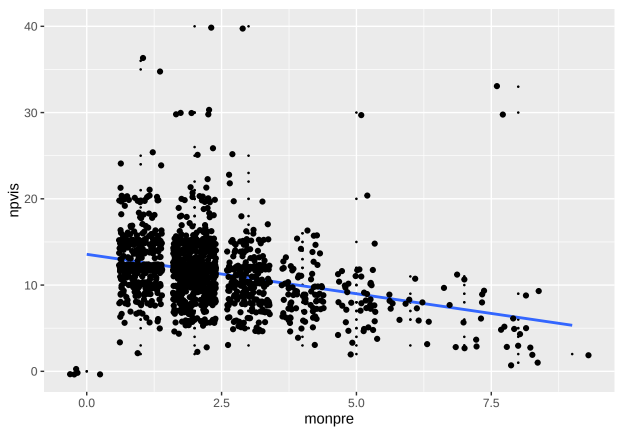
[1] -0.3061006

```
ggplot(data, aes(monpre, npvis)) + geom_point(size = 0.25) +
geom_smooth(method = "lm", se = FALSE) + geom_jitter()
```

Warning: Removed 69 rows containing non-finite values (stat_smooth).

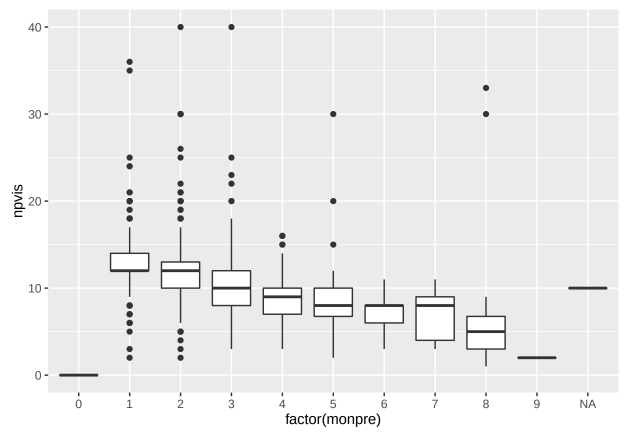
Warning: Removed 69 rows containing missing values (geom_point).

Warning: Removed 69 rows containing missing values (geom_point).



ggplot(data, aes(factor(monpre), npvis)) + geom_boxplot()

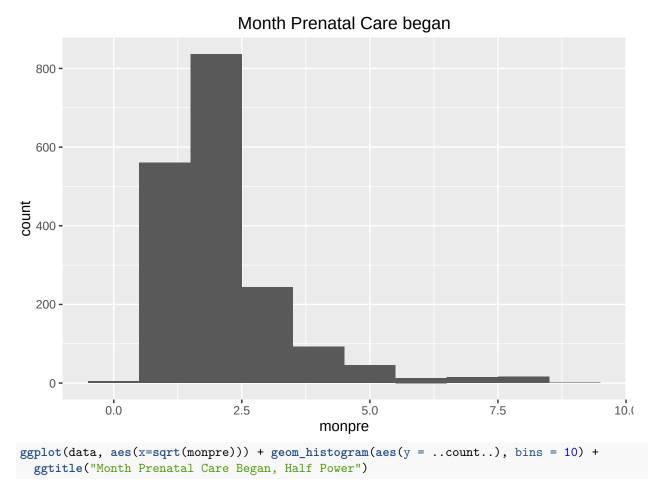
Warning: Removed 68 rows containing non-finite values (stat_boxplot).



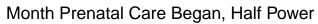
From this set, we can see that the data is not collinear, and indeed we can see that we might have some reporting errors. 5 mothers are listed as starting prenatal care in month 0 of their pregnancy, but they visited the doctor 0 times. These probably denote missing information or an error in reporting. Unfortunately, this data does show a definitive downward trend leading us to suspect that the number of visits is a function of month prenatal care began. This makes sense intuitively; if a mother starts prenatal care in her 2nd month of pregnancy, she has ample time for frequent doctor visits. However, if she starts her prenatal care towards the end of her pregnancy, she does not have enough time to visit the doctor as often as a woman who started in month 2.

