# Incomplete Data Analysis - Assignment 3

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# Question 1

# (a) - Solution

Function cc() in mice package returns the complete cases in a dataset. Thus, we count the number of complete cases in dataset 'nhanes' and further compute the percentage of incomplete cases by  $1 - \frac{\#complete\ cases}{\#all\ cases}$ .

As a result, we obtain the percentage of incomplete cases is 48% (12 cases out of 25).

```
# the percentage of incomplete cases
1 - nrow(cc(nhanes)) / nrow(nhanes)
```

## [1] 0.48

#### (b) - Solution

Recall that to estimate the variance of  $\hat{\theta}^{MI}$ , the multiple imputation of  $\theta$ , we calculate the following statistics:

- 1. between-imputation variance:  $B = \frac{1}{M-1} \sum_{i=1}^{M} \left( \hat{\theta}^{(i)} \hat{\theta}^{MI} \right)^2$
- 2. within-imputation variance:  $\overline{U} = \frac{1}{M} \sum_{i=1}^{M} \hat{U}^{(i)}$ , where  $\hat{U}^{(i)}$  is the estimated variance of  $\hat{\theta}^{(i)}$

Then we obtain the **total variance**  $V^{MI} = \overline{U} + \left(1 + \frac{1}{M}\right)B$ . Note that M denotes the number of imputed datasets by mice (M = 5) by default).

According to the work by Buuren et al. 2011, the proportion of total variance that is attributed to the missing values is

$$\lambda = \frac{B + \frac{B}{M}}{V^{MI}}$$

We yield the following results with the standard MICE procedure - mice(), with() and pool(). Note that in step 2 (with()), we fit the normal linear regression of bmi over age, hyp and cyl.

As mentioned before, we look at the column lambda to get the proportion of variance due to missing data for each parameter (3 decimal places, seed = 1):

- 1.  $\lambda_{age} = 0.686$
- 2.  $\lambda_{hyp} = 0.350$
- 3.  $\lambda_{chl} = 0.304$

Based on the proportion of variance due to missing data, we conclude that age appears to be the most affected by the non-response with the largest value of  $\lambda$ .

```
# impute dataset with mice
imp.list <- mice(nhanes, printFlag = FALSE, seed = 1)

# linear model to predict bmi
bmi.fit <- with(imp.list, lm(bmi ~ age + hyp + chl))

# pool the results
bmi.pool <- pool(bmi.fit)
bmi.pool$pooled</pre>
```

```
##
            term m
                                                                      t dfcom
                      estimate
                                        ubar
                                                        b
## 1 (Intercept) 5 19.61789252 10.588884721 0.8662133972 11.628340797
                                                                           21
             age 5 -3.55287155
## 2
                                0.744810536 1.3585594461
                                                           2.375081872
                                                                           21
                                2.886391704 1.2976511612
## 3
             hyp 5 2.19701748
                                                           4.443573098
                                                                           21
             chl 5 0.05378081
## 4
                                0.000287288 0.0001046083
                                                           0.000412818
                                                                           21
##
                      riv
                              lambda
                                            fmi
## 1 16.936189 0.09816483 0.08938989 0.1807424
     3.528053 2.18884032 0.68640637 0.7824821
     9.035494 0.53949067 0.35043452 0.4583762
## 4 10.228828 0.43694808 0.30408063 0.4092932
```

#### (c) - Solution

We repeat same analysis on  $\lambda$  by using different random seeds from 2 to 6. The result are shown in the data frame below.

We can observe that the conclusions in 1.(b) do not remain the same for different seeds. The values of  $\lambda$  and the parameter most affected by the non-response vary dramatically.

For seed 2, 3 and 6, age has the largest value of  $\lambda$  (0.403, 0.590, 0.655 of each, 3 decimal places) and is considered to be most affected by the non-response, which is consistent with conclusions in 1.(b), using random seed 1. On the other hand, chl and hyp take the largest proportion of variance for seed 4 and 5 respectively, with values 0.331 and 0.594 of each.

Recall that the larger amount of missing data, the larger the variability of values of  $\lambda$  will be. Since nearly half (48%) of cases in the dataset are incomplete, it is reasonable to have different conclusions every time we adjust the random seed.

```
##
                                     chl
     seed
                age
                          hyp
## 1
        1 0.6864064 0.3504345 0.3040806
        2 0.4033924 0.1430995 0.2959966
## 2
## 3
        3 0.5895051 0.4101152 0.5621346
        4 0.2189333 0.1961083 0.3305334
## 4
## 5
        5 0.4511896 0.5942866 0.2346065
## 6
        6 0.6549523 0.2960364 0.5196295
```

#### (d) - Solution

From 1.(c), we observe that the parameter most affected by the non-response varies as random seed changes. Thus, we expect that increasing the number of imputed datasets can lead to more consistent and stable results which has low dependence on the choice of random seed.

We keep using same random seeds from 1 to 6, but change the number of imputed datasets, M, from 5 to 100. Again, we obtain a data frame with values of  $\lambda$  for each parameter with different random seed.

Now we can find that age always takes the largest proportion of variance due to missing cases (largest value of  $\lambda$ ) in 5 of 6 seeds. For seed 3, chl becomes the parameter which is affected by the non-response, with  $\lambda = 0.328$ . It appears that the conclusions become more consistent with M = 100, as we expected.

As far as I am concerned, I prefer the analyses with M = 100.

Recall that the **total variance** of  $\theta$  (bmi in this question) is calculated by

$$V^{MI} = \overline{U} + \left(1 + \frac{1}{M}\right)B$$

where 
$$B = \frac{1}{M-1} \sum_{i=1}^{M} \left( \hat{\theta}^{(i)} - \hat{\theta}^{MI} \right)^2$$
 and  $\overline{U} = \frac{1}{M} \sum_{i=1}^{M} \hat{U}^{(i)}$ .

Therefore, large value of M can reduce the values of both  $\overline{U}$  and B, and further the total variance, increasing the reliability of the estimates. In terms of high accuracy of predictions, larger number of imputed datasets is a better choice.

However, time consumption and low statistical efficiency can be the problems of large value of M. When imputing large scaled dataset with a large number of missing values, large M will relatively increase the running time. Moreover, the imputation procedure arrives at stable and statistically significant conclusions when M reaches a certain value. Increasing M beyond this value does not significantly influence the conclusions but consumes resource and time.

