# Assignment 3 Solution

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All code available on github at: https://github.com/TedOiler/ida\_assignment\_3

## Question 1

#### 1a.

After a first exploration of the dataset we conclude that the percentage of missing cases on the nhanes dataset is 48%

#### 1b.

The proportions of variance due to the missing data for each parameter are given by the lambda term. Specifically:

```
age: 0.6864 hyp: 0.3504 chl: 0.3041
```

Therefore the age parameter seem to be most affected by the nonresponse.

```
# m = 5 (default value)
bmi_hat = pool(with(mice(nhanes, seed=1, printFlag=FALSE), lm(bmi ~ age + hyp + chl)))
bmi_hat[,3][c(1,3,10)]
```

```
## term estimate lambda

## 1 (Intercept) 19.61789252 0.08938989

## 2 age -3.55287155 0.68640637

## 3 hyp 2.19701748 0.35043452

## 4 chl 0.05378081 0.30408063
```

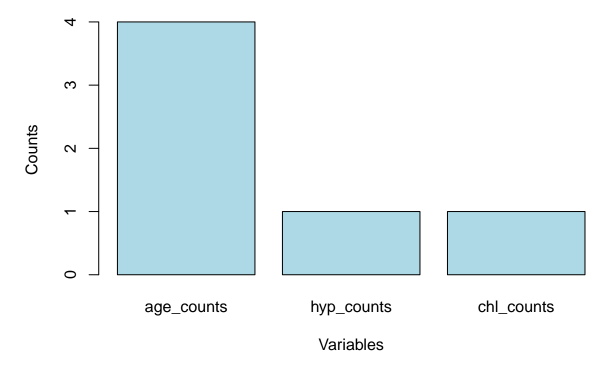
#### 1c.

The script below tries to automate and visualize the process in order to be able to potentially scale it. For the six iteration of random seeds and the tree variables, we notice that on seeds 1, 2, 3 and 6 the age variable is the one with the biggest lambda parameter. On seed = 4 the chl variable has the biggest lambda and on seed = 5 it's the hyp parameter. On my tests up to 1e2 different seeds (not displayed here), the distribution was: age: 46%, hyp: 26%, chl: 28% So finally, the results are dependent on the seed.

```
analyze <- function(data, seeds, models=5){</pre>
  num_of_seeds = seeds
  dataset = data
  m = models
  # Initialize a dataframe to store results
  df <- data.frame(age = rep(OL, num_of_seeds),</pre>
                   hyp = rep(OL, num_of_seeds),
                    chl = rep(OL, num_of_seeds),
                    age_lambda = rep(OL, num_of_seeds),
                   hyp_lambda = rep(OL, num_of_seeds),
                    chl_lambda = rep(OL, num_of_seeds))
  max_lambda <- c(1:num_of_seeds)*0</pre>
  for (i in c(1:num_of_seeds)){
    bmi_hat = pool(with(mice(dataset, seed=i, printFlag=FALSE, m=m), lm(bmi ~ age + hyp + chl)))
    df[i,1:3] <- bmi_hat[2:4,3][1:3,3] # store estimates
    df[i,4:6] <- bmi_hat[2:4,3][1:3,10] # store lambda values
    # Not the cleanest code
    \# Logic loop in order to find the maximum per row of the above dataframe of the lambda parameters s
    max_lambda[i] <- which.max(t(df[4:6])[,i])</pre>
    if (max_lambda[i]==1){
      max_lambda[i] <- 'age'</pre>
    else if (max_lambda[i]==2){
      max_lambda[i] <- 'hyp'</pre>
    }
    else {
      max_lambda[i] <- 'chl'</pre>
    }
  }
  # Counts per variable.
  # Shows on num_of_seeds iterations which variable had the biggest lambda score and adds them up.
  counts <- data.frame(age_counts = length(which(max_lambda == 'age')),</pre>
                        hyp_counts = length(which(max_lambda == 'hyp')),
                        chl_counts = length(which(max_lambda == 'chl')))
  print(df)
  cat('\n')
  print(max_lambda)
  cat('\n')
  print(counts)
  barplot(as.matrix(counts),
          col='lightblue',
          main = 'Biggest lambda per variable',
          xlab = 'Variables',
          ylab = 'Counts')
}
analyze(data=nhanes, seeds=6)
```

```
##
                                chl age_lambda hyp_lambda chl_lambda
                    hyp
           age
## 1 -3.552872 2.197017 0.05378081
                                     0.6864064
                                                0.3504345
                                                            0.3040806
                                                0.1430995
## 2 -4.061509 1.530476 0.06283490
                                     0.4033924
                                                            0.2959966
## 3 -3.857533 1.352812 0.05872834
                                     0.5895051
                                                 0.4101152
                                                            0.5621346
## 4 -3.506034 2.750530 0.04920611
                                     0.2189333
                                                 0.1961083
                                                            0.3305334
## 5 -3.496722 1.509548 0.06081272
                                     0.4511896
                                                0.5942866
                                                            0.2346065
  6 -2.921414 1.224746 0.04949218
                                     0.6549523
                                                0.2960364
                                                            0.5196295
##
##
   [1] "age" "age" "age" "chl" "hyp" "age"
##
##
     age_counts hyp_counts chl_counts
## 1
```

# Biggest lambda per variable



### 1d.

With the above work on the function, now we can just call the function analyze with the extra argument models=10 and we have our results so we can focus on the interpretation.