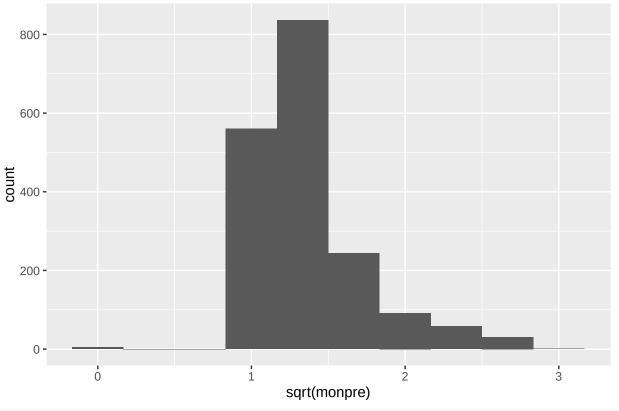
```
ggplot(data, aes(x=monpre)) + geom_histogram(aes(y = ..count..),bins = 10) +
ggtitle("Month Prenatal Care began")
```

Warning: Removed 5 rows containing non-finite values (stat_bin).



Warning: Removed 5 rows containing non-finite values (stat_bin).

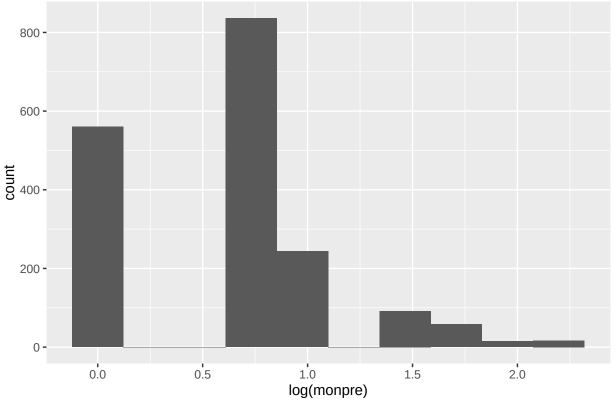




```
ggplot(data, aes(x=log(monpre))) + geom_histogram(aes(y = ..count..), bins = 10) +
   ggtitle("Month Prenatal Care Began, Natural Log")
```

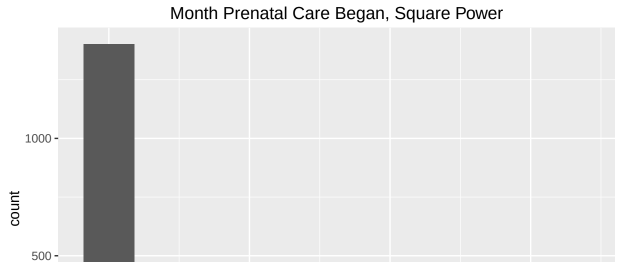
Warning: Removed 10 rows containing non-finite values (stat_bin).





```
ggplot(data, aes(x=(monpre^2))) + geom_histogram(aes(y = ..count..), bins = 10) +
 ggtitle("Month Prenatal Care Began, Square Power")
```

Warning: Removed 5 rows containing non-finite values (stat_bin).



(monpre^2)

ggplot(data, aes(x=npvis)) + geom_histogram(aes(y = ..count..), bins = 15) +
 ggtitle("Number of Prenatal Visits")

50

7₅

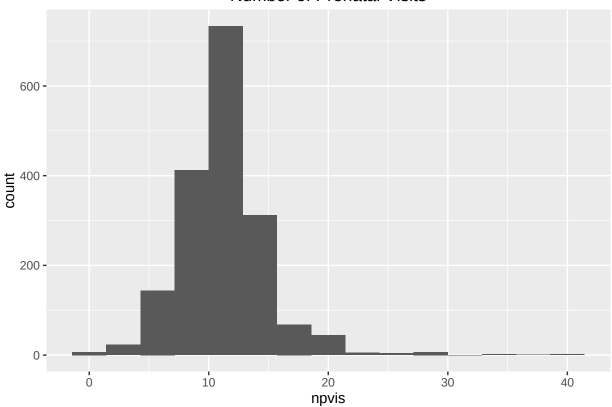
Warning: Removed 68 rows containing non-finite values (stat_bin).

25

0 -

Ö

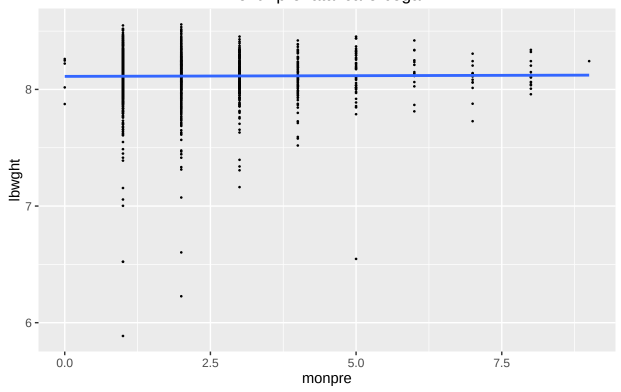
Number of Prenatal Visits



```
ggplot(data, aes(monpre, lbwght)) + geom_point(size = 0.25) +
geom_smooth(method = "lm", se = FALSE) +
ggtitle("Scatterplot of weight against \n month prenatal care began ")
```

- ## Warning: Removed 5 rows containing non-finite values (stat_smooth).
- ## Warning: Removed 5 rows containing missing values (geom_point).

