## Bayesian Final Project

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5/12/2023

## 2.1 Proposal

In this project we will be developing Bayesian models in order to identify the effects of mother's characteristics including her age, gestational length, smoking status, and BMI, on female new-born's birth weight.

#### 2.2 Attribution

Tianyi Li: Project leader, team management, raw data summary, build statistical model

Jade Gu: Raw data summary, model fitting and selection, assisting in prior predictive simulation

Jiashu Liu: Prior predictive simulation, assisting in model fitting and selection

Jeremy Lu: Presentation preparation, assiting in model selection, summarize key convergence diagnostics

## 2.3 Raw Data Summary

#### 2.3.1 Data Origin

Below is a link to a compressed CSV file containing US birth data for the year 2018. The data is provided by the National Center for Health Statistics and can be accessed from the National Bureau of Economic Research website.

#### 2.3.2 Data Clean (Include Variable Selection)

This 2018 birth data contains 3801534 obs. of 240 variables. First, based on our research topic, we will keep girls' data only. Second, we will pick 6 most interesting variables including: dbwt(infant's birthweight in grams), mager(mother's years of age), dmar(martial status), cig\_rec(cigarette recode), bmi(body mass index), combgest(combined gestation weeks). Then, we will exclude any missing values. After that, we will factor all categorical variables.

### 2.3.3 Data Sampling

Since this birth data after filter contains 1591008 obs. which is still extremely large, we will get a random sample of births that includes only 0.1% of all obs, which results in 1591 obs.

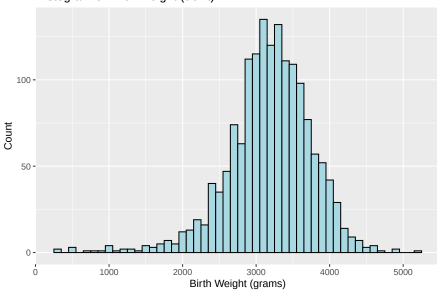
#### 2.3.4 Exploratory data analysis

In the birth data, we have four numerical variables and two factored categorical variables.

Numerical variables:

#### dbwt (Birth weight)

#### Histogram of Birth Weight (dbwt)



Above is a histogram plot of birth weight of female infant babies. Values ranged from 320g (Yes it's possible) to 5220g. We computed some descriptive statistics: Median = 3220; Mean = 3183; SD = 574.74.

#### mager (Mother's age)

Mother's integer years of age. Values ranged from 14 to 47 yrs old. We computed some descriptive statistics: Median = 29; Mean = 28.59; SD = 5.86.

#### hmi

Mother's body mass index. Values ranged from 13.60 to 69.10. We computed some descriptive statistics: Median = 25.70; Mean = 27.29; SD = 7.00.

#### combgest

Mother's combined gestation in weeks. Values ranged from 20.00 to 47.00. We computed some descriptive statistics: Median = 39.00; Mean = 38.55; SD = 2.49.

Categorical variables:

#### dmar

Mother's martial status. "1" stands for married, and "0" stands for unmarried. According to the birth data, 956 out of 1591 observations (60.09%) are married, and the remaining 635 observations (39.91%) are

unmarried. The mean birth weight of married moms is 3260g and median is 3280g, while the mean and median birth weight of unmarried is 3068g and 3105g.

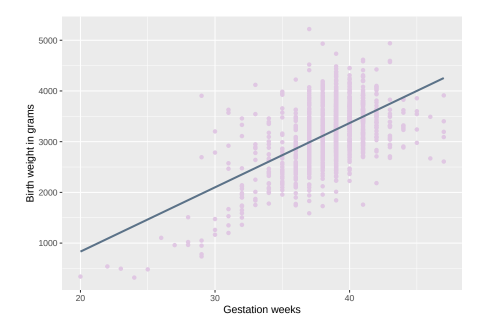
## $cig\_rec$

Whether the mother smokes or not. "1" stands for smoke, and "0" stands for non-smoke. According to the birth data, 106 out of 1591 observations (6.66%) are smokers, and the remaining 1481 observations (93.34) are non-smokers. The mean birth weight of moms who smoke is 2999g and median is 3023g, while the mean and median birth weight of mom who don't smoke is 3197g and 3232g.