Overall, I still consider M=100 as a better choice as the dataset in this question only have 25 observations and 100 imputed datasets can be easily handled.

```
## seed age hyp chl
## 1 1 0.4324680 0.2915346 0.3217837
## 2 2 0.4031077 0.2825108 0.2939693
## 3 3 0.3093072 0.2425105 0.3281911
## 4 4 0.3943223 0.2565132 0.2835232
## 5 5 0.3322570 0.2893046 0.2461956
## 6 6 0.4430300 0.2860700 0.3113085
```

#### Solution

As required, we apply stochastic regression imputation (SRI) and bootstrap sampling to generate the data in step 2. To calculate the empirical coverage probability, according to **NOTE 1**, we define a counter to count the times when the ground truth value of  $\beta_1$ , i.e. 3, is covered in its 95% empirical confidence interval for each method and further obtain the probability with the corresponding frequency.

From the results below, we observe the empirical coverage probability of bootstrap imputation is larger than the probability of SRI (0.95 vs 0.88). The reason for this is that stochastic regression imputation does not incorporate the variability of function weights, which means the uncertainty of imputed values is not considered, unlike bootstrap imputation. Thus, generally the 95% confidence intervals obtained using SRI are more narrow and less likely to contain the ground truth value.

```
# read dataset into R
load("dataex2.Rdata")
# initialize counters for empirical coverage probabilities of two methods
count.sri <- count.boot <- 0</pre>
# MICE standard procedure for each dataset
for (i in 1:100){
  # step 1 - mice()
  imp.sri \leftarrow mice(dataex2[, , i], m = 20, seed = 1,
                 printFlag = FALSE, method = "norm.nob")
  imp.boot \leftarrow mice(dataex2[, , i], m = 20, seed = 1,
                   printFlag = FALSE, method = "norm.boot")
  # step 2 - with()
  fit.sri <- with(imp.sri, lm(Y ~ X))</pre>
  fit.boot <- with(imp.boot, lm(Y ~ X))</pre>
  # step 3 - pool()
  pool.sri <- summary(pool(fit.sri), conf.int = TRUE)</pre>
  pool.boot <- summary(pool(fit.boot), conf.int = TRUE)</pre>
  # if the ground truth, 3, is covered in the 95% CI of beta_1 ...
  if (pool.sri$`2.5 %`[2] <= 3 & 3 <= pool.sri$`97.5 %`[2]){
    count.sri = count.sri + 1
  count.boot = count.boot + 1
  }
}
# report the empirical coverage probability
data.frame(sri = count.sri / 100, bootstrap = count.boot / 100,
          row.names = "prob.")
```

```
## sri bootstrap
## prob. 0.88 0.95
```

#### Solution

We assume a general linear regression model  $y = x\beta + \epsilon$  for the multiple imputation procedure. Here,

- 1.  $y = (y_1, y_2, \dots, y_m)$ : response variable
- 2.  $\mathbf{x} = (\mathbf{1}, \mathbf{x_1}, \mathbf{x_2}, \cdots, \mathbf{x_n})$ : covariates (including intercept term)
- 3.  $\boldsymbol{\beta} = (\beta_0, \beta_1, \beta_2, \cdots, \beta_n)$ : coefficients
- 4.  $\epsilon \sim \mathcal{N}(0, \sigma^2)$ : noise

We start with strategy (i).

Assume that step 1 of the multiple imputation is performed and the number of imputed datasets is M. We fit the regression for each imputed dataset based on the pre-defined linear model and obtain the predicted values (point estimates)  $\hat{\boldsymbol{y}} = (\hat{y}^{(1)}, \hat{y}^{(2)}, \cdots, \hat{y}^{(M)})$ . We pool the predicted values according to Rubin's rule for point estimates:

$$\tilde{y} = \frac{1}{M} \sum_{i=1}^{M} \hat{y}^{(i)}$$

$$= \frac{1}{M} \sum_{i=1}^{M} \left( \beta_0^{(i)} + \sum_{j=1}^{n} \beta_j^{(i)} x_j \right)$$

$$= \tilde{\beta}_0 + \frac{1}{M} \sum_{i=1}^{M} \sum_{j=1}^{n} \beta_j^{(i)} x_j$$

$$= \tilde{\beta}_0 + \sum_{j=1}^{n} \left( x_j \cdot \frac{1}{M} \sum_{i=1}^{M} \beta_j^{(i)} \right)$$

$$= \tilde{\beta}_0 + \sum_{j=1}^{n} \tilde{\beta}_j x_j$$

where  $\tilde{\beta}_0 = \frac{1}{M} \sum_{i=1}^{M} \beta^{(i)}$  and  $\tilde{\beta}_j = \frac{1}{M} \sum_{i=1}^{M} \beta_j^{(i)}$  are the pooled regression coefficients with intercept.

Now we work on strategy (ii) with same notations used in (i).

Before predicting, we pool the regression coefficients from each model, i.e.  $\beta^{(i)} = (\beta_0^{(i)}, \beta_1^{(i)}, \cdots, \beta_n^{(i)}), i = 1, 2, \cdots, M$  in step 2 using Rubin's rule for point estimates. In this way, we also obtain  $\tilde{\beta}_0 = \frac{1}{M} \sum_{i=1}^{M} \beta^{(i)}$  and  $\tilde{\beta}_j = \frac{1}{M} \sum_{i=1}^{M} \beta_j^{(i)}$  and further the same predicting expression  $\tilde{y} = \tilde{\beta}_0 + \sum_{j=1}^{n} \tilde{\beta}_j x_j$  as we derived for strategy (i).

Therefore, the equations above prove that two strategies are mathematically equivalent and lead to same results and conclusions.

#### (a) - Solution

We need to prevent  $x_2$  to be imputed and only impute y and  $x_1$  variables in step 1. To this end, we pre-define the predictor matrix where the row for  $x_2$  are all 0s so that  $x_2$  will not be imputed. Then we apply MICE on the dataset and consider the interaction term  $x_1x_2$  in with() procedure.

According to the pool() procedure, we obtain the estimates and 95% confidence intervals of  $\beta_1$ ,  $\beta_2$  and  $\beta_3$  as follows (3 decimal places):

```
1. \hat{\beta}_1 = 1.411 95%CI = (1.219, 1.603)
2. \hat{\beta}_2 = 1.966 95%CI = (1.861, 2.071)
3. \hat{\beta}_3 = 0.755 95%CI = (0.642, 0.868)
```

Comparing the results above with the ground truth values ( $\beta_1 = 1, \beta_2 = 2, \beta_3 = 1$ ), we observe that only the 95% confidence interval corresponding to  $\beta_2$  covers its true value, with a reasonable estimate.