Scatterplot of weight against month prenatal care began



All in all, the number of visits follows a mostly normal curve, and the square root of the month prenatal care began follow a mostly normal curve. However, we can tell right now that monpre does not have much practical significance with respect to the baby's weight from looking at the graph.

Step 3: Modeling

Model 1: Basic Linear Model

```
model1<-lm(bwght ~ monpre + npvis, data = data)
summary(model1)$r.squared</pre>
```

[1] 0.01123524

6 CLM assumptions:

- 1) Linearity in parameters: We can assume this.
- 2) Random sampling of data: Not random because are not including still births or miscarriages.
- 3) No perfect co-linearity

```
cor(data$monpre, data$npvis, use="complete.obs")
```

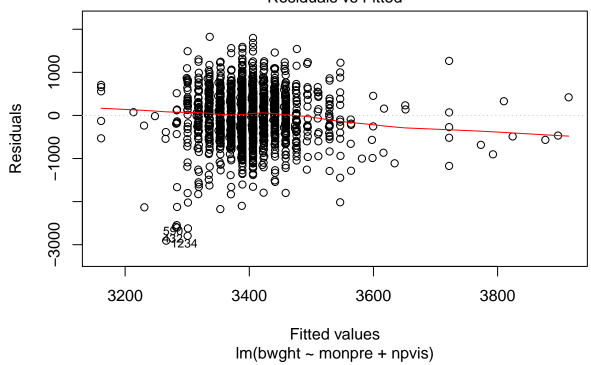
[1] -0.3061006

There is no perfect multicolineraity between our variables. With a correlation of -0.3061006, this shows that the number of prenatal visits is moderately negatively correlated to the month in which prenatal care started.

4) Zero conditional mean

plot(model1, which=1)

Residuals vs Fitted



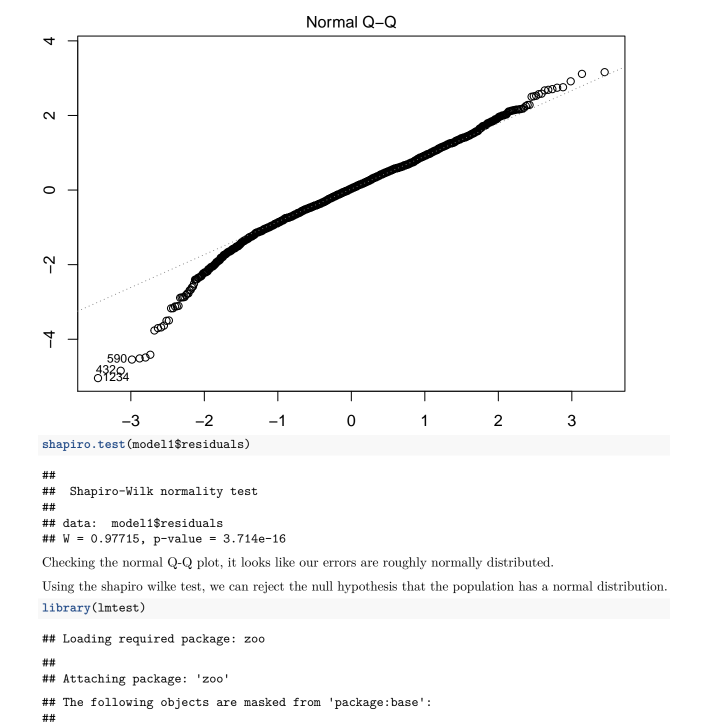
Looking at the Residuals vs. Fitted plot shows that the zero conditional mean is met because the red line is approximately at 0.

5) Homoskedacity of errors

From the residuals vs. fitted plot, we can see that we do not have homoskedacity of erorrs because the data is not in an even band across the plot. This means that we'll have to white standard errors, which are roboust to heteroskadacity.