#### 2.3.5 More visualization

```
ggplot(birth, aes(x = combgest, y = dbwt)) +
geom_point(color = "#e0c7e3") +
geom_smooth(method = "lm", se = FALSE, color = "#5b7288") +
labs(x = "Gestation weeks", y = "Birth weight in grams")
```

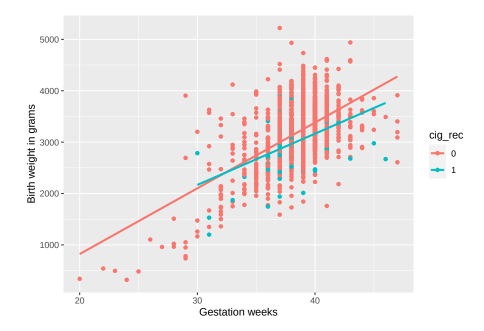
## 'geom\_smooth()' using formula = 'y ~ x'



We can observe a positive linear relationship between the mother's combined gestation length and the female newborn's birth weight. This is expected, mothers with longer combined gestation are expected to get heavier baby girls. However, there might be some variations and potential outliers. Let's add whether or not the mom smokes into account:

```
ggplot(birth, aes(x = combgest, y = dbwt, color = cig_rec)) +
geom_point() +
geom_smooth(method = "lm", se = FALSE) +
xlab("Gestation weeks") +
ylab("Birth weight in grams")
```

```
## 'geom_smooth()' using formula = 'y ~ x'
```



From the scatterplot of birth weight versus gestation weeks of moms who smoke and does not smoke, superimposed with the observed linear relationships in the solid lines, the slope of the lines are different, indicating that there is an interaction effect between the predictor variables and the relationship between gestation weeks and birth weight is different between moms who smoke and those who don't smoke.

#### 2.4 Statistical Model

We aim to model the relationship between the birth weight of female newborns and maternal characteristics, including age, marital status, smoking status, body mass index, and gestation weeks.

From the histogram of birth weight before, we can assume that the birth weight of each female newborn follows a normal distribution. Also, from the scatterplot above, there is a positive linear relationship between the gestation length and birth weight. Let  $Y_i$  denotes the birth weight for observation i. Since  $Y_i$  is numerical and we assume there is a linear relationship between Y and the predictor variables, a Normal Bayesian regression model is the most appropriate model to use. With 5 predictor variables and 1 outcome variable, our model looks like:

data: 
$$Y_{i}|\beta_{0}, \beta_{1}, ..., \beta_{5}, \sigma \overset{ind}{\sim} N\left(\mu_{i}, \sigma^{2}\right) \quad \text{with} \quad \mu_{i} = \beta_{0} + \beta_{1}X_{i1} + ... + \beta_{5}X_{i5}$$
 priors: 
$$\beta_{0} \sim N\left(m_{0}, s_{0}^{2}\right)$$
 
$$\beta_{1} \sim N\left(m_{1}, s_{1}^{2}\right)$$
 ... 
$$\beta_{5} \sim N\left(m_{5}, s_{5}^{2}\right)$$
 
$$\sigma \sim \operatorname{Exp}(l).$$
 
$$(1)$$

For observation of female new-born i,  $Y_i$  is her birth weight,  $X_{i1}$  is her mother's gestation length in week,  $X_{i2}$  is a binary outcome indicating whether her mom smokes,  $X_{i3}$  is her mom's bmi index,  $X_{i4}$  is her mom's age,  $X_{i5}$  is a binary variable of whether her mom is married.

We have to specify the prior mean and sd of each variable. For simplicity, we assume that the prior distributions are independent. First, we center all numerical predictor variables by subtracting the value of each observation by its mean. Now the intercept  $\beta_0$  represents the birth weight of an average female new-born whose mother is unmarried and does not smoke. According to Medical News Today, the average birth weight of a full-term female is 3.2 kg, which normally ranges from 2.5 kg to 4.0 kg; however, it can also

be below 1.5 kg or above 4.5 kg. As a result, we take 3200 and 750 as prior mean and standard deviation for  $\beta_0$ .