On the other hand, the estimates for imputed variables, y and  $x_1$ , are relatively inaccurate and the corresponding 95% confidence intervals fail to cover their true values.

Based on these. we consider that the *impute and then transform* method leads to biased estimates for imputed variables.

```
## term estimate 2.5 % 97.5 %

## 1 (Intercept) 1.5929831 1.404501 1.7814655

## 2 x1 1.4112333 1.219397 1.6030697

## 3 x2 1.9658191 1.860657 2.0709812

## 4 x1:x2 0.7550367 0.642302 0.8677715
```

#### (b) - Solution

## 2

## 3

## 4

Now we consider using passive imputation to impute the missing values in the interaction variable  $x_1x_2$ .

Since the original dataset does not contain the column for interaction variable, we calculate and add the interaction column into the dataset. Note that if  $x_1 = NA$ , the corresponding  $x_1x_2 = NA$  too.

To apply passive imputation, we modify the method argument for  $x_1x_2$  so that its imputation method is the interaction of  $x_1$  and  $x_2$ , i.e.  $\sim I(x_1 * x_2)$ , in mice() procedure. Moreover, we make restrictions to the predictor matrix such that y is not used to impute  $x_1x_2$  and  $x_1x_2$  is not considered when imputing  $x_1$  and  $x_2$ .

With everything ready, we implement multiple imputation on the dataset with pre-defined method and predictor matrix and obtain the estimates of each parameter with their 95% confidence intervals.

```
1. \hat{\beta}_1 = 1.193 \quad 95\%CI = (1.003, 1.382)
2. \hat{\beta}_2 = 1.996 \quad 95\%CI = (1.899, 2.094)
3. \hat{\beta}_3 = 0.874 \quad 95\%CI = (0.762, 0.987)
```

Comparing with *impute and then transform* method, we obtain a more accurate estimate for  $\beta_2$ , with a narrower 95% confidence interval including the true value, indicating higher accuracy of passive imputation.

The estimates for  $\beta_1$  and  $\beta_3$  are also closer to the ground truth values, proving the improved estimation accuracy. However, the problem that the corresponding confidence intervals do not cover the true values of two parameters still arises.

Thus, we consider that the bias caused by the model is reduce by passive imputation but still remains.

```
# calculate interaction variable and add to dataset
dataex4$x1x2 <- dataex4$x1 * dataex4$x2</pre>
# initial mice() to get the method
imp.q4b.0 <- mice(dataex4, m = 50, seed = 1, printFlag = FALSE)</pre>
# specify x1x2 is the interaction of x_1 and x_2
imp.q4b.0$method['x1x2'] <- "~I(x1*x2)"
# x_1 and x_2 should not be imputed by x_1x_2
imp.q4b.0$predictorMatrix[c("x1", "x2"), "x1x2"] <- 0
# x1x2 should not be imputed by y
imp.q4b.0$predictorMatrix["x1x2", "y"] <- 0</pre>
# final MI
imp.q4b <- mice(dataex4, m = 50, seed = 1, printFlag = FALSE,</pre>
                 predictorMatrix = imp.q4b.0$predictorMatrix,
                method = imp.q4b.0$method)
                                                      # mice() procedure
fit.q4b \leftarrow with(imp.q4b, lm(y \sim x1 + x2 + x1x2))
                                                      # with() procedure
                                                      # pool() procedure
pool.q4b <- pool(fit.q4b)</pre>
# report estimates and 95% CIs
summary(pool.q4b, conf.int = TRUE)[, c("term", "estimate", "2.5 %", "97.5 %")]
##
            term estimate
                                2.5 %
                                          97.5 %
## 1 (Intercept) 1.5534782 1.3788626 1.7280939
```

x1 1.1926170 1.0034980 1.3817360

x2 1.9964402 1.8989468 2.0939336

x1x2 0.8740573 0.7615712 0.9865434

#### (c) - Solution

In this case, we consider the interaction variable  $x_1x_2$  as just another variable, meaning that we impute the missing values in  $x_1x_2$  without depending on the missing values in  $x_1$  variable. Thus, we directly construct the linear regression over  $x_1$ ,  $x_2$  and  $x_1x_2$  in with() procedure, without any further steps.

As a result, we obtain the following estimates and 95% confidence intervals of  $\beta_1$ ,  $\beta_2$  and  $\beta_3$  (3 decimal places):

```
1. \hat{\beta}_1 = 1.000 \ 95\%CI = (0.841, 1.166)
2. \hat{\beta}_2 = 2.026 \ 95\%CI = (1.940, 2.113)
3. \hat{\beta}_3 = 1.018 \ 95\%CI = (0.930, 1.105)
```

We observe that the method treating the interaction  $x_1x_2$  as just another variable derives more accurate estimates for each parameter. Moreover, this method significantly improves the problem of model bias as the true value of each parameters are covered in its corresponding confidence interval.

In terms of the data and model in our problem, we regard *just another variable* method as the best performed imputation method among 3 candidates. However, we need to point out that this method leads to the inconsistency of imputed values as we consider the interaction term independent with missing values in other variables when imputing.

```
# consider x1x2 just as another variable
# MI
imp.q4c <- mice(dataex4, m = 50, seed = 1, printFlag = FALSE) # mice() procedure
fit.q4c <- with(imp.q4c, lm(y ~ x1 + x2 + x1x2)) # with() procedure
pool.q4c <- pool(fit.q4c) # pool() procedure

# report the estimates and 95% CIs
summary(pool.q4c, conf.int = TRUE)[, c("term", "estimate", "2.5 %", "97.5 %")]</pre>
```

```
## term estimate 2.5 % 97.5 %

## 1 (Intercept) 1.499714 1.3452011 1.654227

## 2 x1 1.003930 0.8414967 1.166363

## 3 x2 2.026180 1.9398113 2.112548

## 4 x1x2 1.017793 0.9303479 1.105238
```

# (d) - Solution

As mentioned in 4.(c), though just another variable approach obtains unbiased estimates for each parameter in the linear regression, the inner dependence between the interaction term  $x_1x_2$  and  $x_1$  is violated. In other words, we impute the missing values of  $x_1x_2$  independently from  $x_1$ , without considering the deterministic relationship between them.

