MTH-245 Final project Part 2 Fall 2022

Name:

```
library("tidyverse")
library("ggplot2")
library("patchwork")
library("bestglm")
library("EnvStats")
library("car")
library("caret")
library("gGally")
library("gGart")
library("olsrr")
library("gridExtra")
library("boot")
source("https://cipolli.com/students/code/plotResiduals.R")
```

1 Abstract

Background: According to The Washington Post, the average birth weight of American infants has dropped 453.592 grams between 1990 and 2013, making the average birth weight 3247.721 grams. While, this drop in weight may not seem significant, it brings us closer to an average low birth weight which is classified as 2,500 grams or less. Stanford University released a study on how low birth weights can impose health issues on children. Such issues include infection, breathing problems and immature lungs, nervous system problems, bleeding inside the brain, sudden infant death syndrome, and other long term complications such as cerebral palsy, blindness, deafness, developmental delay. Clearly these are extremely high risks and the same study from Stanford listed some social factors of the mothers that influence birth weight such as smoking, not gaining enough weight during pregnancy, African-American background, and the age of the mother being less than 17 or more than 35 years. Awareness of what social factors that influence low birth weights can help with prenatal guidance and care to avoid the potential risks listed above. The purpose of this study is to identify key predictors influencing low birth weights using a sample of infant birth weights and other information collected from North Carolina. Methods: We will use a linear regression to model the relationship between whether the mother was a smoker, weight gained by the mother, the mother's age, ... and the infant's birth weight. Findings: After making adjustments to the initial model such as transformations, centering, and interactions, we determined our final best model to predict which factors have the most influence on birth weight. Our final model had *an 83.6 increase in precision and a 7 increase in predictive ability as compared to the first order additive linear model*.

2 Introduction

According to The World Health Organization, the average weight of a baby born at 37–40 weeks ranges from 5 lb 8 oz to 8 lb 13 oz. This is 2,500 grams to 4,000 grams. Birth weight is something that we don't typically consider when we are projecting the health of our future population, but it plays an extremely important role in influencing the expectancy, quality and health of a person's life. If an infant is born with a low birth weight, they could face immediate and long term health issues. If a child is inflicted with long term health issues, they will require medical care and resources for the rest of their lives. These considerations are important population-wise, because as the population grows and the birth weight continues to decrease, there may be a strain on medical care and some resources available to those with long term health issues. Those who work in the healthcare industry regarding women and children's health, especially Obstetricians, should be informed on what social factors and behaviors within the population strongly influence birth weight so that they can provide the correct medical care and advice for each patient, accordingly. Our data is called NCbirths and comes from the Stat2Data package in R datasets. It was collected by statistician John Holcomb at Cleveland State University, from the North Carolina State Center for Health and Environmental Statistics. NCbirths contains data from births in North Carolina in 2001, with 1450 observations on 15 variables that include social and behavioral characteristics of the mother. The response variable

of our study was BirthWeightGM, which is the baby's birth weight in grams. We hypothesized that the following variables would be the most predictive, after our background research using the Low Birth Weight study published by Stanford University: race of mother, gestation period (weeks), sex of the infant, whether just a single infant was delivered or more than one, if the mother smoked while pregnant, weight gained by mother, the mother's age. We hypothesize that gestation period will be very influential, but our study will determine which other variables are influential. The following code imports our data and alters the type of each variable. We also renamed the levels within our categorical variables, and treated them all as factors. We centered and scaled all of our quantitative variables and created more variables for each transformation conducted on the quantitative variables.

```
prepData <- function() {</pre>
  births <- read_csv("~/GitHub/Mth245Final/dataset/NCbirths.csv")</pre>
  births <- births %>% mutate(Sex = case_when(Sex == 1 ~ "Male",
                                               Sex == 2 ~ "Female"),
                               Marital = case_when(Marital == 1 ~ "Married",
                                                    Marital == 2 ~ "Unmarried"),
                               RaceMom = case_when(RaceMom == 1 ~ "White",
                                                    RaceMom == 2 "Black",
                                                    RaceMom == 3 ~ "Am. Indian",
                                                    RaceMom == 4 ~ "Chinese",
                                                    RaceMom == 5 ~ "Japanese",
                                                    RaceMom == 6 ~ "Hawaiian",
                                                    RaceMom == 7 ~ "Filipino",
                                                    RaceMom == 8 ~ "Other Asian / PI"),
                               Smoke = case_when(Smoke == 1 ~ "Yes",
                                                  Smoke == 0 \sim "No"),
                               Preemie = case_when(Premie == 1 ~ "Yes",
                                                  Premie == 0~ "No"))
  births$Sex <- as.factor(births$Sex)</pre>
  births$Marital <- as.factor(births$Marital)</pre>
  births$Preemie <- as.factor(births$Preemie)</pre>
  births$Smoke <- as.factor(births$Smoke)</pre>
  births$RaceMom <- as.factor(births$RaceMom)</pre>
  births$Plural <- as.factor(births$Plural)</pre>
  births <- births %>% mutate(MomAgeSC = scale(MomAge, center=T, scale=T),
                               MomAgeSq = MomAgeSC<sup>2</sup>,
                               WeeksSC = scale(Weeks, center=T, scale=T),
                               WeeksSq = WeeksSC ^2,
                               GainedSC = scale(Gained, center=T, scale=T),
                               GainedSq = I(GainedSC<sup>2</sup>))
  births <- births %>% filter(!is.na(GainedSC) & !is.na(Smoke))
  # Part 1: First-Order Model and Determinations of Necessary Transformations
  births <- births %>% mutate(WeightGmLog = log(BirthWeightGm))
  births <- births %>% mutate(WeightGmSqrt = BirthWeightGm^.5)
  births <- births %>% mutate(WeightGmS = BirthWeightGm^2)
  births <- births %>% mutate(WeightGmSLog = log(BirthWeightGm)^2)
  births <- births %>% mutate(WeightGmLogLog = log(log(BirthWeightGm)))
  births <- births %>% mutate(WeightGmLogSqr = log(BirthWeightGm^2))
  births <- births %>% mutate(WeightGmLogQuad = log(BirthWeightGm)^4)
  births <- births %>% mutate(WeightGmSqrtLog = log(BirthWeightGm)^.5)
  births <- births %>% mutate(WeightGmInverse = 1/(BirthWeightGm))
  births <- births %>% mutate(WeightGmSC = scale(BirthWeightGm, center=T, scale=T))
  births <- births %>% mutate(WeightLogSC = scale(WeightGmLog, center=T, scale=T))
```

```
births <- births %>% mutate(Twin = (as.character(Plural) == "2"))
  births <- births %>% mutate(Triplet = (as.character(Plural) == "3"))
  births <- births %>% mutate(Filipino = (RaceMom == "Filipino"))
  births <- births %>% mutate(Black = (RaceMom == "Black"))
  births$Twin = as.factor(births$Twin)
  births$Triplet = as.factor(births$Triplet)
  births$Fllipino = as.factor(births$Filipino)
  births$Black = as.factor(births$Black)
  births
births <- prepData()</pre>
## Rows: 1450 Columns: 15
## -- Column specification
## Delimiter: ","
## chr (2): HispMom, MomRace
## dbl (13): ID, Plural, Sex, MomAge, Weeks, Marital, RaceMom, Gained, Smoke, B...
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.
```