6) Errors are normally distributed

```
par(mar = rep(2, 4))
plot(model1, which=2)
```



```
##
## t test of coefficients:
##
## Estimate Std. Error t value Pr(>|t|)
## (Intercept) 3161.2707   74.6049 42.3735 < 2.2e-16 ***</pre>
```

as.Date, as.Date.numeric

coeftest(model1, vcov = vcovHC)

##

library(sandwich)

```
## monpre 17.0622 12.0277 1.4186 0.1561984
## npvis 17.5494 4.8342 3.6302 0.0002913 ***
## ---
## Signif. codes: 0 '*** 0.001 '** 0.05 '.' 0.1 ' ' 1
```

Model 2: An Alternate Main Model

The 1 minute and 5 minute APGAR scores on their own do not tell us much. As we can see from the heatmap on the first scatterplot, a baby who has a low one minute score tends to have a higher five minute score. There are very few examples of a baby having a worse five minute score than a one minute score:

```
nrow(data[!is.na(data$fmaps) < !is.na(data$omaps),])</pre>
```

```
## [1] 3
```

However, we can get some information if we take the product of omaps and fmaps and then normalize it. A baby that goes from 0 to 10 then would have an overal low score compared to a baby who started with a score of 10 and was still at 10 5 minutes later, so the difference doesn't make sense.

```
data$product_apgarscores = data$omaps * data$fmaps
data$normalized_product_apgar =
    (data$product_apgarscores -
        mean(!is.na(data$product_apgarscores)))/sd(!is.na(data$product_apgarscores))
a8 = lm(data$normalized_product_apgar~data$monpre + data$npvis)
a9 = lm(data$normalized_product_apgar~ data$npvis)
AIC(a8)
```

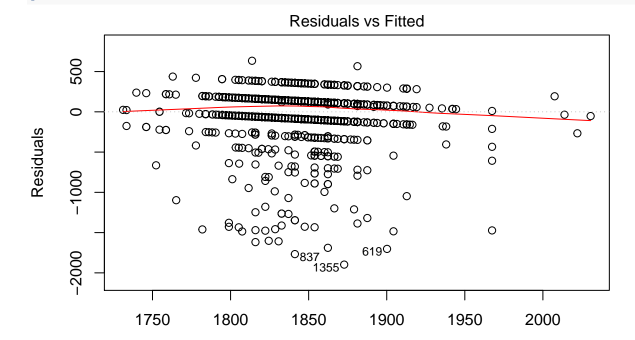
```
## [1] 24885.48
AIC(a9)
```

[1] 24899.94

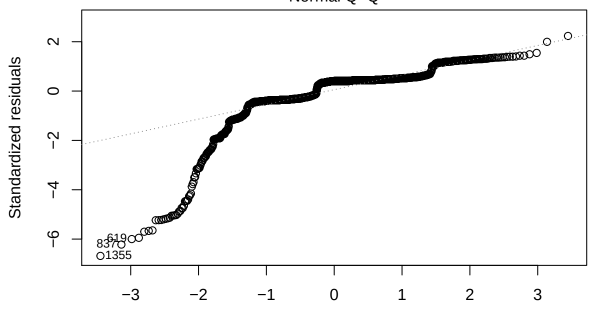
Model a8 has a nominally lower AIC score, so let's continue on with that one.

summary(a8)

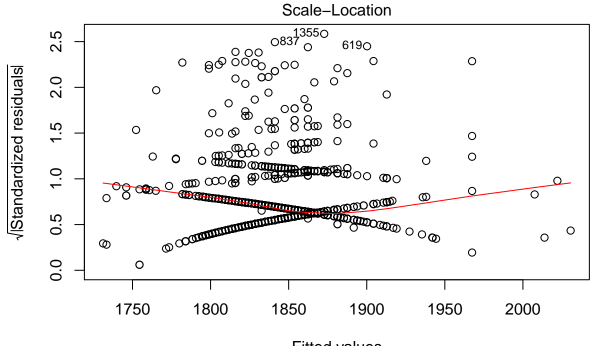
```
##
## Call:
## lm(formula = data$normalized_product_apgar ~ data$monpre + data$npvis)
##
## Residuals:
##
       Min
                  1Q
                       Median
                                    3Q
                                            Max
                       115.74
## -1897.44
             -98.29
                                130.55
                                         634.08
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 1795.067
                            29.566
                                    60.713 < 2e-16 ***
## data$monpre
                 -8.502
                             5.774
                                    -1.472 0.14107
## data$npvis
                  6.313
                             1.936
                                     3.261 0.00113 **
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 284.1 on 1757 degrees of freedom
     (72 observations deleted due to missingness)
## Multiple R-squared: 0.00981,
                                    Adjusted R-squared: 0.008683
```