#### Solution

#### Exploratory Data Analysis

We start our study with inspecting our data. Our dataset consists of 500 cases with 12 variables including 8 continuous variables and 4 factors:

• wgt: weight in kg

• gender: male vs female

• bili: bilirubin concentration in mg/dL

• age: in years

• chol: total serum cholesterol in mg/dL,

• HDL: High-density lipoprotein cholesterol in mg/dL,

• hgt: height in metres,

• educ: educational status; 5 ordered categories,

• race: 5 unordered categories,

• SBP: systolic blood pressure in mmHg,

• hypten: hypertensive status; binary,

• WC: waist circumference in cm

According to the results from summary() function, variables bili, chol, HDL, hgt, educ, SBP, hypten and WC contains missing values. Note that some missing values in the dataset are presented by the string "NaN" rather than NA. Since we are going to use mice package for multiple imputation, which expects missing values coded as NA, we convert "NaN" to NA before any further steps, also for the consistency of notations.

We investigate the missing data patterns intuitively using md\_pattern() function from JointAI package. We can observe that the number of missing values for each variable is not large (47 for bili at most and 1 for educ at least) and the number of fully observed cases is 411 (out of 500). (We'll revisit this fact later when determining the number of imputed datasets, m)

Moreover, we visualize the correlations between variables using corrplot() function from corrplot package. Recall that our model of interest is

$$wqt = \beta_0 + \beta_1 qender + \beta_2 aqe + \beta_3 hqt + \beta_4 WC + \epsilon, \quad \epsilon \sim N(0, \sigma^2)$$

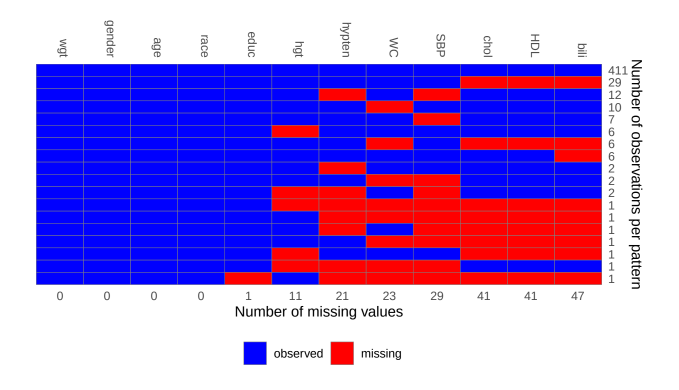
According to the heat map below, we observe that there exists a strong positive correlation between the response variable wgt and WC. gender and hgt also have moderate correlations with wgt. However, the independent variable age shows a relatively low correlation with wgt, with a correlation efficient of 0.069 (3 decimal places), which may lead to poor regression performance later. Another noteworthy fact is that hgt has a strong negative correlation with gender, which will be revisited later as it affects the imputation results of hgt.

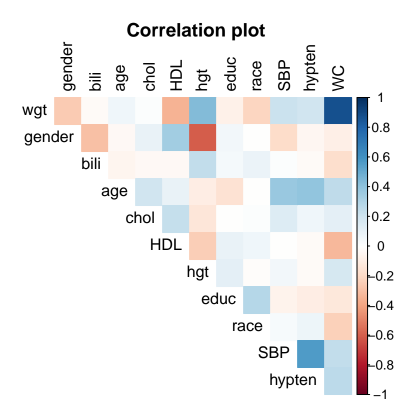
Finally, we explore the distribution of each variable by visualizing it with plot\_all() function from JointAI package. Continuous variables, except hgt, appear to have right-skewed distributions. Thus, using normal distribution for imputation method probably results in poor approximations and we keep using predictive mean matching (pmm) as default. Since we consider hgt can be well approximated by normal distribution, the imputation method of it will be modified to norm, the normal linear stochastic regression imputation regarding the uncertainty of data.

```
# read dataset into R
load("NHANES2.Rdata")
# data structure and summary
str(NHANES2)
                 500 obs. of 12 variables:
## 'data.frame':
## $ wgt : num 78 78 75.3 90.7 112 ...
## $ gender: Factor w/ 2 levels "male", "female": 1 1 2 1 2 1 2 2 1 1 ...
## $ bili : num 1.1 0.7 0.5 0.8 0.6 0.7 1.1 0.8 0.8 0.5 ...
## $ age
          : num 67 39 64 36 33 62 56 63 55 20 ...
## $ chol : num 6.13 4.65 4.14 3.47 6.31 4.47 6.41 5.51 7.01 3.75 ...
## $ HDL
         : num 1.09 1.14 1.29 1.37 1.27 0.85 1.81 2.38 2.79 1.03 ...
         : num 1.75 1.78 1.63 1.93 1.73 ...
## $ educ : Ord.factor w/ 5 levels "Less than 9th grade" < ..: 5 3 5 4 4 3 4 5 4 2 ...
## $ race : Factor w/ 5 levels "Mexican American",..: 5 3 5 3 4 5 4 5 3 3 ...
## $ SBP : num 139 103 NaN 115 107 ...
## $ hypten: Factor w/ 2 levels "no", "yes": 2 1 2 2 1 2 NA 1 2 1 ...
          : num 91.6 84.5 91.6 95.4 119.6 ...
summary(NHANES2)
##
                      gender
                                    bili
                                                                   chol
        wgt
                                                    age
## Min. : 39.01
                   male :252 Min. :0.2000 Min. :20.00 Min. : 2.07
## 1st Qu.: 65.20
                   female:248 1st Qu.:0.6000 1st Qu.:31.00
                                                              1st Qu.: 4.27
## Median : 76.20
                               Median :0.7000 Median :43.00
                                                              Median : 4.86
## Mean : 78.25
                               Mean :0.7404
                                               Mean :45.02
                                                              Mean : 5.00
## 3rd Qu.: 86.41
                                3rd Qu.:0.9000
                                               3rd Qu.:58.00
                                                              3rd Qu.: 5.64
## Max. :167.38
                               Max. :2.9000
                                               Max. :79.00
                                                              Max. :10.68
                                                               NA's
##
                                NA's
                                    :47
                                                                    :41
##
        HDL
                       hgt
                                                  educ
## Min.
                 Min. :1.397 Less than 9th grade : 31
         :0.360
## 1st Qu.:1.110 1st Qu.:1.626
                                 9-11th grade
## Median: 1.320 Median: 1.676 High school graduate: 115
## Mean :1.395
                 Mean :1.687
                                 some college
                                                    :148
                  3rd Qu.:1.753
## 3rd Qu.:1.590
                                 College or above
                                                   :136
## Max. :3.130
                  Max. :1.930
                                 NA's
                                                    : 1
## NA's
         :41
                  NA's :11
##
                  race
                               SBP
                                           hypten
                                                           WC
## Mexican American : 52 Min. : 81.33
                                          no :354 Min. : 61.90
## Other Hispanic
                  : 58 1st Qu.:109.00
                                           yes:125 1st Qu.: 84.80
## Non-Hispanic White:182 Median :118.67
                                           NA's: 21 Median: 95.00
## Non-Hispanic Black:112 Mean
                                :120.05
                                                     Mean
                                                           : 96.07
## other
                   : 96
                           3rd Qu.:128.67
                                                     3rd Qu.:104.80
##
                           Max.
                                 :202.00
                                                     Max.
                                                            :154.70
##
                           NA's
                                  :29
                                                     NA's
                                                            :23
# convert NaN to NA
NHANES2 <- na_if(NHANES2, "NaN")</pre>
# transform factors to numeric
col.factor <- which(sapply(NHANES2, is.factor))</pre>
```