3 Exploratory Data Analysis

a. Graphically summarize the variables in your dataset.

First we visualized our quantitative variables, including our response variable.

And we specifically created a boxplot for the birth weights recorded for mothers who are Black, since our research from the Stanford study indicated that race is influential in birth weight.

```
violin.BirthWeightGm <- ggplot(births, aes(x=BirthWeightGm, y=""))+</pre>
  geom_violin(fill = "lightblue",
              trim = FALSE)+
  geom_boxplot(width = .3,
               fill = "white") +
  theme_bw()+
  xlab("Birth Weights")+
  ylab(" ")+
  ggtitle("Distribution of Birth Weights",
          subtitle = "NCBirths Data")
violin.GestationPeriod <- ggplot(births, aes(x=Weeks, y=""))+</pre>
  geom_violin(fill = "lightblue",
             trim = FALSE)+
  geom_boxplot(width = .3,
               fill = "white") +
  theme_bw()+
  xlab("Weeks")+
  ylab(" ")+
  ggtitle("Distribution of Gestation Period",
     subtitle = "NCBirths Data")
```

Figure 1: Violin plots of each variable.

the shape of the distribution of each variable any unusual looking observations

From 1 We can see that there is variability in almost all of the quantitative variables and they all have and many odd observations.

We also wanted to visualize the distributions of these variables.

```
histogram.BirthWeight<- ggplot(births, aes(x=BirthWeightGm))+
  geom_histogram(fill = "lightblue",
                 color = "black",
                 bins = 5) +
  theme_bw() +
  xlab("Birth Weights")+
  vlab("Count of Weight(gm)")+
  ggtitle("Frequencies of Birth Weights")
histogram.Gestation <- ggplot(births, aes(x=Weeks))+
  geom_histogram(fill = "lightblue",
                 color = "black",
                 bins = 5) +
  theme_bw() +
  xlab("Gestation Period")+
  ylab("Count of Weeks")+
  ggtitle("Frequencies of Gestation Periods")
histogram.MomAge <- ggplot(births, aes(x=MomAge))+
  geom_histogram(fill = "lightblue",
                 color = "black",
                 bins = 5) +
  theme_bw() +
  xlab("Ages of Mothers(years)")+
  ylab("Count of Ages")+
  ggtitle("Frequencies of Ages")
histogram.BirthWeight + histogram.Gestation + histogram.MomAge
```

Figure 2: Grid of histograms for the quantitative variables.

2 shows that the quantitative variables do not follow normal distributions and are all skewed.

We expected the distribution for weights and weeks to be similar in shape because typically babies that are born prematurely have low birth weights. We did find it interesting though, that the distributions showed more preemies that we initially expected.

We weren't sure if the preemie weights in the data set were outlier instances that were heavily skewing the distribution, or if there were just more preemies that we expected. To further investigate this, we created a bootstrap confidence interval to see whether a baby born with a median weight falls within the interval.

```
## Bootstrapping for median weights
median(births$BirthWeightGm)

set.seed(23)
alpha <- 0.05
n <- nrow(births)
R <- 10000
boot.stats <- rep(NA, R)
for (i in 1:R){
  boot.data <- sample(x = births$BirthWeightGm, size = n, replace = TRUE)
  boot.stats[i] <- median(boot.data)
}

quantile(boot.stats, probs = c(alpha/2, 1 - alpha/2))

samp.boot.med <- function(data, indicies){
  median(data[indicies])
}

boot.medians <- boot(data = births$BirthWeightGm, statistic = samp.boot.med, R = 10000)
boot.ci(boot.medians, conf = 0.95)</pre>
```

The median birth weight of our dataset was 3,345.3 grams. Our 95% percentile confidence interval was 3316.95 g. to 3373.65 g. This range contained our median birth weight of 3,345.3 grams, but more than that, our confidence interval was extremely close to the reported statistics from the World Health Organization of median birth weight of 3.3 kg (The WHO did not include any additional significant figures, so we assume that this could range from values anywhere to 3.25 kg to 3.35 kg.). This seems to lend support to our decision to assume that the collected data was representative of the population, despite the lack of specific knowledge of the collection techniques. Further, it indicates to us that we have a reasonable sample even with its non-normal distribution, that was caused by its skew from data primarily relating to babies born prematurely.

b. Numerically summarize the variables in your dataset.

Tableu visual

c. Create a scatterplot matrix and table of correlations.

```
library(GGally)
#ADD IN CORRECT VARIABLE NAMES
```

```
correlationsmatrix <- ggpairs(births, columns = c())
correlationsmatrix</pre>
```

Figure 3: Matrix of ScatterPlots and Correlations for the variables.

d. Other interesting plots.

Plot significant correlations - weeks and birth weight

e. Comment on...

the shape of the distribution of each variable

the relationship between the response and the quantitative predictors

any unusual looking observations

any other interesting takeaways

Before we began with our first order model, we eliminated Low, a catagorical predictor that indicates true if the birth weight was low and false if the birth weight was not low. This attribute is originally calculated off of birth weight, which is our response variable. Excluding this predictor from the model eliminates any skew or strong false influence in our model from data that we could not know without having already had the response variable provided to us.