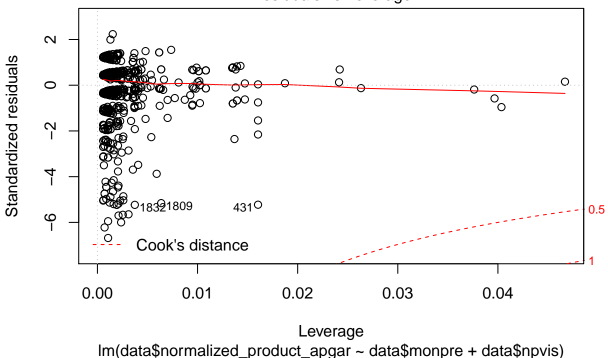
Fitted values
Im(data\$normalized_product_apgar ~ data\$monpre + data\$npvis)
Normal Q-Q



Theoretical Quantiles Im(data\$normalized_product_apgar ~ data\$monpre + data\$npvis)



Fitted values
Im(data\$normalized_product_apgar ~ data\$monpre + data\$npvis)
Residuals vs Leverage



We did not see very good results with the APGAR score variations, but as discussed in the introduction, we were expecting the baby's birth weight would have a better indication.

6 CLM assumptions:

- 1) Linearity in parameters: We can assume this.
- 2) Random sampling of data: This data is not random because stillbirths are omitted.

3) No perfect co-linearity

As previously stated, our regressors do not have perfect collinearity.

4) Zero conditional mean

Looking at the Residuals vs. Fitted plot above shows that the zero conditional mean is met because the red line is approximately at 0 and has very little curvature.

5) Homoskedacity of errors

From the residuals vs. fitted plot, we can see that we do not have homoskedacity of errors because the data is not in an even band across the plot. This means that we'll have to use white standard errors, which are roboust to heteroskadacity.

6) Errors are normally distributed

```
par(mar = rep(2, 4))
shapiro.test(a8$residuals)

##
## Shapiro-Wilk normality test
##
## data: a8$residuals
## W = 0.71096, p-value < 2.2e-16</pre>
```

From normal Q-Q plot, it looks like our errors are roughly normally distributed except at the very highest and very lowest percentiles. This is to be expected in a dataset such as this.

Using the shapiro wilke test, we can reject the null hypothesis that the population has a normal distribution.

```
library(lmtest)
library(sandwich)
coeftest(a8, vcov = vcovHC)
##
## t test of coefficients:
##
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 1795.0673
                           36.0086 49.8510 < 2e-16 ***
## data$monpre
                -8.5024
                            5.7237 -1.4855 0.13760
## data$npvis
                 6.3128
                            2.5166 2.5084 0.01222 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Model 3: Unbiased Covariants

```
model3<-lm(bwght ~ monpre + npvis + cigs + drink + mage + male, data = data)</pre>
```

- 6 CLM assumptions:
 - 1) Linearity in parameters: We can assume this.
 - 2) Random sampling of data: This data is not random because stillbirths are omitted.
 - 3) No perfect co-linearity: As previously stated, our regressors do not have perfect collinearity.

```
## cigs
           0.09905318 -0.03736714 1.00000000 0.185567975 -0.061323113
                                                            0.004413966
## drink
          -0.01031974 0.05263935
                                   0.18556797
                                               1.000000000
          -0.19911595
## mage
                      0.09649250 -0.06132311
                                               0.004413966
                                                           1.000000000
          -0.01868132 -0.02185506 -0.01102578 -0.047648827 -0.039928312
## male
##
                 male
## monpre -0.01868132
## npvis
          -0.02185506
          -0.01102578
## cigs
## drink
          -0.04764883
          -0.03992831
## mage
## male
           1.0000000
```

4) Zero conditional mean

plot(model3, which=1)

Residuals vs Fitted 1000 0 0 0 Residuals 0 800 -10000 0 0 0 3000 3200 3400 3600 3800 Fitted values Im(bwght ~ monpre + npvis + cigs + drink + mage + male)

ing at the Residuals vs. Fitted plot shows that the zero conditional mean is met because the red line is approximately at 0.

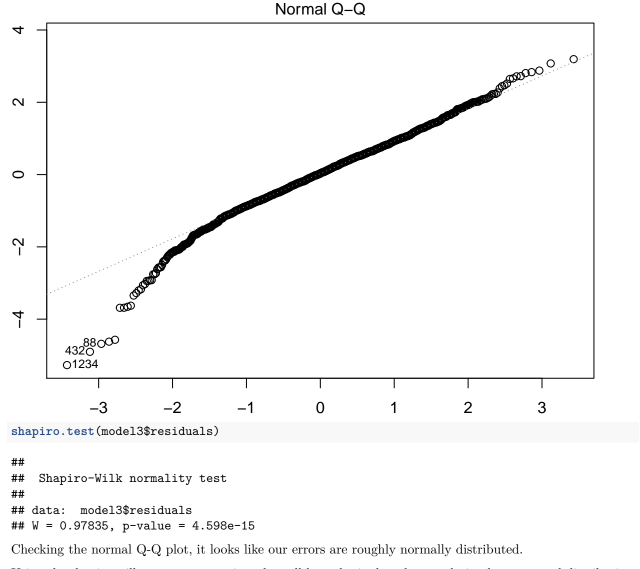
Look-

5) Homoskedacity of errors

From the residuals vs. fitted plot, we can see that we do not have homoskedacity of erorrs because the data is not in an even band across the plot. This means that we'll have to white standard errors, which are roboust to heteroskadacity.

6) Errors are normally distributed

