# IDA Assignment 3

### Callum MacAskill

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Document knitted from an Rmarkdown file. All input code included inline, but if needed the original can be found here.

# Question 1

#### 1a

To get the percentage of rows with NA's, first we check which rows have NAs, then divide by the number of total rows in the data frame, before multiplying by 100 to get to percentages.

```
data = nhanes
#Row checking is done by applying the anyNA function to each row
#This will return true if a row has an NA in any column or false otherwise
#We then sum these trues and divide by the length of the dataframe
(sum(apply(data, 1, function(x) anyNA(x)))/dim(data)[1])*100
```

### ## [1] 48

It's 48% of rows that have NAs.

#### 1b

```
#First impute the data using mices built in defaults, with seed 1
impdata = mice(data, seed = 1, printFlag = FALSE)
#Then create a linear model using mice's with command
lmfit = with(impdata, lm(bmi ~ hyp+ age + chl))
#Finally pool and display resaults, again using built in mice command
poollmfit = pool(lmfit)
poollmfit
## Class: mipo
                                       ubar
                                                       b
                                                                     t dfcom
            term m
                      estimate
## 1 (Intercept) 5 19.61789252 10.588884721 0.8662133972 11.628340797
                                                                          21
## 2
             hyp 5 2.19701748 2.886391704 1.2976511612
                                                                          21
                                                          4.443573098
## 3
             age 5 -3.55287155
                                0.744810536 1.3585594461
                                                          2.375081872
                                                                          21
## 4
                                0.000287288 0.0001046083
                                                          0.000412818
                                                                          21
             chl 5 0.05378081
                              lambda
                      riv
## 1 16.936189 0.09816483 0.08938989 0.1807424
## 2 9.035494 0.53949067 0.35043452 0.4583762
## 3 3.528053 2.18884032 0.68640637 0.7824821
## 4 10.228828 0.43694808 0.30408063 0.4092932
```

Our output poollmfit is of class mipo, where the level of affect from missingness is given by the size of lambda (ref). Here we can see the highest value, and so the variable most effected by missingness in our data,

is age.

#### 1c

First rerun five times with the new seeds

```
impdata2 = mice(data, seed = 2, printFlag = FALSE)
lmfit2 = with(impdata2, lm(bmi ~ hyp+ age + chl))
poollmfit2 = pool(lmfit2)

impdata3 = mice(data, seed = 3, printFlag = FALSE)
lmfit3 = with(impdata3, lm(bmi ~ hyp+ age + chl))
poollmfit3 = pool(lmfit3)

impdata4 = mice(data, seed = 4, printFlag = FALSE)
lmfit4 = with(impdata4, lm(bmi ~ hyp+ age + chl))
poollmfit4 = pool(lmfit4)

impdata5 = mice(data, seed = 5, printFlag = FALSE)
lmfit5 = with(impdata5, lm(bmi ~ hyp+ age + chl))
poollmfit5 = pool(lmfit5)

impdata6 = mice(data, seed = 6, printFlag = FALSE)
lmfit6 = with(impdata6, lm(bmi ~ hyp+ age + chl))
poollmfit6 = pool(lmfit6)
```

We've repeated everything as in Q1b, just with new seeds. Now lets compare results.

```
#The subsetting below pulls out only the relevant columns, term and lambda poollmfit2[[3]][c(1,10)]
```

```
## term lambda

## 1 (Intercept) 0.4144454

## 2 hyp 0.1430995

## 3 age 0.4033924

## 4 chl 0.2959966

poollmfit3[[3]][c(1,10)]
```

```
## term lambda
## 1 (Intercept) 0.2772900
## 2 hyp 0.4101152
## 3 age 0.5895051
## 4 chl 0.5621346
poollmfit4[[3]][c(1,10)]
```

```
## term lambda
## 1 (Intercept) 0.1315114
## 2 hyp 0.1961083
## 3 age 0.2189333
## 4 chl 0.3305334
poollmfit5[[3]][c(1,10)]
```

```
## term lambda
## 1 (Intercept) 0.4855733
## 2 hyp 0.5942866
```

```
## 3     age 0.4511896
## 4     chl 0.2346065

poollmfit6[[3]][c(1,10)]

##     term     lambda
## 1 (Intercept) 0.4168136
## 2     hyp 0.2960364
## 3     age 0.6549523
## 4     chl 0.5196295
```

We can see that although age has the highest lambda most often (in seeds 2,3 and 6, not counting the intercept), that in seed 3 cholesterol is most affected by missingness, as is hypertension in seed 5.