4 First-Order Model and Model Selection.

Before we began generating models, we first built several functions that would make analysis of the data more straightforward. One of these was to measure a statistic we devised and termed the "quantile departure" (QD) statistic. After scaling the residuals and sorting them in order, it measures how much the actual quantile positioning of each residual in a model departs from the theoretical quantiles of the model, calculated using q-norm. The value it returns is equivalent to:

$$\frac{\sum_{i=1}^{n} |R_i - T_i|}{n}$$

where R_i is equal to the i^{th} lowest actual residual and T_i is the i^{th} lowest theoretical residual. A higher value would indicate a higher departure from the theoretical quantiles, while a lower value would indicate residuals more consistent with the theoretical quantiles.

```
#Create functions
#Calculates the residuals departure from the theoretical quantiles, returning the mean abs() value of the
quantDepart <- function(model) {
   residuals <- sort(scale(model$residuals, scale=T))
   i <- 1:length(residuals)
   fi <- (i - 0.5) / length(residuals)
      x.norm <- qnorm(fi)

   mean(abs(residuals - x.norm))
}
#Displays a variety of summary stats and graphics of the model tailored based off of the specific options</pre>
```

```
modelSummary <- function(model, coef=T, stat=T, plot=T){</pre>
    print(round(summary(model)$coefficients,10))
 if(stat) {
    print(paste("R-squared:", summary(model)$r.squared))
    print(paste("Adjusted R-Squared:", summary(model)$adj.r.squared))
    print(paste("RMSE:", summary(model)$sigma))
    print(paste("AIC:", AIC(model)))
    print(paste("BIC:", BIC(model)))
    print(paste("Quantile Departure:", quantDepart(model)))
  if(plot) {
   plotResiduals(model)
#Calculates the R squared of predicted vs actual values
r_squared <- function(actual, predicted) {</pre>
  cor(actual, predicted)^2
#If provided with a model and a number of subsets of data, this will generate summary statistics for those
predictForSubsets <- function(model, class.attr, ..., names=c()) {</pre>
 x <- list(...)
  i <- 1
  residPlots <- list()</pre>
  predPlots <- list()</pre>
  for (v in x) {
    v$predict <- predict(model, v)</pre>
    v$resid <- v[[class.attr]] - v$predict
    subsetName <- ""
    if (length(names) >= i) {
      subsetName <- names[i]</pre>
    } else {
      subsetName <- paste("Subset", i)</pre>
    print(paste(subsetName, "R-Squared:", r_squared(v$predict, v[[class.attr]])))
    print(paste(subsetName, "Mean Abs. Error:", mean(abs(v$resid))))
    ggplot(data=v, aes(x=predict, y=resid)) +
      geom_point(size=1,
                 shape=16)+
      theme_bw()+
      xlab("Predicted")+
      vlab("Residuals")-> residuals
    ggplot(data=v, aes(x=predict, y=get(class.attr))) +
      geom_point(size=1,
                 shape=16)+
      theme_bw()+
      xlab("Predicted")+
      vlab("Actual") -> predictions
    if (length(x) != 3) {
      print((predictions + ggtitle("Predicted versus Actual",
           subtitle=subsetName)) + (residuals + ggtitle("Predicted versus Residuals",
           subtitle=subsetName)))
```

4.1 First Order Model

Fitting a first-order linear model with all our acceptable predictor variables, we calculated the following estimated linear regression equation:

Table 1: Summary of first order regression model R-squared, Adj R-squared, RSE, AIC, and BIC.

As Table ?? indicates, Plural2 (Twins), Plural3 (Triplets), Sex = Male, MomAge, Weeks, RaceMom = Filipino, Gained, Smoke, and Preemie are all significant predictors. Table 1 demonstrates that the R^2 is fairly high, meaning the model is quite predictive; additionally, the quantile departure value is very close to 0, meaning it does well to satisfy the requirements for equal distribution of residuals. This is further emphasized in Figure 4. However, AIC and BIC values are also high, meaning that the number of predictor variables is not predictive enough to justify the variety of parameters we are using.

```
Marital + GainedSC + Smoke + Preemie, births) -> model.sqrt
# Squared of Weight
lm(WeightGmS ~ Plural + Sex + MomAgeSC + WeeksSC + RaceMom +
     Marital + GainedSC + Smoke + Preemie, births) -> model.s
# Square of the Log of Weight
lm(WeightGmSLog ~ Plural + Sex + MomAgeSC + WeeksSC + RaceMom +
     Marital + GainedSC + Smoke + Preemie, births) -> model.slog
# Log of the Log of Weight
lm(WeightGmLogLog ~ Plural + Sex + MomAgeSC + WeeksSC + RaceMom +
     Marital + GainedSC + Smoke + Preemie, births) -> model.loglog
# Log of the Square Root of Weight
lm(WeightGmLogSqr ~ Plural + Sex + MomAgeSC + WeeksSC + RaceMom +
     Marital + GainedSC + Smoke + Preemie, births) -> model.logsqr
# Square Root of the Log of Weight
lm(WeightGmSqrtLog ~ Plural + Sex + MomAgeSC + WeeksSC + RaceMom +
     Marital + GainedSC + Smoke + Preemie, births) -> model.sqrtlog
# Inverse of the Weight
lm(WeightGmInverse ~ Plural + Sex + MomAgeSC + WeeksSC + RaceMom +
     Marital + GainedSC + Smoke + Preemie, births) -> model.inverse
```

Before eliminating the non-significant variables from our model we wanted to test transformations of BirthWeight, to see if any of them improved our model performance statistics or residual distributions. The model statistics for each transformation are listed in Table ??.

Transformation	R^2	R_{adj}^2	RSE	AIC	BIC	QD
No Transformation	0.4570	0.4512	464.7	21322	21412	0.0472
log of Weight	0.5243	0.5192	0.1756	-885.3	-796	0.1573
Square Root of Weight	0.4994	0.4940	4.3160	8137	8227	0.0787
Weight Squared	0.3699	0.3631	$3.04*10^{6}$	46086	46175	0.0974
log of Weight Squared	0.5208	0.5156	2.7	6812	6901	0.1340
log of the log of Weight	0.5249	0.5198	0.0231	-6597	-6508	0.1824
log of the Square Root of Weight	0.5243	0.5192	0.3512	1068	1157	0.1573
Square Root of the <i>log</i> of Weight	0.5250	0.5199	0.0318	-5699	-5610	0.1695
Inverse of the Weight	0.4507	0.4447	0.0001	-21462	-21373	0.3578

Table 2: Summary of all adjusted first order regression model R-squared, Adj R-squared, RSE, AIC, and BIC.