- For  $\beta_1$  of  $X_1$ , the gestation week, from prior scientific understanding, on average, one week longer in gestation length is associated with an increase in birth weight by 10 to 225 grams, so we take 108 and 54 as mean and sd for  $\beta_1$ .
- For  $\beta_2$  of  $X_2$ , whether mom smokes or not, from prior scientific research, on average, the birth weight of female baby whose mother smokes could be 200 to 300 grams lower than baby whose mother doesn't smoke. We take -250 and 25 for prior mean and sd of  $\beta_2$ .
- For  $\beta_3$  of  $X_3$ , the bmi index of baby's mother, from prior scientific research findings, on average, one unit increase in the bmi index is associated with 20 to 30 grams increase in birth weight. We take 25 and 2.5 as prior mean and sd for  $\beta_3$ .
- For  $\beta_4$  of  $X_4$ , the mom's age. we employ weakly informative criteria and set 0 and 2.5 as prior mean and sd.
- For  $\beta_5$  of  $X_5$ , the marriage status of baby's mom, prior scientific study suggests that the average birth weight of female infants born to married mothers may be around 50 grams to 100 grams higher. We take 75 and 12.5 as prior mean and sd for  $\beta_5$
- Finally, for the error term, a plausible exponential rate parameter is 0.003. We have  $\sigma \sim Exp(0.003)$ .

#### 2.5 Prior Predictive Simulation

Now let's start simulate some data based on our prior parameters. First we define our prior model function in R.

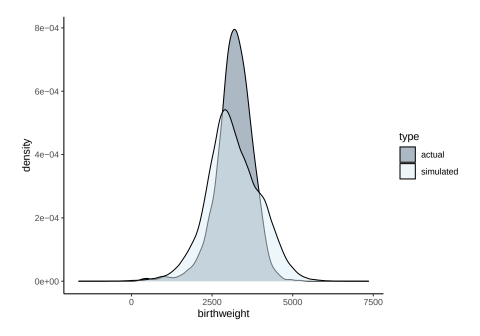
```
set.seed(84735)
# Define function
prior_pred <- function(data) {</pre>
  beta_0 <- rnorm(1, 3200, 750)
  beta 1 <- rnorm(1, 108, 54)
  beta_2 \leftarrow rnorm(1, -250, 25)
  beta_3 \leftarrow rnorm(1, 25, 2.5)
  beta_4 \leftarrow rnorm(1, 0, 2.5)
  beta_5 <- rnorm(1, 75, 12.5)
  sigma \leftarrow rexp(1, 0.003)
  1 <- nrow(data)</pre>
  y <- numeric(1)
  for (i in 1:1) {
    # Convert factors to numeric
    cig_rec_num <- as.numeric(data$cig_rec[i])</pre>
    dmar_num <- as.numeric(data$dmar[i])</pre>
    mu <- beta 0 +
      beta_1 * data$centered_combgest[i] +
      beta_2 * cig_rec_num +
      beta_3 * data$centered_bmi[i] +
      beta 4 * data$centered mager[i] +
      beta_5 * dmar_num
    y[i] <- rnorm(1, mu, sigma)
```

```
}
  return(y)
}

# Simulate 100 times
n <- 100
pr_p <- replicate(n = n, prior_pred(birth))
dim(pr_p)</pre>
```

#### ## [1] 1591 100

To get a sense of how our prior parameters perform, we draw a density plot of the original data and our simulated data.



The model looks good. However, the simulated data indicates a higher variability compared to the actual data. So we decide to lower the sd of our prior parameters.

## 2.6 Modeling Fitting, PPCs, and Model Selection

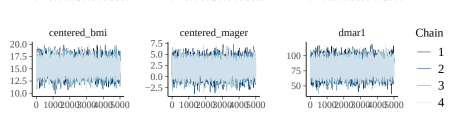
```
# Set prior
prior_intercept <- normal(3200, 750, autoscale = FALSE)</pre>
prior_beta <- normal(c(108, -250, 25, 0, 75),</pre>
                      c(50, 25, 2, 2, 12),
                      autoscale = FALSE)
prior_sigma <- exponential(0.003)</pre>
# Fit the model
birth_model <- stan_glm(dbwt ~</pre>
                            centered_combgest +
                            cig_rec +
                            centered_bmi +
                            centered_mager +
                            dmar,
                          data = birth,
                          prior_intercept = prior_intercept,
                          prior = prior_beta,
                          prior_aux = prior_sigma,
                          chains = 4,
                          iter = 5000*2,
                          seed = 84735)
```