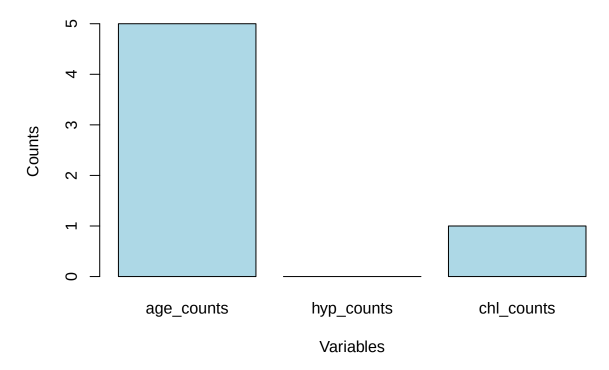
For M=100 we see exactly what we expected, which is the reduction of the variability since age is most effected by the nonresponse. However, even for a small number of seeds there is a noticeable increase in time until execution of the models, since we are now calculating 3 coefficients for 6 seeds and 100 models. The overall  $\mathcal{O}$  is  $\mathcal{O}(CSM)$  where C: coefficients, S: number of seeds,M: number of models, which is expensive. Our dataset is small so the calculations of all the coefficients don't take that much time, but in order to calculate the coefficients which are given from:  $b = (X^TX)^{-1}X^TY$  we need  $\mathcal{O}(N^3)$  where N: number or rows in dataset X which can get be very expensive very quickly.

When comparing between the model with M=5 and M=100 for this specific dataset, I would prefer the M=100. But I think that in reality I would chose a more moderate M=50.

```
analyze(data=nhanes, seeds=6, models=100)
```

```
hyp
                                chl age_lambda hyp_lambda chl_lambda
           age
## 1 -3.641255 1.685794 0.05449706
                                    0.4324680
                                                0.2915346
                                                           0.3217837
## 2 -3.633789 1.719915 0.05352120
                                    0.4031077
                                                           0.2939693
## 3 -3.557061 1.557562 0.05445409
                                    0.3093072
                                                0.2425105
                                                           0.3281911
## 4 -3.648153 1.659470 0.05511595
                                    0.3943223
                                                0.2565132
                                                           0.2835232
## 5 -3.762972 1.802832 0.05534382
                                    0.3322570
                                                0.2893046
                                                           0.2461956
## 6 -3.593092 1.862196 0.05319323
                                    0.4430300
                                                0.2860700
                                                           0.3113085
##
  [1] "age" "age" "chl" "age" "age" "age"
##
##
     age_counts hyp_counts chl_counts
## 1
              5
```

# Biggest lambda per variable



# Question 2

We use the method argument in order to distinguish the two imputation methods form each other, the Stochastic Regression Imputation as well as the bootstrap method. We have them taken the attributes of 2.5 % and 97.5 % Confidence intervals of the two methods for each dataset and stored them. It the true value of 3 is contained within the interval then we can check it with 2 if statements for each method. We

observe that the bootstraps method CI is typically wider than the SRI's CI, so it 'misses' the true value of the distribution more. sri = 88% boot = 95%

```
load('dataex2.Rdata')
sri = 0
boot = 0
models = 20 # fixed value.
set_seed = 1 # fixed value for reproducability of the result.
beta_1 = 3
for(i in c(1:100)){
  sri_imp = mice(dataex2[,,i],
                 m=models,
                 seed=set seed,
                 printFlag=FALSE,
                 method="norm.nob")
  boot_imp = mice(dataex2[,,i],
                  m=models,
                  seed=set_seed,
                  printFlag=FALSE,
                  method="norm.boot")
  sri_results = pool(with(sri_imp, lm(Y ~ X)))
  boot_results = pool(with(boot_imp, lm(Y ~ X)))
  ci_sri_low = summary(sri_results , conf.int=TRUE)$'2.5 %'[2]
  ci_sri_hi = summary(sri_results , conf.int=TRUE)$'97.5 %'[2]
  ci_boot_low = summary(boot_results, conf.int=TRUE)$'2.5 %'[2]
  ci_boot_hi = summary(boot_results, conf.int=TRUE)$'97.5 %'[2]
  if(ci_sri_low<=beta_1 & ci_sri_hi>beta_1){
   sri = sri + 1
  }
  if(ci_boot_low<=beta_1 & ci_boot_hi>beta_1){
   boot = boot + 1
 }
}
sri
```

```
## [1] 88
```

boot

## [1] 95

# Question 3

In order to answer this question we need to carefully define the problem in a mathematical way. After that the solution should follow.

• Case (i)

$$\begin{split} \hat{y} &= \frac{1}{M} \Sigma_{m=1}^{M} \hat{y_i}^{(m)} \quad (1) \\ \text{where} \\ \hat{y_i}^{(1)} &= \hat{\beta_0}^{(1)} + \hat{\beta_1}^{(1)} x_{1i} + \hat{\beta_2}^{(1)} x_{2i} + \ldots + \hat{\beta_k}^{(1)} x_{ki} + \epsilon_i \\ \hat{y_i}^{(2)} &= \hat{\beta_0}^{(2)} + \hat{\beta_1}^{(2)} x_{1i} + \hat{\beta_2}^{(2)} x_{2i} + \ldots + \hat{\beta_k}^{(2)} x_{ki} + \epsilon_i \\ & \\ & \\ \vdots \\ \hat{y_i}^{(M)} &= \hat{\beta_0}^{(M)} + \hat{\beta_1}^{(M)} x_{1i} + \hat{\beta_2}^{(M)} x_{2i} + \ldots + \hat{\beta_k}^{(M)} x_{ki} + \epsilon_i \\ \text{and} \quad \epsilon_i \overset{\text{iid}}{\sim} N(0, \sigma_\epsilon^2) \end{split}$$