Based on Table 2, we have several potential transformation options. First is the un-transformed weight, which better satisfies the assumptions behind linear regression modeling, namely the equal distribution of residuals. On the other hand, we have the log weight model, which has higher R^2 values and lower RMSE, AIC, and BIC values than most other models, meaning it is more accurate. This accuracy is, however, gained only at the expense of making residuals less equally distributed. As shown by the QD values, and emphasized by the comparisons between Figure 4 and 5, the log transformation results in a quantile departure nearly four times the size of the non-transformed. The residual disparities also seem to come much more from the lower end of the birth weights in the logarithmic model. Because of this, it does not seem to as strongly satisfy the assumptions necessary to do things like conduct statistical inference on parameters. Thus, we will be proceeding with two models, the assumptions model and the accuracy model, where one will seek to minimize departure from assumptions and the other will seek to maximize accuracy. Assumption models will use the scaled and centered gram weight as their target, while accuracy models will use the scaled and centered log of gram weight.

Figure 4: Residual Plots of Untransformed Model.

Figure 5: Residual Plots of Logarithmic Model.

Below are our two first-order models, one with no transformations and the other with the *log* of the weight taken. Both have all numeric variables, including the response variables, scaled and centered.

4.2 First Order Model Pruning

Our next step was to reduce the number of variables in the two models to only those that had an affect. For both of our models, we ran AIC-based elimination, BIC-based elimination, and regression subset optimization techniques to determine which variables should be removed.

```
x <- model.matrix(model.2.assu)[,-1]
## Error in model.matrix(model.2.assu): object 'model.2.assu' not found
y <- births$WeightGmSC

xy <- as.data.frame(cbind(x,y))
## Error in cbind(x, y): object 'x' not found
best.subsets.aic <- bestglm(xy, IC="AIC", TopModels = 5)
## Error in is.data.frame(Xy): object 'xy' not found
best.model.aic <- best.subsets.aic$BestModel
## Error in eval(expr, envir, enclos): object 'best.subsets.aic' not found
modelSummary(best.model.aic, plot=F)
## Error in summary(model): object 'best.model.aic' not found
best.subsets.bic <- bestglm(xy, IC="BIC", TopModels = 5)
## Error in is.data.frame(Xy): object 'xy' not found
best.model.bic <- best.subsets.bic$BestModel</pre>
```

```
## Error in eval(expr, envir, enclos): object 'best.subsets.bic' not found
modelSummary(best.model.bic, plot=F)
## Error in summary(model): object 'best.model.bic' not found
regsubsets.out <- regsubsets(WeightGmSC ~ Plural + Sex + MomAgeSC + WeeksSC + RaceMom +
                            Marital + GainedSC + Smoke + Preemie,
                           data=births, nbest = 1, nvmax=15)
fit.stats <- data.frame(num.variables=1:15,</pre>
                      adjr2 = summary(regsubsets.out)$adjr2,
                      bic=summary(regsubsets.out)$bic)
cbind(fit.stats, as.data.frame(summary(regsubsets.out)$outmat))
           num.variables
                            adjr2 bic Plural2 Plural3 SexMale MomAgeSC
## 1 (1)
                      1 0.3419165 -576.0584
## 2 (1)
                      2 0.3659468 -622.2229
## 3 (1)
                      3 0.3936644 -678.9560
## 4 (1)
                      4 0.4139286 -720.6032
## 5 (1)
                      5 0.4262316 -744.2497
## 6 (1)
                      6 0.4342432 -757.8164
## 7 (1)
                     7 0.4397681 -765.3984
## 8 (1)
                      8 0.4453965 -773.3810
## 9 (1)
                      9 0.4508074 -780.9513
## 10 (1)
                     10 0.4518552 -777.3989
## 11 ( 1 )
                      11 0.4524400 -772.6604
## 12 ( 1 )
                     12 0.4523523 -766.1931
## 13 (1)
                     13 0.4519692 -758.9668
## 14 ( 1 )
                     14 0.4515905 -751.7534
## 15 (1)
                      15 0.4512188 -744.5593
##
            WeeksSC RaceMomBlack RaceMomChinese RaceMomFilipino RaceMomJapanese
## 1 ( 1 )
## 2 (1)
## 3 (1)
## 4 ( 1 )
## 5 (1)
## 6 (1)
## 7 (1)
## 8 (1)
## 9 (1)
## 10 (1)
## 11 ( 1 )
## 12 ( 1 )
## 13 (1)
## 14 ( 1 )
## 15 (1)
##
            RaceMomOther Asian / PI RaceMomWhite MaritalUnmarried GainedSC
## 1 (1)
## 2 (1)
## 3 (1)
## 4 (1)
## 5 (1)
## 6 (1)
## 7 (1)
## 8 (1)
## 9 (1)
```

Figure 6: "Residual Plots of Pruned Non-Transformed Model"

```
??
```

```
## 10
     (1)
     (1)
## 11
     (1)
## 12
     (1)
## 13
## 14
     (1)
## 15 (1)
##
           SmokeYes PreemieYes
## 1
     (1)
## 2
     (1)
## 3
     (1)
## 4
     (1)
## 5
     (1)
## 6
     (1)
## 7
     (1)
## 8
     (1)
## 9
     (1)
## 10 (1)
## 11
     (1)
      (1)
## 12
## 13
      (1)
## 14
     (1)
## 15
     (1)
```

For the assumption model, we found that the AIC-based technique, the BIC-based technique, and the regression subset technique all agreed that all racial attributes except for Black and Filipino should be removed. BIC also recommended removing the Marital measure and the Filipino measure. We decided to remove all of the variables recommended for removal by the algorithms; the decision to exclude Filipino was fairly easy, as only a single value in our dataset was classified as Filipino. We also were confident in removing Marital because the effect size was extremely low and the significance levels quite high in the AIC-based model that had included it. We thus ended up with the following model:

xtable(model.3.assu, caption="Predictor Statistics and Significance for Pruned First-Order Non-Transformed

	Estimate	Std. Error	t value	$\Pr(> t)$
(Intercept)	0.0948	0.0337	2.81	0.0050
Twin	-1.1201	0.1220	-9.18	0.0000
Triplet	-1.4721	0.3785	-3.89	0.0001
Male	0.1522	0.0396	3.85	0.0001
MomAge (SC)	0.1186	0.0204	5.81	0.0000
Weeks (SC)	0.4187	0.0292	14.35	0.0000
Black	-0.1917	0.0481	-3.98	0.0001
Gained (SC)	0.1738	0.0200	8.70	0.0000
Smoke	-0.3343	0.0564	-5.92	0.0000
Preemie	-0.3577	0.0848	-4.22	0.0000