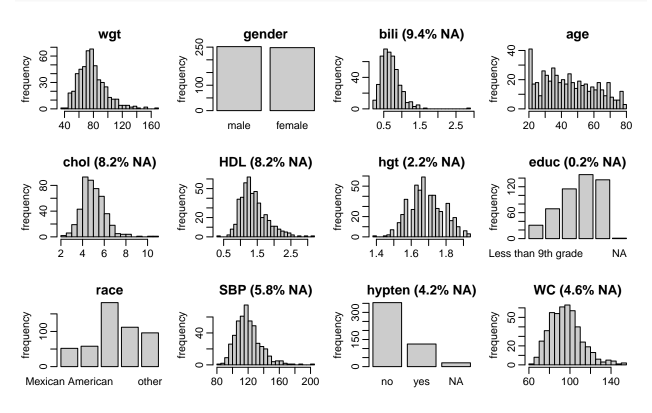
```
NHANES2.numeric <- NHANES2
NHANES2.numeric[, col.factor] <- sapply(NHANES2[, col.factor], as.numeric)
```

```
# visualize missing pattern
mdp <- md_pattern(NHANES2, pattern = TRUE, color = c("blue", "red"))
mdp$plot</pre>
```





# visualize the distribution of observed values in each variable
par(mar = c(3, 3, 2, 1), mgp = c(2, 0.6, 0))
plot\_all(NHANES2, breaks = 30, ncol = 4)



#### Multiple Imputation

We begin multiple imputation with a dry/set-up run of mice() procedure. From the imputation summary, we observe the default imputation method for continuous variables is pmm. As mentioned before, we modify this imputation method for hgt to norm. Moreover, since hgt represents the individual's height in metres, we need to restrict the support the it. Here, we set a reasonable range of hgt to [0.5, 2.8].

For factors, the methods are different depending on the number of classes in the factor. logreg is used for hypten as it is a binary variable, while polr is used for educ since this variable consists of multiple categories.

Before any further imputation checks, we determine a proper number of imputed datasets, i.e. m argument in mice() function, based on both accuracy and statistical efficiency. To this end, we run complete multiple imputation procedures including mice(), with() and pool() with a set of candidates of  $m \in \{5, 10, 15, 20, 25\}$  and repeat the procedures for  $seed \in \{1, 2, 3, 4, 5, 6\}$ . We report the estimates for the regression coefficients as well as the running time using different seed for each candidate of m.

As the data frames shown below, the variance of estimated regression coefficients gets smaller as m increases. However, this improvement becomes insignificant from m=20 to m=25. Meanwhile, we notice that the running time of each multiple imputation increases by around 10 seconds as the value of m increases by 5 (50 seconds for m=25).

Therefore, we prefer a multiple imputation on our dataset with m = 20, which obtains both computationally efficient and robust to random variation due to seed choice. Meanwhile, we set maxit argument to 20 and keep random seed as 1 for model reproducibility, without the loss of generality.