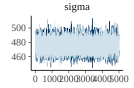
Then, we will create and interpret both visual and numerical diagnostics of our MCMC simulation.

```
# Visual diagnostics of MCMC simulation
mcmc_trace(birth_model, size = 0.1)
                              (Intercept)
                                                       centered_combgest
                                                                                          cig_rec1
                    3210
                                                                                      depthal distrigent the district all trans
                                                  140
                                                       amelika da kumaken yan bel
                    3180
                                                                              -200
                                                  130
                    3150
                                                                              -250
                                                  120
                    3120
                                                                              -300
                                                  110
```

0 10002000300040005000

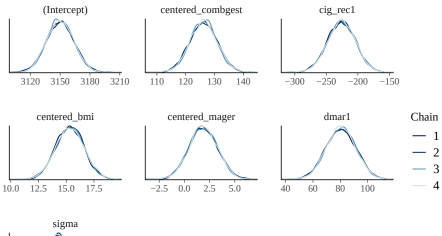
0 10002000300040005000

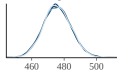




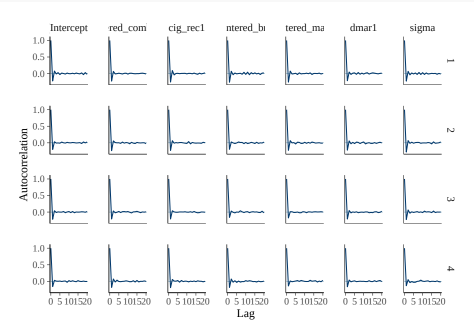
0 10002000300040005000

mcmc\_dens\_overlay(birth\_model)





## mcmc\_acf(birth\_model)



# # Numerical diagnostics of MCMC simulation neff\_ratio(birth\_model)

##	(Intercept)	centered_combgest	cig_rec1	centered_bmi
##	1.48540	1.45490	1.43985	1.51560
##	centered_mager	dmar1	sigma	
##	1.49440	1.49680	1.59830	

## rhat(birth\_model)

## (Intercept) centered\_combgest cig\_rec1 centered\_bmi

##	0.9998547	0.9998940	1.0000289	0.9999481
##	centered_mager	dmar1	sigma	
##	0.9998671	0.9999052	0.9999623	

Regarding the assessment of our MCMC simulation, the visual diagnostics provide valuable insight into the behavior of the Markov chains. In particular, we have examined the trace plots, density plots, and autocorrelation plots for each parameter. Based on the trace plots, we can observe that the lines for each parameter fluctuate randomly without exhibiting any clear patterns. Additionally, all four chains appear to mix well, which is a positive indication that the simulation is properly exploring the parameter space. Furthermore, the density plots show that each parameter has a smooth, bell-shaped distribution, which is a desirable property of a well-fitting model. Moreover, the autocorrelation plots exhibit a quick decay in correlation as the lag increases for each parameter, which is indicative of good autocorrelation properties.

Moving onto the numerical diagnostics of our MCMC simulation, we have calculated the neff\_ratio and R-hat statistics for each parameter. In terms of numerical diagnostics, the neff\_ratio statistic shows that all parameters have a ratio larger than 0.5, indicating a good effective sample size. Additionally, the R-hat statistic shows that all parameters have a value close to 1, indicating good convergence of the chains. These numerical diagnostics further confirm that our posterior simulation has stabilized.

Taken together, the results of both the visual and numerical diagnostics provide strong evidence that our posterior simulation has sufficiently stabilized, and that we can trust the results of our Bayesian analysis.