Table 3: Predictor Statistics and Significance for Pruned First-Order Non-Transformed Model

In our new model, according to Table 3, all remaining variables have coefficients with p-values < 0.05, indicating

that the values for β_i all fall either above or below 0. R^2 sees a slight decrease, but R^2_{adj} is now closer to R^2 , as $R^2 - R^2_{adj}$ goes from 0.0058 to 0.0035. Further, according to Figure ??, our residuals have remained roughly constant, compared to 4, and our quantile departure is approximately the same.

```
x <- model.matrix(model.2.accur)[,-1]
## Error in model.matrix(model.2.accur): object 'model.2.accur' not found
y <- births$WeightLogSC
xy <- as.data.frame(cbind(x,y))</pre>
## Error in cbind(x, y): object 'x' not found
best.subsets.aic <- bestglm(xy, IC="AIC", TopModels = 5)</pre>
## Error in is.data.frame(Xy): object 'xy' not found
best.model.aic <- best.subsets.aic$BestModel</pre>
## Error in eval(expr, envir, enclos): object 'best.subsets.aic' not found
modelSummary(best.model.aic)
## Error in summary(model): object 'best.model.aic' not found
best.subsets.bic <- bestglm(xy, IC="BIC", TopModels = 5)</pre>
## Error in is.data.frame(Xy): object 'xy' not found
best.model.bic <- best.subsets.bic$BestModel</pre>
## Error in eval(expr, envir, enclos): object 'best.subsets.bic' not found
modelSummary(best.model.bic)
## Error in summary(model): object 'best.model.bic' not found
regsubsets.out <- regsubsets(WeightLogSC ~ Plural + Sex + MomAgeSC + WeeksSC + RaceMom +
                              Marital + GainedSC + Smoke + Preemie,
                            data=births, nbest = 1, nvmax=15)
fit.stats <- data.frame(num.variables=1:15,
                       adjr2 = summary(regsubsets.out)$adjr2,
                       bic=summary(regsubsets.out)$bic)
cbind(fit.stats, as.data.frame(summary(regsubsets.out)$outmat))
            num.variables
                              adjr2
                                          bic Plural2 Plural3 SexMale MomAgeSC
## 1 (1)
                       1 0.4419739 -808.4394
## 2 (1)
                        2 0.4600565 -848.6047
## 3 (1)
                       3 0.4817581 -900.1570
                       4 0.4980496 -938.9143
## 5 (1)
                       5 0.5048386 -951.8545
## 6 (1)
                        6 0.5112379 -963.9368
## 7 (1)
                       7 0.5152965 -969.4404
## 8 (1)
                       8 0.5180355 -971.1805
## 9 (1)
                       9 0.5190927 -968.0308
## 10 (1)
                       10 0.5197597 -963.7431
## 11 ( 1 )
                       11 0.5201610 -958.6786
## 12 (1)
                       12 0.5201212 -952.3202
## 13 ( 1 )
                       13 0.5198713 -945.3456
## 14 ( 1 )
                       14 0.5195516 -938.1675
## 15 ( 1 )
                       15 0.5192084 -930.9220
            WeeksSC RaceMomBlack RaceMomChinese RaceMomFilipino RaceMomJapanese
##
```

```
## 1 (1)
## 2 (1)
## 3
    (1)
## 4
    (1)
## 5
    (1)
## 6
    (1)
## 7
    (1)
## 8
    (1)
## 9
    (1)
## 10 (1)
## 11
     (1)
## 12
     (1)
## 13
     (1)
## 14 ( 1 )
## 15 (1)
##
           RaceMomOther Asian / PI RaceMomWhite MaritalUnmarried GainedSC
## 1 (1)
## 2 (1)
    (1)
## 3
## 4
    (1)
## 5
    (1)
## 6
    (1)
## 7
    (1)
    (1)
## 8
## 9
    (1)
## 10 (1)
## 11
     (1)
## 12 (1)
## 13 (1)
## 14 ( 1 )
## 15
     (1)
##
           SmokeYes PreemieYes
## 1 (1)
## 2 (1)
    (1)
## 4
    (1)
## 5
    (1)
    (1)
## 6
## 7
    (1)
    (1)
## 8
## 9
    (1)
## 10 (1)
## 11 (1)
## 12 (1)
## 13 (1)
## 14
     (1)
## 15 (1)
```

Just as with the model for the non-transformed weights, the same variables - all levels of race except for Black, as well as marital status - were recommended to be removed in the model with the log transformation of the weights. Having done so, in Table 4, we found that all remaining variables were significant with p-values < 0.05.

```
model.3.accur <- lm(WeightLogSC ~ Twin + Triplet + Sex + MomAgeSC + WeeksSC + Black +
GainedSC + Smoke + Preemie, births)

xtable(model.3.accur, caption="Predictor Statistics and Significance for Pruned First-Order Log-Transforme
```

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	0.0671	0.0315	2.13	0.0336
Twin	-1.0119	0.1141	-8.87	0.0000
Triplet	-1.5454	0.3542	-4.36	0.0000
Male	0.1100	0.0370	2.97	0.0030
MomAge (SC)	0.1120	0.0191	5.86	0.0000
Weeks (SC)	0.5553	0.0273	20.33	0.0000
Black	-0.1582	0.0450	-3.51	0.0005
Gained (SC)	0.1523	0.0187	8.15	0.0000
Smoke	-0.2547	0.0528	-4.82	0.0000
Preemie	-0.1602	0.0793	-2.02	0.0436

Table 4: Predictor Statistics and Significance for Pruned First-Order Log-Transformed Model

```
modelSummary(model.3.assu, coef=F, plot=F)
```

Just as before, our R^2 values decreased as we removed values, but our R^2_{adj} actually remained exactly the same, going from 0.5191 to 0.5192.