[Note: There is no doubt to use larger value of m (e.g. 50, 100) to improve the model accuracy. However, in this case, the percentage of missing values in each variable is low (9.4% for bili at most) and over 80% cases in the dataset are complete. Thus, we believe smaller m can satisfy the requirements of both less variation and statistical efficiency.]

```
# multiple imputation
# dry mice() procedure
imp0 <- mice(data = NHANES2, maxit = 0)
imp0</pre>
```

```
## Class: mids
## Number of multiple imputations:
##
   Imputation methods:
                gender
                                                              HDL
##
         wgt
                             bili
                                         age
                                                  chol
                                                                        hgt
                                                                                  educ
                                                 "pmm"
##
                            "pmm"
                                                           "pmm"
                                                                      "pmm"
                                                                               "polr"
                                          WC
##
                   SBP
                          hypten
        race
                 "pmm" "logreg"
##
                                      "pmm"
## PredictorMatrix:
##
           wgt gender bili age chol HDL hgt educ race SBP hypten WC
## wgt
             0
                                1
                                           1
                                                1
                                                           1
                                                                1
                                                                        1
                                                                            1
                      1
                            1
                                      1
                                                     1
                      0
                                           1
## gender
              1
                            1
                                1
                                      1
                                                1
                                                     1
                                                           1
                                                                1
                                                                        1
                                                                            1
## bili
              1
                      1
                            0
                                1
                                      1
                                           1
                                                1
                                                     1
                                                           1
                                                                1
                                                                        1
                                                                            1
## age
              1
                      1
                            1
                                0
                                      1
                                           1
                                                1
                                                     1
                                                           1
                                                                1
                                                                        1
                                                                           1
## chol
              1
                      1
                                1
                                      0
                                           1
                                                1
                                                     1
                                                           1
                                                                1
                                                                        1
                                                                           1
                            1
## HDL
                      1
                                           0
              1
                            1
                                1
                                      1
                                                1
                                                     1
                                                           1
                                                                1
                                                                        1
```

```
# modify the imputation method of 'hgt'
imp0$method["hgt"] <- "norm"
# restrict the support of 'hgt'
imp0$post["hgt"] <- "imp[[j]][,i] <- squeeze(imp[[j]][,i], c(0.5, 2.8))"</pre>
```

```
\# MI for m = \{5, 10, 20\} with seed = \{1, 2, 3, 4, 5, 6\}
m.set \leftarrow c(5, 10, 15, 20, 25)
seed.set \leftarrow c(1, 2, 3, 4, 5,6)
for (i in m.set){
  est.df <- NULL
  for (j in seed.set){
    time.start <- Sys.time()</pre>
    imp.q5 <- mice(NHANES2, maxit = 20, m = i, seed = j,</pre>
                   method = imp0$method,
                   post = imp0$post, printFlag = FALSE)
    fit.q5 <- with(imp.q5, lm(wgt ~ gender + age + hgt + WC))
    pool.q5 <- pool(fit.q5)</pre>
    time.end <- Sys.time()</pre>
    df <- data.frame(seed = j,</pre>
                      genderfemale = pool.q5$pooled[2, "estimate"],
                      age = pool.q5$pooled[3, "estimate"],
                     hgt = pool.q5$pooled[4, "estimate"],
                      WC = pool.q5$pooled[5, "estimate"],
                      running.time = time.end - time.start)
    est.df <- rbind(est.df, df)
  cat("regression estimates (m = ", i, "): \n", sep = "")
  print(est.df)
## regression estimates (m = 5):
     seed genderfemale
                                        hgt
                                                   WC running.time
                               age
## 1
             -1.369652 -0.1588398 52.46588 1.026546 9.878979 secs
## 2
             -1.286194 -0.1565471 53.26393 1.026603 9.476264 secs
## 3
             -1.363540 -0.1606729 52.83642 1.023533 9.574917 secs
## 4
             -1.391903 -0.1611415 52.35877 1.028648 9.667157 secs
## 5
             -1.331400 -0.1588171 52.89258 1.025179 9.555487 secs
             -1.334292 -0.1569351 52.97270 1.025940 9.628549 secs
## 6
        6
## regression estimates (m = 10):
     seed genderfemale
                               age
                                        hgt
                                                   WC running.time
## 1
             -1.318411 -0.1567395 52.58736 1.026748 19.14589 secs
             -1.305026 -0.1589174 52.90901 1.027096 19.08154 secs
## 2
## 3
             -1.362416 -0.1546543 52.36162 1.024514 18.83958 secs
## 4
             -1.420964 -0.1583875 52.39275 1.025138 18.77708 secs
             -1.306057 -0.1576264 52.85176 1.024285 20.72711 secs
## 5
        5
             -1.368114 -0.1558868 52.13114 1.026035 19.81665 secs
## 6
## regression estimates (m = 15):
     seed genderfemale
                                                   WC running.time
                                        hgt
                               age
## 1
             -1.416849 -0.1592273 52.25315 1.026394 28.42731 secs
## 2
             -1.361419 -0.1571292 52.56111 1.025913 28.36120 secs
## 3
             -1.324025 -0.1561752 52.43867 1.025362 29.15734 secs
             -1.331270 -0.1579811 52.66479 1.024820 28.81583 secs
             -1.400804 -0.1581106 52.03306 1.025732 29.88852 secs
## 5
        5
        6
             -1.351421 -0.1570642 52.42774 1.025911 29.38083 secs
## regression estimates (m = 20):
```

hgt

-1.323852 -0.1555430 52.49801 1.024904 39.87659 secs

age

WC running.time

seed genderfemale

## 1

```
## 2
            -1.363742 -0.1569720 52.70480 1.025105 38.79402 secs
## 3
            -1.346148 -0.1575175 52.72683 1.025153 39.43095 secs
        3
## 4
            -1.347170 -0.1561035 52.44223 1.025811 38.07592 secs
## 5
             -1.368667 -0.1584354 52.42954 1.025529 39.34228 secs
        5
             -1.373698 -0.1594694 52.28251 1.026873 38.87875 secs
## 6
## regression estimates (m = 25):
     seed genderfemale
                                                  WC running.time
                              age
                                       hgt
             -1.354632 -0.1591073 52.48408 1.025561 47.51230 secs
## 1
## 2
        2
             -1.319659 -0.1570791 52.70967 1.025361 49.73243 secs
## 3
            -1.358916 -0.1568233 52.43961 1.025951 48.72103 secs
## 4
          -1.375729 -0.1576124 52.16183 1.025448 48.49303 secs
## 5
        5
             -1.333896 -0.1579284 52.52684 1.025330 48.53448 secs
## 6
             -1.311311 -0.1584013 52.75073 1.024948 48.31162 secs
\# MI with m = 20 and seed = 1
imp.q5 \leftarrow mice(NHANES2, maxit = 20, m = 20, seed = 1,
               method = imp0$method,
               post = imp0$post, printFlag = FALSE)
fit.q5 <- with(imp.q5, lm(wgt ~ gender + age + hgt + WC))</pre>
pool.q5 <- pool(fit.q5)</pre>
```

#### Imputation Check

After applying a multiple imputation with proper arguments to our data, we conduct necessary checks.

We first check if mice() found any problems during imputation with object loggedEvents and obtain the result NULL, indicating no problems detected. Furthermore, we check the Monte Carlo chains to ensure the convergence across all 12 variables. From the plots with 20 MC chains for each variable, the there exists no obvious pattern among chains, suggesting good convergence for all variables. Note that the plot for the standard deviation of educ is blank as there is only 1 missing values in this column.

Next, we inspect if the distribution of the imputed values agree with the distribution of the observed values, using densityplot() as well as bwplot() functions from mice package for continuous variables and propplot() function for categorical variables.

For continuous variables, except hgt, comparing with the distribution of their observed values (blue), their imputed values follow the similar distribution (red). On the other hand, we can observe a clear left-shifted distribution of the imputed values of hgt.

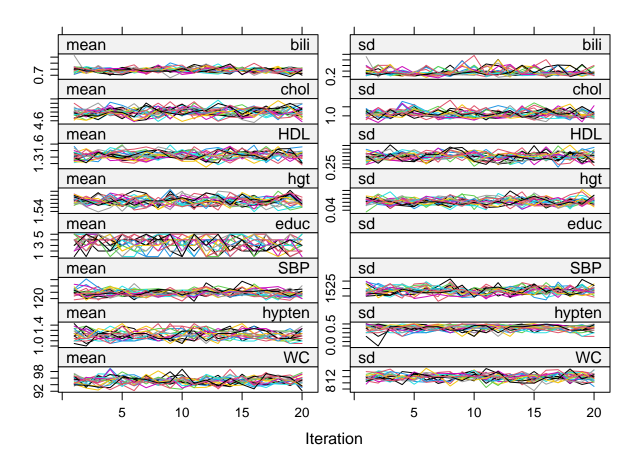
Recall that we observed high correlation between hgt and gender from the heat map. We guess this is the main reason for the shifted distribution of imputed hgt. To prove this, we next make a density plot of hgt, conditional on gender. As a result, we observe that gender does strongly influences the imputed datasets. Specifically, values in male group has much more narrow, concentrated distributions while the distribution in female group is wider.