## 2.6.2 Tidy summary

Next, we will produce a tidy() summary of this model. In addition, we will interpret the non-intercept coefficients' posterior median values in context.

```
## # A tibble: 8 x 5
##
     term
                         estimate std.error
                                               conf.low conf.high
##
     <chr>>
                            <dbl>
                                       <dbl>
                                                  <dbl>
                                                             <dbl>
## 1 (Intercept)
                          3150.
                                       13.8
                                             3133.
                                                           3168.
                           126.
                                              120.
## 2 centered_combgest
                                        4.79
                                                            132.
## 3 cig_rec1
                          -226.
                                       22.4
                                              -255.
                                                           -198.
                                        1.30
                                                             17.0
## 4 centered_bmi
                            15.3
                                                13.7
## 5 centered mager
                             1.85
                                                 0.0304
                                                              3.72
                                        1.43
## 6 dmar1
                                       11.0
                            80.6
                                                66.5
                                                             94.7
## 7 sigma
                           476.
                                        8.60
                                               465.
                                                            487.
## 8 mean_PPD
                          3184.
                                       16.9 3162.
                                                           3205.
```

Interpretations of coefficients of some key variables:

- The posterior median value of the intercept in the model is 3150, indicating that on average, the expected weight of a female newborn whose mother has average age, bmi, gestation length, is not married and doesn't smoke is 3150 grams.
- The posterior median value of the centered\_combgest coefficient is 126, suggesting that for each additional gestational week, the expected weight of the female newborn increases by 126 grams, holding all other predictors in the model constant.
- The posterior median value of the cig\_rec1 coefficient is -226, indicating that mothers who smoke during pregnancy are expected have female newborns that weigh around 226 grams less on average than non-smoking mothers, holding all other predictors in the model constant.

## 2.6.3 Posterior Summary

Here, we will use posterior\_interval() to produce 95% credible intervals for the model parameters. After that, we will check any association between independent variables and dependent variables.

```
# posterior summarizes
posterior_interval(birth_model, prob = 0.95)
```

```
##
                             2.5%
                                         97.5%
## (Intercept)
                     3123.5490609 3177.283226
## centered_combgest 116.7908392
                                  135.733853
## cig rec1
                     -270.2001644 -182.386082
## centered_bmi
                       12.8151132
                                     17.885912
## centered mager
                       -0.9388254
                                      4.671981
## dmar1
                       59.1699660
                                   101.648477
## sigma
                      459.2321583 492.998295
```

When controlling for the other predictors in the model, the 95% posterior credible intervals for the coefficient of centered\_combgest, centered\_bmi, and dmar1 lie entirely above 0, suggesting that centered\_combgest, comtered\_bmi, and dmar1 have significant positive associations with dbwt. The 95% posterior credible intervals for the coefficient of centered\_mager contains 0, suggesting that centered\_mager has no association with dbwt. The 95% posterior credible intervals for the coefficient of cig\_rec1 lie entirely below 0, suggesting that cig\_rec1 has significant negative association with dbwt. In other words, the model suggests that increasing the gestational period (centered\_combgest) and maternal BMI (centered\_bmi) are associated with higher birthweights of female newborns, as is being married (dmar1), and smoking (cig\_rec1) is associated with a significant decrease in the birthweight.

## 2.6.4 More models

Next, we will bring more models and see which ones work the best. For our new model birth\_model2, we are going to add two interaction terms: cig\_rec&bmi and mager&dmar. We believe that mother's age and her marital status should be correlated, and smokers probably would have lower BMI. To fit birth\_model2, we will use weakly informative priors throughout.

```
# Use weakly informative criteria throughout
# Add interaction: ciq_rec*centered_bmi, centered_mager*dmar
birth_model2 <- stan_glm(dbwt ~</pre>
                            centered_combgest +
                            cig_rec +
                            centered_bmi +
                            centered_mager +
                            dmar +
                            cig_rec:centered_bmi +
                            centered mager:dmar,
                         data = birth,
                         family = gaussian,
                         prior intercept = normal(3200, 750),
                        prior = normal(0, 2.5, autoscale = TRUE),
                        prior aux = exponential(1, autoscale = TRUE),
                         chains = 4, iter = 5000*2, seed = 84735,
                         prior_PD = FALSE)
```