Using these two models, we decided to conduct run cross validation tests to make sure that the data was not simply being overfit, and that our \mathbb{R}^2 and MAE values were accurate to the data as a whole. We ran cross validation on both models, using both k-fold cross validation and leave-one-out cross validation.

```
specs <- trainControl(method="LOOCV")</pre>
model.cval.LOO.accur <- train(WeightLogSC ~ Twin + Triplet + Sex + MomAgeSC + WeeksSC + Black +
                      GainedSC + Smoke + Preemie, data=births,
                      method = "lm",
                      trControl = specs,
                      na.action = na.omit)
model.cval.L00.accur$results[["Method"]] <- "L00CV"</pre>
specs <- trainControl(method="CV", number=10)</pre>
model.cval.cv.accur <- train(WeightLogSC ~ Twin + Triplet + Sex + MomAgeSC + WeeksSC + Black +
                      GainedSC + Smoke + Preemie, data=births,
                      method = "lm",
                      trControl = specs,
                      na.action = na.omit)
model.cval.cv.accur$results[["Method"]] <- "K-Fold"</pre>
rbind(model.cval.LOO.accur$results,
      model.cval.cv.accur$results %>% select(intercept, RMSE, Rsquared, MAE, Method)) %>%
  select(Method, RMSE, Rsquared, MAE) %>%
 xtable(caption="Comparison of Cross-Validation Techniques on Logarithmic Model", label="cross.val.log")
```

Method	RMSE	R^2	MAE
None	0.69	0.52	0.49
LOOCV	0.70	0.51	0.50
K-Fold	0.69	0.49	0.50

Table 5: Comparison of Cross-Validation Techniques on Logarithmic Model

```
trControl = specs,
                      na.action = na.omit)
model.cval.L00.assu$results[["Name"]] <- "L00CV"</pre>
specs <- trainControl(method="CV", number=10)</pre>
model.cval.cv.assu <- train(WeightGmSC ~ Twin + Triplet + Sex + MomAgeSC + WeeksSC + Black +
                      GainedSC + Smoke + Preemie, data=births,
                      method = "lm",
                      trControl = specs,
                      na.action = na.omit)
model.cval.cv.assu$results[["Name"]] <- "K-Fold"</pre>
rbind(model.cval.LOO.assu$results,
      model.cval.cv.assu$results %>% select(intercept, RMSE, Rsquared, MAE, Name)) %>%
    select(RMSE, Rsquared, MAE, Name) %>%
  xtable(caption="Comparison of Cross-Validation Techniques", label="cross.val.un")
## % latex table generated in R 4.2.1 by xtable 1.8-4 package
## % Thu Dec 8 01:53:43 2022
## \begin{table}[ht]
## \centering
## \begin{tabular}{rrrrl}
    \hline
## & RMSE & Rsquared & MAE & Name \\
    \hline
##
## 1 & 0.74 & 0.45 & 0.58 & LOOCV \\
     2 & 0.74 & 0.45 & 0.58 & K-Fold \\
##
      \hline
## \end{tabular}
## \caption{Comparison of Cross-Validation Techniques}
## \label{cross.val.un}
## \end{table}
```

Method	RMSE	R^2	MAE
Orig.	0.74	0.45	0.57
LOOCV	0.74	0.45	0.58
K-Fold	0.74	0.44	0.58

Table 6: Comparison of Cross-Validation Techniques

In both Table 10 and 6, it is apparent that the cross validation yields values of RMSE, R^2 , and MAE that are directly in line with the values we calculate using the full training data.

After obtaining our new model, we were curious as to how some of the predictors influenced the data set. Premature births means that the baby was born weeks before the predicted due date, and its commonly known that "Preemies" have low birth weights. They are also some of the more important subject to have accurate weight predictions of, because of their often-delicate health condition and the importance that greater levels of development can bring with it. We calculated the residuals of our models for the entire dataset, just the premature births, and just the non-premature births, and created plots comparing the distribution of residuals for each of the subsets of the data.

As Figures 7 and 8 demonstrate, both model tend to have higher residual departures for premature births than they do for the dataset as a whole or for the subset of non-premature births. This is also reflected in the MAE values, as seen in Table 7, where the premature birth subset is always lower than the non-premature birth or the full dataset. However, Table 7 also shows that premature births also have a much higher correlation with the

Figure 7: Comparison of Residual Distributions Between Datasets for Untransformed Model

Figure 8: Comparison of Residual Distributions Between Subsets for Logarithmic Model

predicted values from the model, meaning that in some sense they are both more predictive of the general trends in premature birth weights than of non-premature birth weights. Further, the logarithmic model is much better at predicting births that are at the higher ends of the weight range than at the lower end of the range, while the non-transformed model is more consistent across weight ranges.

	R^2		M	AE
Subset	Orig.	log	Orig.	log
Births	0.4543	0.5221	0.5732	0.4956
Preemies	0.5521	0.6559	0.7119	0.8867
Not Preemies	0.1706	0.1490	0.5528	0.4380

Table 7: Summary of Comparison Between Subset Evalution of Nontransformed and Logarithmic Models

4.3 Interaction Model

Our next goal was to see if we could improve accuracy by fitting a full interaction model. Since the assumptions model is intended to be used for interpreting the regression term coefficients, we decided that we would only fit interaction for the accuracy-optimization model. Fitting the full interaction model yielded the following:

The accuracy of this model is much improved; our R^2 value is nearly 30% higher than it was previously, and our RMSE, AIC and BIC values each have seen modest improvements. However, there are far too many terms, leading to a much lower proportion of significant estimations for the linear regression coefficients, and pruning is necessary to reduce the size of the model to only the most important terms.

4.4 Interaction Model Pruning

To prune our interaction model, we used the step-wise reduction method to reduce terms on the basis of AIC values. Doing so yields the following model:

Figure 9: Comparison of Residual Distributions Between Subsets for Pruned Interaction Logarithmic Model

```
Black:Smoke + Black:Preemie, data = births)

xtable(step.reduced.accur, caption="Predictor Statistics and Significance for Log-Transformed Pruned Inter

modelSummary(step.reduced.accur, coef=F, plot=F)

## [1] "R-squared: 0.655809585063165"

## [1] "Adjusted R-Squared: 0.651352442999235"

## [1] "RMSE: 0.59046384902106"

## [1] "AIC: 2534.78504921359"

## [1] "BIC: 2639.79775945157"

## [1] "Quantile Departure: 0.0933319382706169"
```