• Case (ii)

$$\hat{\beta_0}^* = \frac{1}{M} \sum_{m=1}^M \hat{\beta_0}^{(m)}, ..., \hat{\beta_k}^* = \frac{1}{M} \sum_{m=1}^M \hat{\beta_k}^{(m)} \Rightarrow \hat{y_i}^* = \hat{\beta_0}^* + \hat{\beta_1}^* x_{1i} + \hat{\beta_2}^* x_{2i} + ... + + \hat{\beta_k}^* x_{ki} + u_i \quad (2)$$
and  $u_i \stackrel{\text{iid}}{\sim} N(0, \sigma_u^2)$ 

So if we work our way from (1) to (2) algebraically then we can show that Case (i) and Case (ii) coincide:

$$\stackrel{(1)}{\Rightarrow} \hat{y} = \frac{1}{M} \sum_{m=1}^{M} \hat{y_i}^{(m)} = \frac{1}{M} \sum_{m=1}^{M} [\hat{\beta_0}^{(m)} + \hat{\beta_1}^{(m)} x_{1i} + \hat{\beta_2}^{(m)} x_{2i} + \dots + \hat{\beta_k}^{(m)} x_{ki} + \epsilon_i] = \frac{1}{M} \sum_{m=1}^{M} \hat{\beta_0}^{(m)} + \frac{1}{M} \sum_{m=1}^{M} \hat{\beta_1}^{(m)} x_{1i} + \frac{1}{M} \sum_{m=1}^{M} \hat{\beta_2}^{(m)} x_{2i} + \dots + \frac{1}{M} \sum_{m=1}^{M} \hat{\beta_k}^{(m)} x_{ki} + m \epsilon_i$$
 (3)

We set 
$$m\epsilon_i = u_i \Rightarrow u_i \stackrel{\text{iid}}{\sim} N(mE[\epsilon_i], \sigma_{\epsilon}^2) = N(0, \sigma_u^2)$$
, with  $\sigma_u^2 = \sigma_{\epsilon}^2$  (4)  
(3), (4)  $\Rightarrow \hat{y}_i = \hat{\beta_0}^* + \hat{\beta_1}^* x_{1i} + \hat{\beta_2}^* x_{2i} + ... + \hat{\beta_k}^* x_{ki} + u_i = \frac{1}{M} \sum_{m=1}^M y_i^{*(m)} = \hat{y}_i^* \Rightarrow \hat{y} = \hat{y}^* \square$ 

#### Question 4

4a.

On this Naive model where we impute x1 and then use this values in order to create the  $x_1 * x_2$  variable. We see a good mixing with m=20 on seed=1 on the trace plots, which is a good indication that everything goes along smoothly. We are not changing anything on the predictor matrix and we are using pmm for y and x1.

Our final 95% confidence intervals for the coefficients of the independent variables for the naive case are:

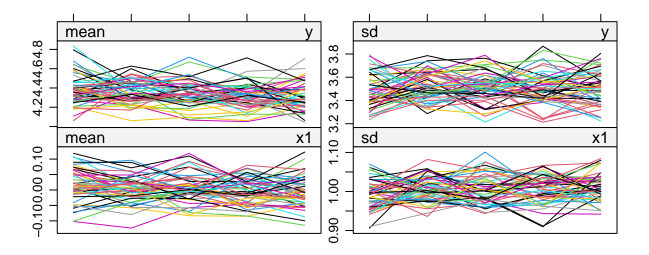
```
b_0 = 1.59, (1.40, 1.78)

b_1 = 1.41, (1.22, 1.60)

b_2 = 1.96, (1.86, 2.07)

b_3 = 0.75, (0.64, 0.87)
```

```
maxit=0)
pred <- imp0$predictorMatrix</pre>
meth <- imp0$method</pre>
imp_a = mice(dataex4,
            m=models,
            seed=set_seed,
            printFlag=FALSE,
            predictorMatrix=pred,
            method=meth)
estimates_a = summary(pool(with(imp_a, lm(y~x1+x2+(x1*x2)))), conf.int=TRUE)
estimates_a
##
           term estimate std.error statistic
                                                    df p.value
                                                                  2.5 %
## 1 (Intercept) 1.5929831 0.09541331 16.69561 154.5617 0 1.404501
             x1 1.4112333 0.09732912 14.49960 216.0125
                                                            0 1.219397
## 2
                                                           0 1.860657
             x2 1.9658191 0.05323220 36.92913 153.5344
## 3
## 4
         x1:x2 0.7550367 0.05701458 13.24287 138.0530 0 0.642302
##
       97.5 %
## 1 1.7814655
## 2 1.6030697
## 3 2.0709812
## 4 0.8677715
plot(imp_a)
```



# Iteration

#### 4b.

In order to set a passive imputation method for  $x_3 = x_1 * x_2$ . To achieve that we need to change the predicting method and the predictorMatrix accordingly. Our 95% CI for our coefficient is:

```
b_0 = 1.55, (1.38, 1.73)

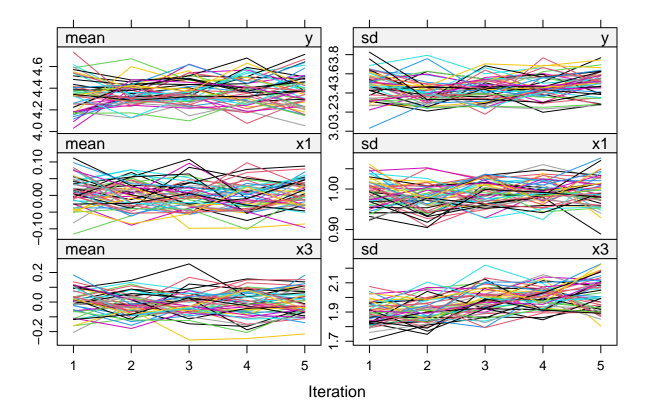
b_1 = 1.19, (1.00, 1.38)

b_2 = 2.00, (1.90, 2.09)

b_3 = 0.87, (0.76, 0.99)
```