For categorical variables, the distribution of imputed values of **educ** appears to be abnormal simply because we only imputed 1 value each time. The same reason can also explain the discrepancies between the observed and imputed values for hypten.

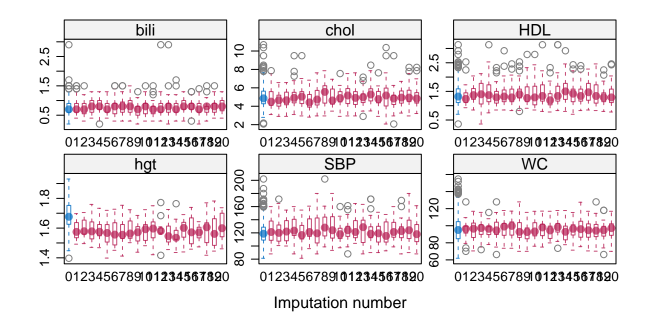
```
# necessary checks for mice() procedure
# problems check during mice() procedure
imp.q5$loggedEvents
```

## NULL

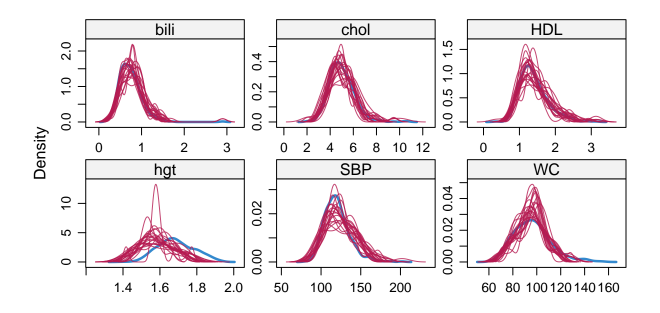
```
# convergence check
plot(imp.q5, layout = c(2, 8))
```



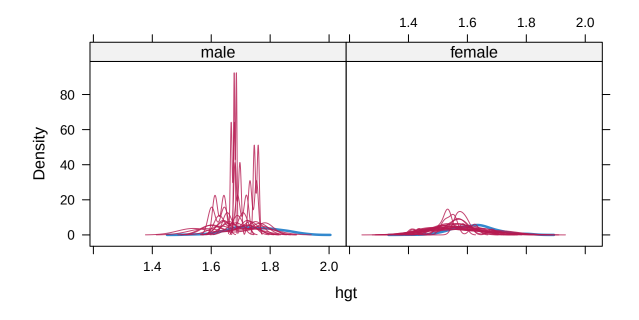
```
# distribution check
# case for continuous variables
bwplot(imp.q5)[c(2, 4, 5, 6, 7, 8)]
```



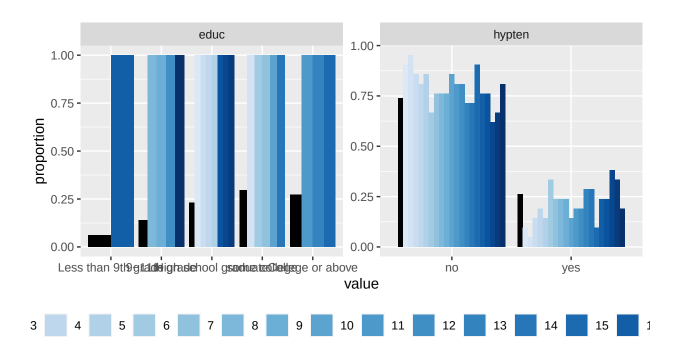
# densityplot(imp.q5)



# distribution of hgt conditional on gender
densityplot(imp.q5, ~ hgt | gender)



# case for categorised variables
propplot(imp.q5)



#### Regression Check

Before we reach the final conclusions, we also need to check if our model of interest fits well the *completed* data. Moreover, we check if model performances across different subsets are consistent. Thus, we display the regression summaries for both subset 1 and 20 with plots.

According to the regression summaries, we observe that all independent variables, except genderfemale (gender), are significant with relatively low p-values. Moreover,  $R^2$  around 85 indicates good model explanation on response variable. Looking at the residual plots, we cannot observe obvious pattern of residuals, suggesting the linear assumption is met. The well-performed normal Q-Q plots shows the normality of error terms is satisfied. Thus, we consider our model to be well fitted.

Note that the insignificance of genderfemale (gender) mainly because its high correlation with hgt. High correlation indicates there exists high collinearity between two variables, where one covers most information of the other. As a result, one variable becomes insignificant when including both of them into regression model.

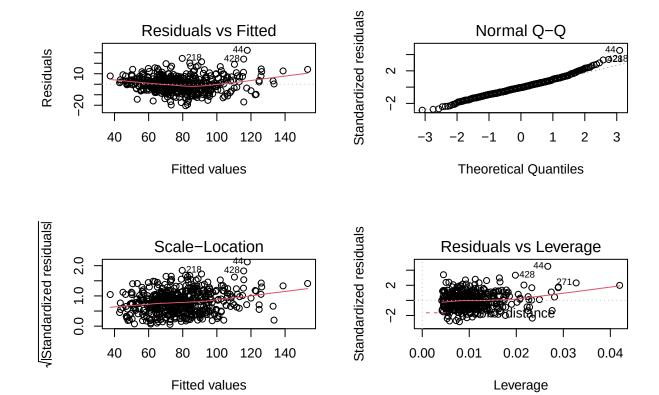
The strong linear relationship between two variables influence not only the distribution of imputed values of hgt, but also the contribution to wgt of gender. Though the regression model appears to be good, we still need to focus on solving this problem in future research to improve model performance.