This leaves us with a total of 18 regression attributes with associated estimated linear regression coefficients, of which all but 3 have significant p-values.

```
specs <- trainControl(method="LOOCV")</pre>
model.cval.LOO <- train( WeightLogSC ~ Twin + Triplet + Sex + MomAgeSC + WeeksSC +
                  Black + GainedSC + Smoke + Preemie + Twin: WeeksSC + Triplet: WeeksSC + MomAgeSC: Smoke +
                  WeeksSC:Black + WeeksSC:GainedSC + WeeksSC:Smoke + WeeksSC:Preemie +
                  Black: Smoke + Black: Preemie, data=births,
                      method = "lm",
                      trControl = specs,
                      na.action = na.omit)
model.cval.L00$results[["Method"]] <- "L00CV"</pre>
specs <- trainControl(method="CV", number=10)</pre>
model.cval.cv <- train( WeightLogSC ~ Twin + Triplet + Sex + MomAgeSC + WeeksSC +
                  Black + GainedSC + Smoke + Preemie + Twin: WeeksSC + Triplet: WeeksSC + MomAgeSC: Smoke +
                  WeeksSC:Black + WeeksSC:GainedSC + WeeksSC:Smoke + WeeksSC:Preemie +
                  Black: Smoke + Black: Preemie, data=births,
                      method = "lm",
                      trControl = specs,
                      na.action = na.omit)
model.cval.cv$results[["Method"]] <- "K-Fold"</pre>
rbind(model.cval.L00$results,
      model.cval.cv$results %>% select(intercept, RMSE, Rsquared, MAE, Method)) %>%
  select(Method, RMSE, Rsquared, MAE) %>%
  xtable(caption="Comparison of Cross-Validation Techniques on Logarithmic Model", label="cross.val.log")
```

After conducting k-fold and leave-one-out cross validation, we find that they are in general agreement with our calculations of RMSE, R^2 , and MAE; the k-fold cross validation has a value of R^2 that is somewhat lower than the original calculation and the LOOCV value, but not by much, and at a level that is still much higher than our original model.

Similar to our estimates with the earlier model, we wanted to see how the model performed on the three sections of the dataset, our full dataset, the premature births, and the non-premature births.

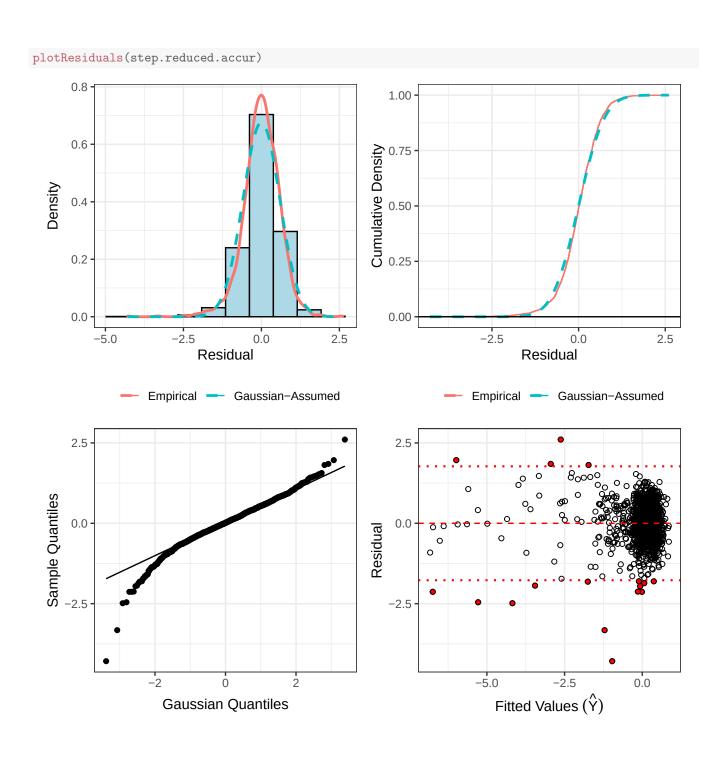


Figure 10: Distribution of Residuals for Pruned Interaction Logarithmic Model

As Figure 9 demonstrates when compared to Figure 8, the residuals seem more evenly distributed. Premature births and birth weights around the lower end of the estimation tend to be overestimated, but predictions, especially around the main range of values, seems much more accurate, with the only notable area with large departures coming from reduction of mid-ranged values of the non-premature babies. The Residual-Density plot from Figure 10 indicates that, if anything, residuals in the neighborhood of 0 are somewhat over-represented.

4.5 Variance Inflation Factor Assessment

Before finalizing our models, we checked to ensure that variance inflation did not play into the model's accurace using the VIF test.

In Table 11, we see that all VIF values fall below 3, and are thus very acceptable; for Table 12, all but one value falls below 5, which is the highly-acceptable limit, and the last remaining value still falls below 10, which is the moderatly-acceptable limit. We are thus confident that variance inflation is not a particularly heavy factor in our model, and collinearity is not a significant problem.

5 Final Model(s) and Conclusions.

```
assessModel <- function(model, p) {</pre>
  print(modelSummary(model, plots=F))
  cbind(vif(model), vif(model)[,3]^2)
  print(summary(model$residual))
  print(confint(model))
  lev <- model$model %>% mutate(h.values = hatvalues(model))
  print(summary(lev$h.values))
  n <- nrow(model$model)</pre>
  high.lev <- lev %>% filter(h.values > 2*p/n)
  print(paste("High Lev.:", nrow(high.lev)))
  v.high.lev <- lev %>% filter(h.values > 3*p/n)
  print(paste("Very High Lev.:", nrow(v.high.lev)))
  new.resid <- model$model %>% mutate(stdres = rstandard(model),
                                       stures = rstudent(model))
  print("Standard Residual Quant.:")
  print(summary(new.resid$stdres))
  print("Studentized Residual Quant.:")
  print(summary(new.resid$stures))
  s.outliers.stdres <- new.resid %>% filter(abs(stdres)>3)
  print(paste("Strong Standard Residual Outliers:", nrow(s.outliers.stdres)))
  print(s.outliers.stdres)
  s.outliers.stures <- new.resid %>% filter(abs(stures)>3)
  print(paste("String Studentized Residual Outliers:", nrow(s.outliers.stures)))
```

```
print(s.outliers.stures)
  cooks.values <- model$model %>% mutate(cooks = cooks.distance(model))
  print("Cook's Values:")
  print(summary(cooks.values$cooks))
  cooks.strong <- cooks.values %>% filter(cooks>1)
  print(paste("Strong C. Values:", nrow(cooks.strong)))
}

assessModel(mod.accur.final)
assessModel(mod.accur.final.part)
assessModel(min.vif.model)
```

Highest Accuracy Model:

```
\hat{y} = 0.1813 - 1.2273 \cdot \text{I(Twin} = \text{TRUE)} - 1.0309 \cdot \text{I(Triplet} = \text{TRUE)} + 0.1110 \cdot \text{I(Sex} = \text{Male)} + 0.0912 \cdot \text{MomAgeSC} + 0.1681 \cdot \text{MomAgeSC}
```