Now our trace plots are showing evidence of good mixing for both  $x_1$  and  $x_3$ , so we are clear to move to the next analysis. This is a much better model as we don't use y as a predictor of x3 which would introduce a cyclic relation of y effecting x3 and x3 effecting y. We also prevent x3 to be used to impose x1 and x2. This results in much better  $\beta_0$ ,  $\beta_1$ ,  $\beta_2$ ,  $\beta_3$ , that are close to the values given on the problem definition.

```
pred_b <- imp1$predictorMatrix</pre>
pred_b[c('x1','x2'), 'x3'] <- 0</pre>
pred_b['x3', 'y'] <- 0</pre>
vis_seq_b <- imp1$visitSequence</pre>
vis_seq_b #x3 is already the last variable imputed so we are fine.
## [1] "y" "x1" "x2" "x3"
imp_b = mice(dataex4,
             m=models,
              seed=set_seed,
             printFlag=FALSE,
              predictorMatrix=pred_b,
             visitSequence=vis_seq_b,
             method=meth_b)
estimates_b = summary(pool(with(imp_b, lm(y~x1+x2+x3))), conf.int=TRUE)
estimates_b
##
            term estimate std.error statistic
                                                         df p.value
                                                                          2.5 %
## 1 (Intercept) 1.5534782 0.08842211 17.56889 161.1274 0 1.3788626
              x1 1.1926170 0.09584345 12.44339 180.3188 0 1.0034980 x2 1.9964402 0.04936582 40.44175 159.8398 0 1.8989468
## 2
## 3
             x3 0.8740573 0.05678521 15.39234 114.4704 0 0.7615712
## 4
##
        97.5 %
## 1 1.7280939
## 2 1.3817360
## 3 2.0939336
## 4 0.9865434
plot(imp_b)
```



#### 4c.

On this example we want to predict the composite variable x3 by pmm, and use it to also impose the missing values on x1 as well. So we make the changes and the final 95% CI is:

```
b_0 = 1.50, (1.34, 1.65)

b_1 = 1.00, (0.84, 1.16)

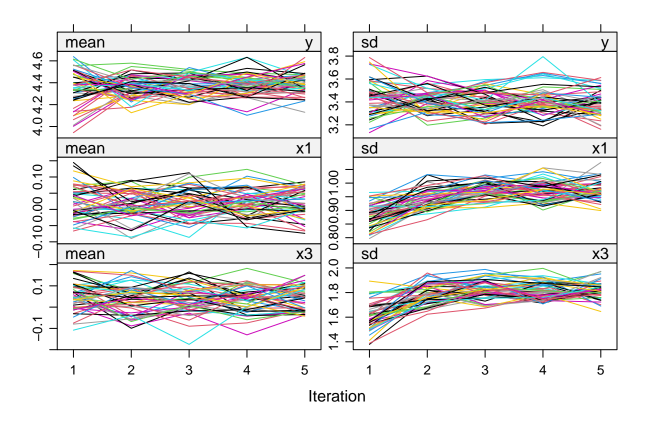
b_2 = 2.02, (1.93, 2.11)

b_3 = 1.01, (0.93, 1.10)
```

However our trace plot have a noticeably big trend on both x1 and x3 imputed values. This is not a good indication and it's going to be further explained below.

## term estimate std.error statistic df p.value 2.5 % 97.5 %

```
plot(imp_c)
```



#### 4d.

The reason we observe this behavior when we impute x3 as just another variable, and the obvious conceptual drawback is that we have information  $x_3 = x_1 * x_2$  but we disregard it. This introduced bias in our model, and our chains don't converge. So for all the cases where  $x_1$  and  $x_2$  are observed, our  $x_3$  variable follows the above relation and for all the other, it's imputed randomly using pmm. The method that makes more sense according to the definitions given and the analysis we want to do is the second approach.

# Question 5

We want to find how weight is affected by gender, height and waist circumference from the NHANES2 dataset.

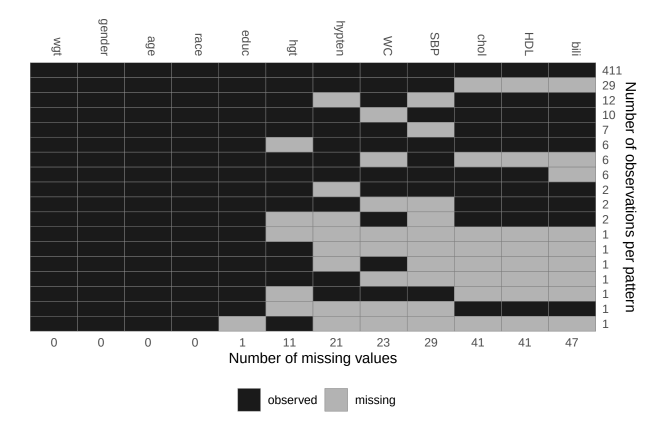
wht = 
$$\beta_0 + \beta_1$$
gender +  $\beta_2$ age +  $\beta_3$ hgt +  $\beta_4$ WC +  $\epsilon$ ,  $\epsilon \sim N(0, \sigma^2)$ 

So at first we explore the data.