Next, to determine the necessity of interaction terms, we will check the 80% posterior credible interval for interaction coefficients.

```
# Posterior summary statistics
tidy(birth_model2, effects = c("fixed", "aux"))
## # A tibble: 10 x 3
##
      term
                             estimate std.error
##
      <chr>
                                <dbl>
                                          <dbl>
    1 (Intercept)
                                          20.9
##
                             3145.
##
    2 centered_combgest
                              125.
                                           4.88
##
  3 cig_rec1
                              -139.
                                          48.1
## 4 centered_bmi
                                7.73
                                           1.78
## 5 centered_mager
                                7.68
                                           3.20
##
  6 dmar1
                                90.8
                                          26.1
  7 cig_rec1:centered_bmi
                                 6.40
                                           7.31
                                -7.01
                                           4.29
  8 centered_mager:dmar1
## 9 sigma
                              473.
                                           8.48
## 10 mean_PPD
                             3183.
                                          16.7
# Posterior credible interval for the interaction terms
posterior_interval(birth_model2, prob = 0.80,
                   pars = "cig_rec1:centered_bmi")
##
                                10%
                                         90%
## cig_rec1:centered_bmi -2.909131 15.60692
posterior_interval(birth_model2, prob = 0.80,
                   pars = "centered_mager:dmar1")
##
                              10%
                                         90%
## centered_mager:dmar1 -12.49847 -1.431468
```

The 80% posterior credible interval for interaction coefficient  $\beta 6$  contains 0, suggesting that the association between smoking and BMI is not significant. Thus, this interaction term is not necessary. The 80% posterior credible interval for interaction coefficient  $\beta 7$  is entirely and well below 0, suggesting a negative association between mother's age and marriage status. And we believe that we should include this interaction term in our new model birth\_model3.

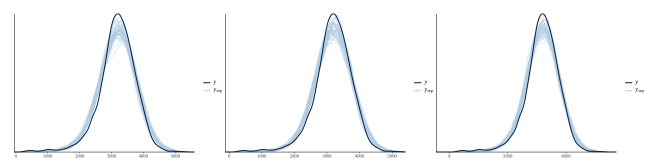
```
prior = normal(0, 2.5, autoscale = TRUE),
prior_aux = exponential(1, autoscale = TRUE),
chains = 4, iter = 5000*2, seed = 84735,
prior_PD = FALSE)
```

## 2.6.5 Model Comparison

To compare our 3 models and decide which one is the best, we will use three different approaches, including posterior predictive checks, cross-validation, and ELPD.

The first method is to evaluate predictive accuracy using pp\_check.

```
pp_check(birth_model)
pp_check(birth_model2)
pp_check(birth_model3)
```



According to pp\_check() output, all three models preform pretty good. Although the simulations are not perfect, but they do reasonably capture the features of the observed data.

The second method is to evaluate predictive accuracy using cross-validation.

```
# Run a 10-fold cross-validation
cv_procedure1 <- prediction_summary_cv(model = birth_model, data = birth, k = 10)
cv_procedure2 <- prediction_summary_cv(model = birth_model2, data = birth, k = 10)
cv_procedure3 <- prediction_summary_cv(model = birth_model3, data = birth, k = 10)
# Compare 3 model's mean
cv_procedure1$cv</pre>
### mae mae_scaled within_50 within_95
```

```
cv_procedure2$cv
```

mae mae\_scaled within\_50 within\_95

## 1 319.9812 0.6721165 0.5116234 0.9515841

##

```
cv_procedure3$cv
```

```
## mae mae_scaled within_50 within_95
## 1 308.9813  0.6504977 0.5172799 0.9522327
```

Based on the cross-validated metrics provided, birth\_model3 has the lowest mae, lowest mae\_scaled, highest within\_50, and highest within\_95 than the other two models. Its lowest mae and mae\_scaled value indicate that the this model on average makes most accurate predictions, and its highest within\_50 and within\_95 value indicate that the this model is more likely to make accurate predictions within the selected interval. Therefore, the best model is birth model3.

The third method is to evaluate predictive accuracy using ELPD.

```
# Calculate ELPD for the 3 models
set.seed(84735)
loo_1 <- loo(birth_model)</pre>
loo_2 <- loo(birth_model2)</pre>
loo_3 <- loo(birth_model3)</pre>
# Results
c(loo_1$estimates[1], loo_2$estimates[1], loo_3$estimates[1])
## [1] -12068.69 -12060.64 -12060.01
# Compare the ELPD for the 3 models
loo_compare(loo_1, loo_2, loo_3)
                 elpd_diff se_diff
                            0.0
## birth_model3 0.0
## birth_model2 -0.6
                            0.9
## birth_model -8.7
                            5.2
```

According to loo\_compare() output, birth\_model3 has the highest ELPD, indicating it is the best model out of three.

#### 2.7 Discussion