```
par(mar = rep(2, 4))
plot(model3, which=2)
```



Using the shapiro wilke test, we can reject the null hypothesis that the population has a normal distribution.

```
coeftest(model1, vcov=vcovHC)
```

##

```
##
## t test of coefficients:
##
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 3161.2707
                           74.6049 42.3735 < 2.2e-16 ***
## monpre
                17.0622
                           12.0277 1.4186 0.1561984
                17.5494
                            4.8342 3.6302 0.0002913 ***
## npvis
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
coeftest(model3, vcov=vcovHC)
##
## t test of coefficients:
##
```

Estimate Std. Error t value Pr(>|t|)

```
## (Intercept) 2999.6248
                          124.7351 24.0480 < 2.2e-16 ***
                           12.0531 1.7341 0.083091 .
## monpre
                20.9010
## npvis
                            4.6619 3.3258 0.000901 ***
                15.5046
               -11.2291
                            3.6793 -3.0520 0.002310 **
## cigs
## drink
               -14.0495
                           33.0106 -0.4256 0.670451
                 5.3168
                            3.1399 1.6933 0.090592 .
## mage
## male
                80.9374
                           28.2671 2.8633 0.004246 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
AIC(model1)
## [1] 27428.8
AIC(model3)
## [1] 25582.07
```

Model 4: Problematic Covariants

We will select the attributes of baby's gender and parent's race as well. In the United States, it is a sad fact that minorities such as African Americans do not have adequate access to proper health care as often as non-minorities. Their babies might not fare as well, and their mothers may not get the proper prenatal care.

From all of the summaries, we can tell that the t-statistic for the monpre variable is not significant. Thus, we cannot trust this particular regressor, and will omit it from this test.

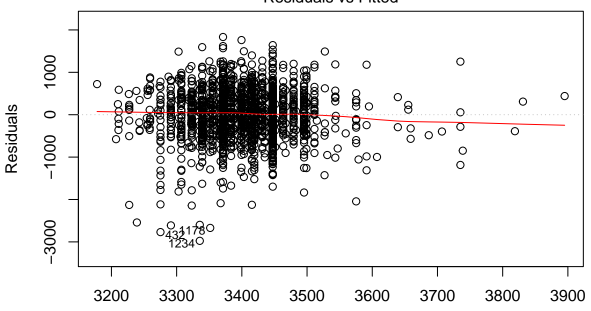
```
##
## Call:
## lm(formula = data$bwght ~ data$npvis + data$male + data$mblck +
##
       data$fblck)
##
## Residuals:
##
       Min
                 1Q
                      Median
                                    3Q
                                            Max
## -2975.51 -336.55
                       31.69
                               360.92
                                       1832.85
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 3179.315
                           48.188 65.977 < 2e-16 ***
## data$npvis
                15.986
                            3.735
                                    4.280 1.97e-05 ***
## data$male
                76.262
                           27.534
                                     2.770 0.00567 **
## data$mblck
               -97.221
                          126.174
                                   -0.771 0.44109
## data$fblck
                48.729
                          127.179
                                    0.383 0.70166
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 576.7 on 1759 degrees of freedom
     (68 observations deleted due to missingness)
## Multiple R-squared: 0.01479,
                                   Adjusted R-squared: 0.01255
## F-statistic: 6.6 on 4 and 1759 DF, p-value: 2.857e-05
```

AIC(c1)

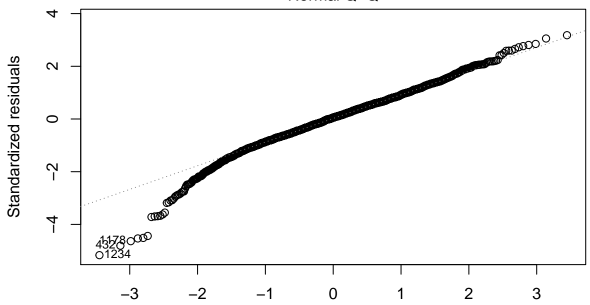
[1] 27441.54

plot(c1)

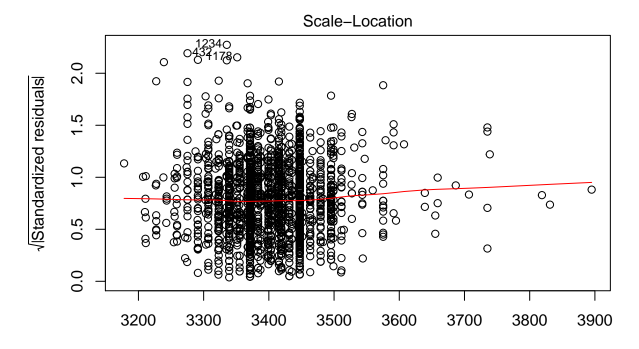




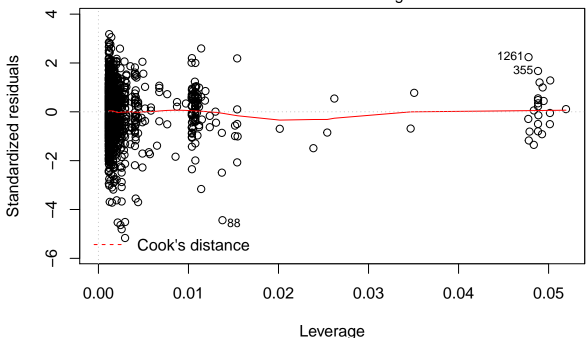
Fitted values
Im(data\$bwght ~ data\$npvis + data\$male + data\$mblck + data\$fblck)
Normal Q-Q



Theoretical Quantiles
Im(data\$bwght ~ data\$npvis + data\$male + data\$mblck + data\$fblck)



Fitted values
Im(data\$bwght ~ data\$npvis + data\$male + data\$mblck + data\$fblck)
Residuals vs Leverage



Im(data\$bwght ~ data\$npvis + data\$male + data\$mblck + data\$fblck)

6 CLM assumptions:

- 1) Linearity in parameters: We can assume this.
- 2) Random sampling of data: This data is not random because stillbirths are omitted.
- 3) No perfect co-linearity in regressors:

```
cor(data[,c('npvis', 'mblck', 'fblck', 'male')], use="complete.obs")
```

```
##
              npvis
                           mblck
                                       fblck
                                                    male
## npvis 1.00000000 -0.03379275 -0.03133149 -0.02635585
## mblck -0.03379275
                     1.00000000
                                 0.88963736
                                              0.04743914
## fblck -0.03133149
                     0.88963736
                                  1.00000000
                                              0.02402644
## male -0.02635585 0.04743914 0.02402644
                                              1.00000000
```

As previously stated, our regressors do not have perfect collinearity.

4) Zero conditional mean

Looking at the Residuals vs. Fitted plot above shows that the zero conditional mean has not been met because the red line shows curvature for larger babies.

5) Homoskedacity of errors

From the residuals vs. fitted plot, we can see that we do not have homoskedacity of erorrs because the data is not in an even band across the plot. This means that we'll have to use white standard errors, which are roboust to heteroskadacity.

6) Errors are normally distributed

```
par(mar = rep(2, 4))
shapiro.test(c1$residuals)
```

```
##
## Shapiro-Wilk normality test
##
## data: c1$residuals
## W = 0.97639, p-value < 2.2e-16</pre>
```

From normal Q-Q plot, it looks like our errors are roughly normally distributed except at the very lowest percentiles. This is to be expected in a dataset such as this.

Using the shapiro wilke test, we can reject the null hypothesis that the population has a normal distribution.

```
library(lmtest)
library(sandwich)
coeftest(c1, vcov = vcovHC)
```

```
##
## t test of coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) 3179.3155
                           56.2484 56.5228 < 2.2e-16 ***
## data$npvis
                 15.9863
                            4.3518
                                    3.6735 0.0002464 ***
## data$male
                76.2618
                            27.4392
                                    2.7793 0.0055056 **
## data$mblck
                -97.2213
                          121.8744 -0.7977 0.4251425
## data$fblck
                 48.7286
                          118.4882 0.4113 0.6809371
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

As we were hoping with such biased data, we can see that the race of the parents is not statistically significant so it is inappropriate to include it in our model.