# load('NHANES2.Rdata') nhanes2 <- NHANES2 summary(nhanes2)</pre>

```
gender
##
                                       bili
                                                                         chol
         wgt
                                                        age
         : 39.01
                     male :252
                                                          :20.00
                                                                    Min. : 2.07
##
                                         :0.2000
   Min.
                                  Min.
                                                   Min.
##
   1st Qu.: 65.20
                     female:248
                                  1st Qu.:0.6000
                                                   1st Qu.:31.00
                                                                    1st Qu.: 4.27
                                  Median :0.7000
                                                                    Median : 4.86
##
   Median : 76.20
                                                   Median :43.00
   Mean : 78.25
                                  Mean
                                         :0.7404
                                                   Mean
                                                          :45.02
                                                                    Mean : 5.00
   3rd Qu.: 86.41
                                  3rd Qu.:0.9000
                                                                    3rd Qu.: 5.64
##
                                                   3rd Qu.:58.00
##
   Max. :167.38
                                  Max.
                                         :2.9000
                                                   Max.
                                                           :79.00
                                                                    Max.
                                                                          :10.68
                                  NA's
                                                                    NA's
                                                                           :41
##
                                         :47
##
        HDL
                         hgt
                                                      educ
##
   Min. :0.360
                    Min. :1.397
                                    Less than 9th grade: 31
                                    9-11th grade
##
   1st Qu.:1.110
                    1st Qu.:1.626
                                                        : 69
                                    High school graduate:115
##
   Median :1.320
                    Median :1.676
                                    some college
   Mean :1.395
                    Mean
                         :1.687
                                                        :148
##
   3rd Qu.:1.590
                    3rd Qu.:1.753
                                    College or above
                                                         :136
##
   Max.
          :3.130
                    Max.
                           :1.930
                                    NA's
                                                         : 1
   NA's
           :41
                    NA's
                           :11
##
##
                                  SBP
                                               hypten
                                                               WC
                    race
   Mexican American : 52
                                                                : 61.90
##
                             Min.
                                    : 81.33
                                              no :354
                                                         Min.
##
   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
                                    :120.05
                                                         Mean
                                                                : 96.07
                             Mean
##
   other
                      : 96
                             3rd Qu.:128.67
                                                         3rd Qu.:104.80
##
                             Max.
                                    :202.00
                                                         Max.
                                                                 :154.70
##
                             NA's
                                    :29
                                                         NA's
                                                                 :23
```

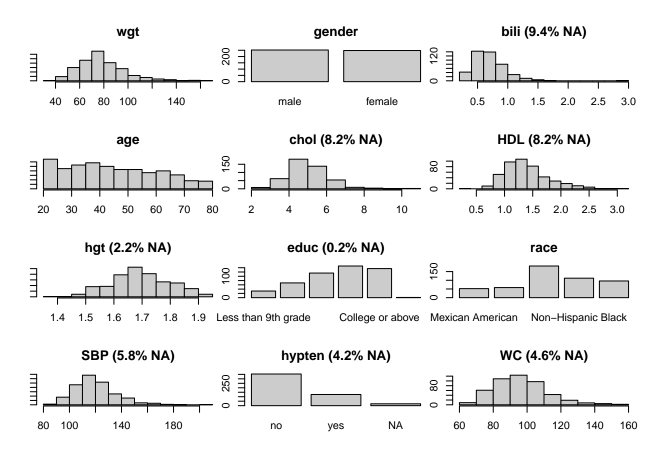
md\_pattern(nhanes2, pattern = FALSE)



We quickly realize that we have a lot of missing values on the important variables so we need to impute them.

Firstly we will try to visualize the distributions of all the variables in order to get a first feeling for our dataset. We can extract some meaning full information form the graphs, such as age looks quite Uniform(20,80), maybe with a heave decreasing tail, hgt can be approximated by Normal around 1.7 wgt and SBP look like they follow some kind of shifted Beta distribution etc.

```
par(mar=c(3,1,3,1))
plot_all(nhanes2)
```



Before starting with imposing missing values, we need to inspect what methods are automatically attributed to the missing data values. We notice that mice has chosen pmm for the height variable while from our graphs we see that we could change that for a Normal.

```
imp0 <- mice(nhanes2, maxit=0)</pre>
imp0
## Class: mids
   Number of multiple imputations:
                                           5
##
   Imputation methods:
##
                 gender
                              bili
                                                    chol
                                                                HDL
                                                                                     educ
         wgt
                                          age
                                                                           hgt
                                            11 11
           11 11
                             "pmm"
##
                                                   "pmm"
                                                              "pmm"
                                                                         "mmm"
                                                                                   "polr"
                    SBP
                                           WC
##
                           hypten
        race
                  "pmm" "logreg"
           11 11
##
                                        "pmm"
##
   PredictorMatrix:
##
            wgt
                gender bili age chol HDL hgt educ race SBP hypten WC
              0
                                  1
                                             1
                                                  1
                                                              1
                                                                           1
##
                       1
                             1
                                        1
                                                       1
                                                                   1
                                                                               1
   wgt
   gender
              1
                       0
                             1
                                  1
                                        1
                                             1
                                                  1
                                                       1
                                                              1
                                                                   1
                                                                           1
                                                                               1
##
              1
                       1
                                             1
                                                              1
                                                                           1
## bili
                             0
                                  1
                                        1
                                                  1
                                                        1
                                                                   1
                                                                               1
              1
                       1
                                  0
## age
                             1
                                        1
                                             1
                                                  1
                                                        1
                                                              1
                                                                   1
                                                                           1
                                                                               1
## chol
              1
                       1
                             1
                                  1
                                        0
                                             1
                                                  1
                                                        1
                                                              1
                                                                   1
                                                                           1
                                                                               1
## HDL
              1
                       1
                             1
                                        1
                                             0
                                                  1
                                                        1
                                                              1
                                                                   1
                                                                           1
                                                                               1
```