Finally we pool the regression results and report the estimates of regression coefficients with their 95% confidence intervals and p-values.

```
# with() procedure (regression) check
summary(fit.q5$analyses[[1]])
```

```
##
## Call:
## lm(formula = wgt ~ gender + age + hgt + WC)
##
## Residuals:
##
       Min
                1Q
                   Median
                                3Q
                                       Max
                    -0.385
##
  -20.463
           -4.585
                             4.162
                                    32.323
##
## Coefficients:
##
                  Estimate Std. Error t value Pr(>|t|)
## (Intercept) -104.98644
                              7.55808 -13.891 < 2e-16 ***
  genderfemale
                  -1.23947
                              0.83155
                                       -1.491
                                                  0.137
                              0.02106
                                       -7.288 1.25e-12 ***
## age
                  -0.15348
## hgt
                  55.12737
                              4.30910
                                       12.793 < 2e-16 ***
## WC
                   1.01901
                              0.02225
                                       45.792 < 2e-16 ***
##
                  0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Signif. codes:
##
## Residual standard error: 7.259 on 495 degrees of freedom
## Multiple R-squared: 0.8543, Adjusted R-squared: 0.8531
## F-statistic: 725.6 on 4 and 495 DF, p-value: < 2.2e-16
par(mfrow = c(2, 2))
```

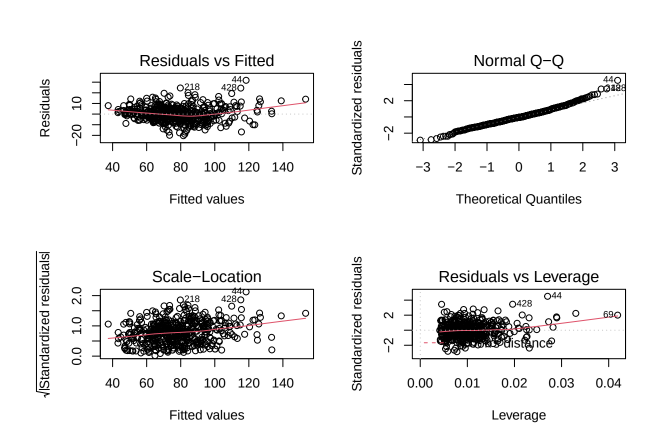
```
par(mfrow = c(2, 2))
plot(fit.q5$analyses[[1]])
```



#### summary(fit.q5\$analyses[[20]])

```
##
  lm(formula = wgt ~ gender + age + hgt + WC)
##
## Residuals:
       Min
                1Q
                   Median
                                3Q
                                       Max
## -20.321 -4.552
                    -0.353
                             3.933
                                    31.751
##
## Coefficients:
                  Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                -101.63310
                              7.39209 -13.749
                                              < 2e-16 ***
## genderfemale
                  -1.31788
                              0.81265
                                       -1.622
                                                  0.106
                  -0.15988
                              0.02073
                                       -7.714 6.76e-14 ***
## age
                  52.72579
                                       12.494
## hgt
                              4.22001
                                               < 2e-16 ***
## WC
                                       46.843
                                               < 2e-16 ***
                   1.02899
                              0.02197
##
                   0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Signif. codes:
##
## Residual standard error: 7.138 on 495 degrees of freedom
## Multiple R-squared: 0.8591, Adjusted R-squared: 0.858
## F-statistic: 754.6 on 4 and 495 DF, p-value: < 2.2e-16
```

```
par(mfrow = c(2, 2))
plot(fit.q5$analyses[[20]])
```



# # pool() procedure check pool.q5

```
## Class: mipo
##
                   m
                        estimate
                                          ubar
                                                                        t dfcom
## 1 (Intercept) 20 -100.960341 5.526751e+01 3.949302e+00 5.941428e+01
                                                                            495
## 2 genderfemale 20
                       -1.323852 6.738407e-01 1.415847e-02 6.887071e-01
                                                                            495
## 3
                       -0.155543 4.346036e-04 2.988707e-05 4.659850e-04
                                                                            495
              age 20
                       52.498011 1.805798e+01 1.323640e+00 1.944780e+01
## 4
              hgt 20
                                                                            495
## 5
               WC 20
                        1.024904 4.878153e-04 1.249150e-05 5.009314e-04
                                                                            495
           df
                     riv
                              lambda
## 1 410.3546 0.07503083 0.06979412 0.07429488
## 2 476.7304 0.02206217 0.02158594 0.02566495
## 3 414.3349 0.07220702 0.06734429 0.07181387
## 4 407.6217 0.07696442 0.07146422 0.07598680
## 5 471.9279 0.02688739 0.02618339 0.03028429
summary.q5 <- summary(pool.q5, conf.int = TRUE)</pre>
summary.q5
```

## term estimate std.error statistic df p.value

```
## 1 (Intercept) -100.960341 7.70806594 -13.098012 410.3546 0.000000e+00
## 2 genderfemale
                    -1.323852 0.82988381 -1.595226 476.7304 1.113244e-01
## 3
                    -0.155543 0.02158669 -7.205505 414.3349 2.753797e-12
              age
## 4
                    52.498011 4.40996593 11.904403 407.6217 0.000000e+00
              hgt
## 5
               WC
                     1.024904 0.02238150 45.792460 471.9279 0.000000e+00
##
            2.5 %
                       97.5 %
## 1 -116.1125628 -85.8081195
                    0.3068302
## 2
       -2.9545344
## 3
      -0.1979761 -0.1131099
## 4
       43.8288964 61.1671255
## 5
        0.9809240
                   1.0688835
df <- data.frame(estimate = summary.q5[, 2],</pre>
                 lower = summary.q5[, 7],
                 upper = summary.q5[, 8],
                 p.value = summary.q5[, 6],
                 row.names = c("$\\beta_0$", "$\\beta_1$",
                               "$\\beta_2$", "$\\beta_3$", "$\\beta_4$"))
colnames(df) <- c("estimate", "2.5% quantile", "97.5% quantile", "p-value")</pre>
knitr::kable(df, escape = FALSE, digits = 3,
             caption = "Regression coefficients, 95% CIs and p-values")
```

Table 1: Regression coefficients, 95% CIs and p-values

	estimate	2.5% quantile	97.5% quantile	p-value
$\beta_0$	-100.960	-116.113	-85.808	0.000
$\beta_1$	-1.324	-2.955	0.307	0.111
$\beta_2$	-0.156	-0.198	-0.113	0.000
$\beta_3$	52.498	43.829	61.167	0.000
$\beta_4$	1.025	0.981	1.069	0.000