Non-Interact Assumptions Model

```
\hat{y} = 0.0948 - 1.1201 \cdot \text{I(Twin} = \text{TRUE)} - 1.4721 \cdot \text{I(Triplet} = \text{TRUE)} + 0.1522 \cdot \text{I(Sex} = \text{Male)} \\ + 0.1186 \cdot \text{MomAgeSC} + 0.4186 \cdot \text{MomAgeSC
```

	Estimate	Std. Error	t value	$\Pr(> t)$
(Intercept)	0.1686	0.0324	5.21	0.0000
Twin	-1.2867	0.2003	-6.42	0.0000
Triplet	-1.6956	1.6094	-1.05	0.2923
Male	0.1304	0.0432	3.02	0.0026
MomAge (SC)	0.0955	0.0278	3.43	0.0006
Weeks (SC)	0.1804	0.0410	4.40	0.0000
Black	-0.1775	0.0632	-2.81	0.0050
Gained (SC)	0.1436	0.0282	5.08	0.0000
Smoke	-0.2597	0.0780	-3.33	0.0009
Preemie	0.8388	0.1400	5.99	0.0000
Twin:Male	0.1786	0.2021	0.88	0.3770
Triplet:Male	0.5711	0.8682	0.66	0.5108
Twin:MomAge (SC)	-0.0038	0.1098	-0.03	0.9725
Triplet:MomAge (SC)	0.1514	1.0843	0.14	0.8890
Twin:Weeks (SC)	-0.2927	0.1009	-2.90	0.0038
Triplet:Weeks (SC)	-0.2034	0.2702	-0.75	0.4518
Twin:Black	-0.3361	0.3158	-1.06	0.2873
Twin:Gained (SC)	-0.0882	0.0954	-0.92	0.3554
Twin:Smoke	0.3392	0.5266	0.64	0.5196
Twin:Preemie	0.0904	0.2789	0.32	0.7458
Male:MomAge (SC)	0.0140	0.0330	0.42	0.6723
Male:Weeks (SC)	-0.0221	0.0497	-0.44	0.6568
Male:Black	-0.0883	0.0778	-1.13	0.2566
Male:Gained (SC)	0.0139	0.0326	0.43	0.6699
Male:Smoke	-0.0280	0.0927	-0.30	0.7629
Male:Preemie	-0.0164	0.1385	-0.12	0.9056
MomAge (SC):Weeks (SC)	0.0003	0.0256	0.01	0.9905
MomAge (SC):Black	-0.0342	0.0399	-0.86	0.3916
MomAge (SC):Gained (SC)	-0.0041	0.0174	-0.23	0.8157
MomAge (SC):Smoke	-0.1053	0.0474	-2.22	0.0265
MomAge (SC):Preemie	-0.0329	0.0697	-0.47	0.6371
Weeks (SC):Black	0.1516	0.0545	2.78	0.0055
Weeks (SC):Gained (SC)	-0.0641	0.0250	-2.56	0.0105
Weeks (SC):Smoke	-0.1369	0.0627	-2.18	0.0293
Weeks (SC):Preemie	1.1095	0.0663	16.74	0.0000
Black:Gained (SC)	-0.0330	0.0372	-0.89	0.3759
Black:Smoke	0.1584	0.1152	1.37	0.1694
Black:Preemie	0.5377	0.1605	3.35	0.0008
Gained (SC):Smoke	-0.0404	0.0433	-0.93	0.3507
Gained (SC):Preemie	-0.0642	0.0761	-0.84	0.3992
Smoke:Preemie	0.0710	0.1898	0.37	0.7083

Table 8: Predictor Statistics and Significance for Log-Transformed Interaction Model

	Estimate	Std. Error	t value	$\Pr(> t)$
(Intercept)	0.1813	0.0279	6.49	0.0000
Twin	-1.2273	0.1369	-8.97	0.0000
Triplet	-1.0309	0.6270	-1.64	0.1004
Male	0.1110	0.0316	3.51	0.0005
MomAge (SC)	0.0912	0.0178	5.14	0.0000
Weeks (SC)	0.1688	0.0321	5.26	0.0000
Black	-0.2197	0.0471	-4.66	0.0000
Gained (SC)	0.1250	0.0160	7.80	0.0000
Smoke	-0.2696	0.0516	-5.22	0.0000
Preemie	0.8473	0.1045	8.10	0.0000
Twin:Weeks (SC)	-0.2970	0.0700	-4.25	0.0000
Triplet:Weeks (SC)	-0.0969	0.2116	-0.46	0.6470
MomAge (SC):Smoke	-0.1005	0.0466	-2.16	0.0313
Weeks (SC):Black	0.1588	0.0521	3.05	0.0023
Weeks (SC):Gained (SC)	-0.0422	0.0169	-2.50	0.0127
Weeks (SC):Smoke	-0.1502	0.0422	-3.56	0.0004
Weeks (SC):Preemie	1.1080	0.0601	18.45	0.0000
Black:Smoke	0.1590	0.1121	1.42	0.1564
Black:Preemie	0.5283	0.1537	3.44	0.0006

Table 9: Predictor Statistics and Significance for Log-Transformed Pruned Interaction Model

Method	RMSE	R^2	MAE
Orig.	0.59	0.66	0.44
LOOCV	0.60	0.64	0.45
K-Fold	0.62	0.61	0.45

Table 10: Comparison of Cross-Validation Techniques on Logarithmic Model

	VIF
Twin	1.13
Triplet	1.04
Sex	1.00
MomAge (SC)	1.07
Weeks (SC)	2.11
Black	1.05
Gained (SC)	1.02
Smoke	1.02
Preemie	2.06

Table 11: VIF Values for Pruned Nontransformed Regression Model

	VIF
Twin	2.24
Triplet	4.50
Sex	1.01
MomAge (SC)	1.27
Weeks (SC)	4.02
Black	1.59
Gained (SC)	1.04
Smoke	1.34
Preemie	4.95
Twin:Weeks (SC)	2.98
Triplet:Weeks (SC)	4.64
MomAge (SC) Smoke	1.24
Weeks (SC) Black	3.51
Weeks (SC) Gained (SC)	1.30
Weeks (SC) Smoke	1.40
Weeks (SC) Preemie	7.91
Black:Smoke	1.44
Black:Preemie	3.64

Table 12: VIF Values for Pruned Logarithmic Interaction Regression Model