Which is what we do with the following chunk of code.

```
methods <- imp0$method
methods['hgt'] <- 'norm'
methods</pre>
```

```
##
                gender
                                                              HDL
                             bili
                                         age
                                                   chol
                                                                         hgt
                                                                                   educ
         wgt
##
                            "pmm"
                                                 "pmm"
                                                            "pmm"
                                                                                "polr"
                                                                      "norm"
##
                    SBP
                                          WC
                           hypten
        race
                  "pmm" "logreg"
##
                                       "mmmg"
```

Next step to our analysis, is to set the minimum and the maximum for all the numeric variables that we will try to imputing On this specific example, I tried to automate the procedure by finding the min and the max for all the variables and then setting those values to be the barriers for each variable.

```
min bili <- min(nhanes2$bili)
max bili <- max(nhanes2$bili)</pre>
min_chol <- min(nhanes2$chol)
max chol <- max(nhanes2$chol)</pre>
min_HDL <- min(nhanes2$HDL)
max HDL <- max(nhanes2$HDL)</pre>
min hgt <- min(nhanes2$hgt)
max_hgt <- max(nhanes2$hgt)</pre>
min_SBP <- min(nhanes2$SBP)
max_SBP <- max(nhanes2$SBP)</pre>
min_WC <- min(nhanes2$WC)
max_WC <- max(nhanes2$WC)</pre>
post <- imp0$post</pre>
post['bili'] <- 'imp[[j]][,i] <- sqeeze(imp[[j]][,i], c(min_bili, max_bili)'</pre>
post['chol'] <- 'imp[[j]][,i] <- sqeeze(imp[[j]][,i], c(min_chol, max_chol)'</pre>
post['HDL'] <- 'imp[[j]][,i] <- sqeeze(imp[[j]][,i], c(min_HDL , max_HDL )'</pre>
post['hgt'] <- 'imp[[j]][,i] <- sqeeze(imp[[j]][,i], c(min_hgt , max_hgt )'</pre>
post['SBP'] <- 'imp[[i]][,i] <- sqeeze(imp[[i]][,i], c(min SBP , max SBP )'</pre>
post['WC']
              <- 'imp[[j]][,i] <- sqeeze(imp[[j]][,i], c(min_WC , max_WC )'
```

Now our analysis can begin. On this part, we will try to impute the missing values by using 30 copies of the dataset and running them for 20 steps. This is a critical point in our analysis because in order to go further and still trust our result we need to somehow convene ourselves that our chains have converged. Unfortunately we know that we cannot *prove* convergence, we can only examine various plots and graphs in order to stop any hard patterns or trends. In this example we are going to examine the trace and histogram plots.

On this point I would like to spend a little time on the hyperparameters  $\mathtt{maxit}$  and  $\mathtt{m}$ , as well as the  $\mathtt{seed}$ . A good way to set these hyperparameters would be through a grid search method, where we would loop through  $\mathtt{maxit} \in (10,100)$ ,  $\mathtt{m} \in (10,100)$  and repeat the process for some random seeds in order to exclude the possibility of the random seed affecting the results. On each iteration we would store for each chain the Brooks-Gelman-Rubin statistic, which measures the proportion of between to within variability. Then after the grid search we would average on all  $\mathtt{seeds}$  the BGR statistic and finally we would pick the one where it's closest to one. This was out of the scope of the exercise but I wanted to mention it as a potentially improvement and automation of this algorithm.

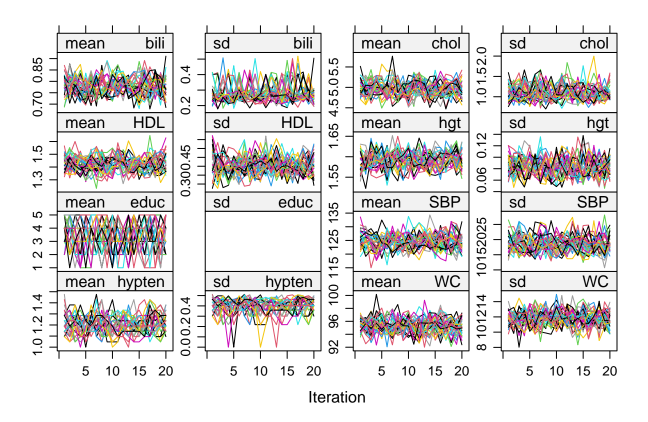
Finally we see that the loggedEvents are NULL.

## NULL

```
# TODO: grid search
```

So by looking at the chains we don't spot anything too out of the ordinary. We might be able to see some strong patterns on the educ variable, but that it to be expected since it's a categorical variable with only a few levels and only 0.2% of their entries missing. So for the moment every chain looks to be normal and well behaved.