Step 4: CLM and the Models

Step 5: Regression Tables and Model Analysis

% Table created by stargazer v.5.2 by Marek Hlavac, Harvard University. E-mail: hlavac at fas.harvard.edu % Date and time: Fri, Dec 09, 2016 - 20:29:42

See table 1 on the next page.

```
AIC(model1)

## [1] 27428.8

AIC(model3)

## [1] 25582.07

AIC(c1)
```

[1] 27441.54

From the Akaike Information Criterion test, we see that model3 is the best option for a linear model predicting the health of the baby. Model3 has the highest adjusted R^2, showing that virutally 2% of all variability in the baby's health indicators can be determined by the months prenatal visits started, number of prenatal visits, the mother's smoking and driking habits, the mother's age, and the baby's gender. As always, monpre was not a statistically significant regressor, and neither was the mother's age or drinking habits. In words, we can say if the baby is a boy we can expect he will weigh 80.937 grams more than if he is a girl, for every year older his mother is, he will weight 5.317 grams more, for every alcoholic drink his mother inbibes per week he will weigh 14.050 grams less, for every cigarette his mother smokes per day, he will weigh 11.229 grams less, for each prenatal visit, he will weight 15.505 more, and for each month the mother waits to to start her prenatal care, the baby weight 20.901 grams more. Just writing it out what the model means stresses even more that we should ignore the monpre variable in modeling a baby's health.

Table 1:

	Dependent variable:							
	bwght	normalized_product_apgar	bwght	bwght				
	(1)	(2)	(3)	(4)				
monpre	17.062 (12.028)		20.901 (12.053)					
npvis	17.549*** (4.834)		15.505*** (4.662)					
monpre		-8.502 (5.724)						
npvis		$6.313^* $ (2.517)		15.986*** (4.352)				
cigs			-11.229** (3.679)					
drink			-14.050 (33.011)					
mage			5.317 (3.140)					
male			80.937** (28.267)					
male				76.262** (27.439)				
mblck				-97.221 (121.874)				
fblck				48.729 (118.488)				
Constant	3,161.271*** (74.605)	1,795.067*** (36.009)	2,999.625*** (124.735)	3,179.315*** (56.248)				
Observations	1,763	1,760	1,647	1,764				
\mathbb{R}^2	0.011	0.010	0.023	0.015				
Adjusted R ² Residual Std. Error	$0.010 \\ 577.470 \text{ (df} = 1760)$	0.009 284.115 (df = 1757)	0.019 569.408 (df = 1640)	0.013 576.683 (df = 1759)				

Note:

*p<0.05; **p<0.01; ***p<0.001

Step 6: Causality

We choose to operationalize infant health by birthweight. There are many other factors that influence birthweight that are not captured in this data set, which leads to omitted variable bias.

- 1) Mother's weight is a strong predictor for newborn weight.
- 2) Socioeconomic status of mother.
- 3) Having more than one baby at a time reduces the weight of each baby. (E.g. twins will be smaller)

4)

Biases and Limitation

This data is extremely biased in that no still births were included in our dataset. It is a sad fact in the United States that over 2 in 1,000 births are stillbirths[5]. Since we do not know the prenatal care data for stillbirths, we cannot completely guage how much prenatal care contributes to a child's health at birth.

In addition, it appears that there is little correlation between the Appar score and the later health of the baby. The Apar is only meant to be used in the context of emergency situations. In this manner, looking at a baby's weight will give us deeper insight into the baby's overall health.

No miscarriages were included in the data, so this further biases our data.

Using birthweight as a proxy for infant health was the best that we could do given our data set, but is by no means a comprehensive view on an infants' health.

Step 7: Conclusion

Prenatal care, as shown by number of prenatal care visits has a positive impact on birthweight. Other explanatory factors are mother's cig consumption, which has a negative impact on birthweight. Being male has a positive impact on birthweight.

References

[1]https://www.nichd.nih.gov/health/topics/pregnancy/conditioninfo/pages/prenatal-care.aspx

[2]https://www.ncbi.nlm.nih.gov/pubmed/7543353

[3] https://www.ncbi.nlm.nih.gov/pmc/articles/PMC1595023/~[4] http://ije.oxfordjournals.org/content/30/6/1233.long

[5] https://www.washingtonpost.com/news/wonk/wp/2014/09/29/our-infant-mortality-rate-is-a-national-embarrassment/? utm term=.58 dedfd178 fd