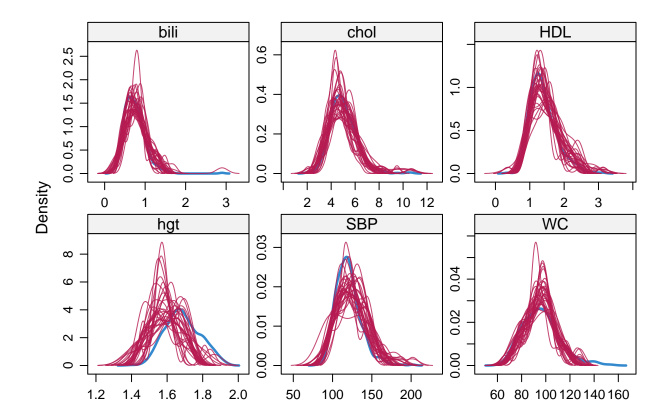
```
# checking convergence
plot(imp, layout=c(4,4))
```



However, after seeing the Histograms we start to see that something might be wrong with the hgt variable after all. We can clearly see that the imposed lines are not following very well the blue line, indicating that the imposed values don't have the same distribution as the complete cases on the original dataset. Furthermore, we see that the blue line looks like a mixture model of two Normals, or a *Binormal* distribution, which

makes sense, because the distribution of the whole population consists males and females and we can assume that they have statistically significant average heights. Our imposed data not only missed the mean of the distribution, but they also missed the structure of the distribution. hgt was the only variable that we changed the imposing method, so it's clear not that we should change it back.

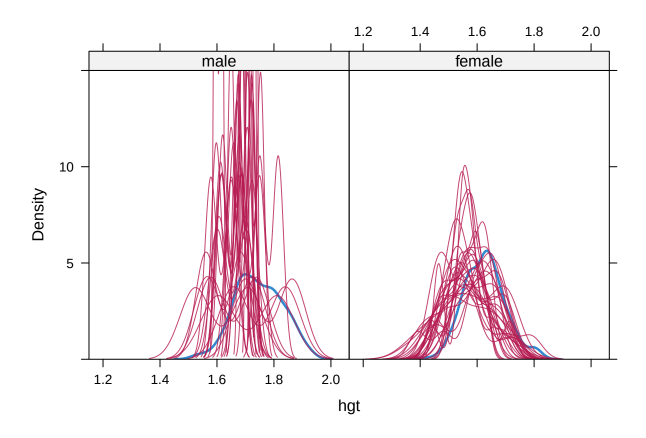
```
# maxit = 20, m=30 no strong patterns visible
densityplot(imp)
```



# height's actual distribution appears to be biNormal which makes sense since height has a big correlat # it appears as if the imputed values for height are a bit biased towards lower numbers, however they d # we should not be to worries about this.

Taking a more granular look we see the problem clearly on the males. The imposed distributions for height of the M=30 datasets when gender=male are very poorly approximating the complete cases.

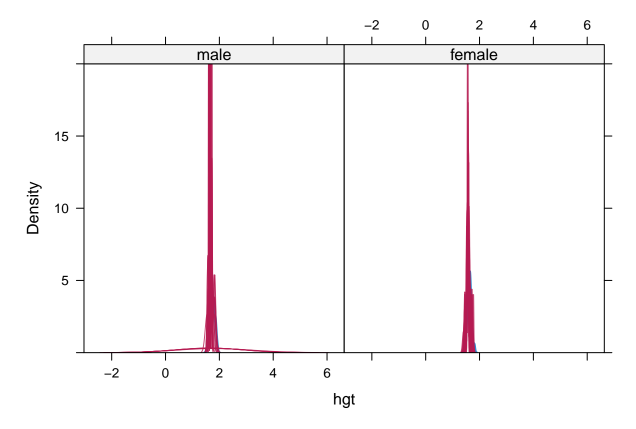
densityplot(imp, ~hgt|gender, ylim=c(0,15))



#here we can see more clearly where the problem on the above diagram comes from. It comes from the esti # it doesn't really cover very well the spectrum since it over attributes the median ~1.7...

After changing the method back to pmm from norm we see a much better coverage hense it should stay like that.

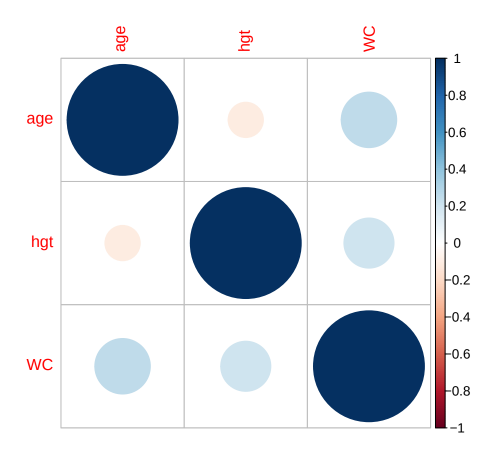
## NULL



So after changing the method back, we can start our analysis and see our model. Before that I would like to see the correlation plot for our dependent variables in order have a feeling of multicolinearity. It's very evident that the WC is correlated to age and to hgt. From the theory of OLS we know that the OLS method will give unbiased estimators in the presence of multicolinearity, but not efficient ones. Hence the  $R^2$  produced will be slightly overestimated. Whit that in mind the final model shows that Weight is mostly influenced by height since our  $\hat{\beta}_3 = 52$ , which makes total sense. All other variables coefficients have relatively little influence on the weight. An important observation is also that genders coefficient, has a p-values of 0.122 which is statistically insignificant. That result combined with the multicolinearity we observed from the correlation diagram could indicate that a simpler model, one without gender as an independent variable could be a better one.

```
comp1 <- complete(imp, 1)

# proceed to analysis
model_data <- comp1[,c(4,7,12)]
M <- cor(model_data)
corrplot(M)</pre>
```



```
fit <- with(imp, lm(wgt ~ gender + age + hgt + WC))
summary(fit$analyses[[1]])</pre>
```

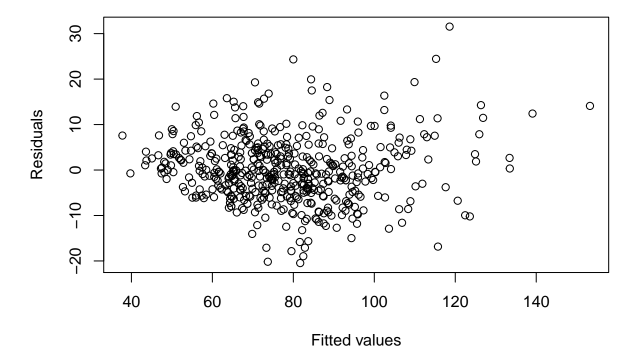
```
##
## Call:
## lm(formula = wgt ~ gender + age + hgt + WC)
##
## Residuals:
##
        Min
                  1Q
                       Median
                                    3Q
                                            Max
   -20.4638 -4.5537 -0.4955
                                3.8854
                                        31.5403
##
## Coefficients:
                  Estimate Std. Error t value Pr(>|t|)
                              7.50652 -13.390 < 2e-16 ***
## (Intercept) -100.51035
  genderfemale
                  -1.26815
                              0.81952
                                       -1.547
                                                 0.122
##
## age
                              0.02085
                                       -7.590
                  -0.15827
                                               1.6e-13 ***
## hgt
                  52.15392
                              4.29615
                                       12.140
                                               < 2e-16 ***
                              0.02213
## WC
                   1.02795
                                       46.452 < 2e-16 ***
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
## Residual standard error: 7.179 on 495 degrees of freedom
## Multiple R-squared: 0.8575, Adjusted R-squared: 0.8563
## F-statistic: 744.6 on 4 and 495 DF, p-value: < 2.2e-16
```

Finally we need to calculate our confidence intervals for our coefficients and compute the adj  $R^2$ . so we need

to make our final checks in order to trust the analysis. We need to check the second big disease a linear regression model might have, heteroskedasticity.

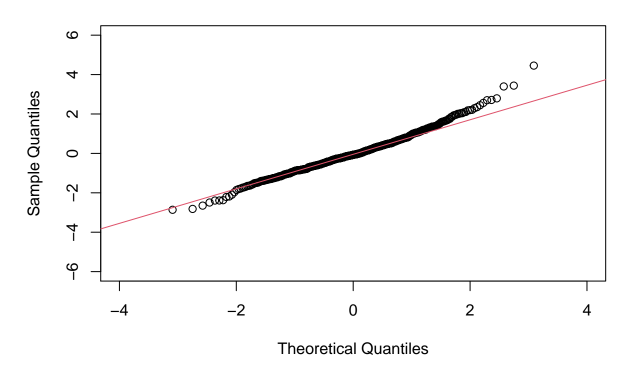
By plotting the residuals against the fitted values, we see that they are centered around 0 which is a good thing. However we also notice that the variance doesn't remain the same as it get's bigger the further right we move on the x-axis. It looks mostly like a cloud though which is a good sign so I would say that no further action is required. In order to fix this we need to model the heteroskedasticity with a model on  $\epsilon_i$  and then introduce this on our Linear model in order to cure the disease. Similar conclusions can be reached when checking the QQ-plot with the tails being a bit of which is always acceptable to that extend.

```
plot(fit$analyses[[1]]$fitted.values, residuals(fit$analyses[[1]]),
xlab = "Fitted values", ylab = "Residuals")
```



```
# QQ plot
qqnorm(rstandard(fit$analyses[[1]]), xlim = c(-4, 4), ylim = c(-6, 6))
qqline(rstandard(fit$analyses[[1]]), col = 2)
```

# Normal Q-Q Plot



```
# pooling
pooled_ests <- pool(fit)</pre>
summary(pooled_ests, conf.int = TRUE)
##
                      estimate std.error
                                                                     p.value
             term
                                            statistic
                                                             df
## 1
      (Intercept) -100.8520702 7.67289217 -13.143945 449.6454 0.000000e+00
                                            -1.659810 469.9080 9.761988e-02
## 2 genderfemale
                    -1.3850796 0.83448095
## 3
              age
                    -0.1576829 0.02141424
                                           -7.363458 451.6777 8.562040e-13
## 4
              hgt
                    52.4292200 4.39636603 11.925581 444.3719 0.000000e+00
               WC
                     1.0260613 0.02232811 45.953795 481.3369 0.000000e+00
## 5
##
            2.5 %
                       97.5 %
## 1 -115.9312510 -85.7728894
## 2
       -3.0248557
                    0.2546965
## 3
       -0.1997668
                   -0.1155990
## 4
       43.7889680
                   61.0694720
## 5
        0.9821887
                    1.0699340
pool.r.squared(pooled_ests, adjusted = TRUE)
```

We present here the final confidence intervals per variable coefficient and the final  $adj R^2 = 85.6$  which is a good indication that our model is a good fit to the data. However of course as said earlier we have to keep

hi 95 fmi

lo 95

est

## adj R^2 0.8559941 0.8304596 0.8779613 NaN

##

in mind that we have observed some multicolinearity on the model and potentially some heterosked asticity, which both have an effect on the  $adj~R^2$ 

# Merry Christmas and a happy new year