#### 1d

Now we repeat the analysis again with a higher M (m = 100)

```
#The previous 4 lines of code for each seed can be condensed down to 1
pool(with(mice(data, seed = 2, printFlag = FALSE, m = 100), lm(bmi ~ hyp+ age + chl)))[[3]][c(1,10)]
##
                    lambda
            term
## 1 (Intercept) 0.1882474
             hyp 0.2825108
             age 0.4031077
## 3
## 4
             chl 0.2939693
pool(with(mice(data, seed = 3, printFlag = FALSE, m = 100), lm(bmi ~ hyp+ age + chl)))[[3]][c(1,10)]
                    lambda
##
            term
## 1 (Intercept) 0.2199607
## 2
             hyp 0.2425105
## 3
             age 0.3093072
## 4
             chl 0.3281911
pool(with(mice(data, seed = 4, printFlag = FALSE, m = 100), lm(bmi ~ hyp+ age + chl)))[[3]][c(1,10)]
            term
                    lambda
## 1 (Intercept) 0.2144722
## 2
             hyp 0.2565132
## 3
             age 0.3943223
             chl 0.2835232
pool(with(mice(data, seed = 5, printFlag = FALSE, m = 100), lm(bmi ~ hyp+ age + chl)))[[3]][c(1,10)]
            term
                    lambda
## 1 (Intercept) 0.2294356
## 2
             hyp 0.2893046
## 3
             age 0.3322570
             chl 0.2461956
pool(with(mice(data, seed = 6, printFlag = FALSE, m = 100), lm(bmi ~ hyp+ age + chl)))[[3]][c(1,10)]
            term
                    lambda
## 1 (Intercept) 0.2472607
## 2
             hyp 0.2860700
## 3
             age 0.4430300
## 4
             chl 0.3113085
```

This would seem to be a good idea. A higher m, meaning more database copies created and imputed before being pooled back together, would reduce the effect of random chance through a larger number of receptions. Given that in q1c we can see the lambda for age has a range of (0.219, 0.654), a difference of over 300%, this would definitely seem to be an area we can improve on. Now, when repeating for m = 100, we can see our lambda for age now has a range of (0.309, 0.443), much more consistent results.

The pattern does not just apply to lambda for age. See below the summary of resulst for default m and m = 100, with seed 2.

```
#Recalculate and name the m=100 dataframe for seed 2
m100 = pool(with(mice(data, seed = 2, printFlag = FALSE, m = 100), lm(bmi ~ hyp+ age + chl)))
#First display with default m
summary(poollmfit2)
##
                   estimate std.error statistic
                                                        df
                                                               p.value
## 1 (Intercept) 19.9464142 4.36349993
                                       4.571196
                                                 7.595481 0.002083932
             hyp 1.5304762 2.01529855
                                       0.759429 15.210844 0.459204412
## 3
             age -4.0615093 1.27092690 -3.195706 7.827560 0.013061972
## 4
             chl 0.0628349 0.02215369
                                        2.836317 10.450088 0.016956575
#Then after display with the increased m
summary(m100)
##
                   estimate
                             std.error
                                        statistic
                                                        df
                                                                p.value
## 1 (Intercept) 20.4677977 3.63913732
                                        5.6243543 15.53932 4.241712e-05
## 2
             hyp 1.7199145 2.14123878
                                        0.8032334 13.65957 4.355947e-01
## 3
             age -3.6337887 1.26258958 -2.8780443 11.27749 1.468267e-02
## 4
                 0.0535212 0.02123996
                                       2.5198358 13.43174 2.511358e-02
```

We can see that the standard error is down for all values except cholesterol (where it has seen a slight increase). This would suggest we get a more a accurate estimation from m = 100, however in a real life setting more tests would need to be carried out (change of seeds, m, ect).

# Question 2

If we're investigating the effects of acknowledging or not parameter uncertainty on confidence intervals then the first step is to impute the missing values, using our given methods (stochastic regression imputation and bootstrap variant), calculate and pool our results. Here we are only interested in the results for  $\beta_1$ .

The second step would then be to check if the correct value of our parameter is included in the confidence interval, the correct value being  $\beta_1 = 3$ . The number of times the correct value (out of 100 possible) was then counted, and is displayed below.

```
#First load the data
load('dataex2.Rdata')
#And create a 100x2 matrix to store our boolean interval values in
conf_mat = matrix(OL, nrow = 100, ncol = 2)
#Ttart a for loop to calculate the intervals for all datasets
for (i in 1:100) {
  #Multiple imputation steps 1-4 condensed into one line, along with taking the summary of the informat
  sto = as.matrix(summary(pool(with(mice(dataex2[,,i], m = 20, seed = 1, printFlag = FALSE, method = "n
  boot = as.matrix(summary(pool(with(mice(dataex2[,,i], m = 20, seed = 1, printFlag = FALSE, method = ":
  #We then check if our value of beta is between the intervals, and record as true or false
  #Note as it's being stored in a matrix, true or false becomes 1 or 0
  conf_mat[i,1] = (sto[1] <= 3 & sto[2] >= 3)
  conf_mat[i,2] = (boot[1] \le 3 \& boot[2] > = 3)
}
#Sum each column to give the required totals
list(sum(conf_mat[,1]), sum(conf_mat[,2]))
## [[1]]
## [1] 88
##
## [[2]]
## [1] 95
```

In the list above, entry one is the number of times our confidence interval included the correct value using the stochastics regression imputation, 88 times. The second for bootstrapping, 95 times.

Given we were calculating the 95% confidence interval we would expect the true value of  $\beta_1$  to be included in the interval 95% of the time (at least over a large enough sample), as we can see with the bootstrapping method.

We also see the expected results of using improper multiple imputation, namely less accurate confidence intervals. This is expected as a consequence of using stochastic regression imputation, in which after imputing each missing observation we treat it the same as fully observed variable. Although both methods may converge to the same expected value, by treating imputed variables as observed we fail to fully account for the variation in the missing variables, something the bootstrapping method does. SRI on the other hand displays narrower confidence, and ultimately less accurate, confidence intervals, hence improper multiple imputation should be avoided.

# Question 3

What do we mean by compute predicted values then pool? Assume we are using a linear model with n terms, with m imputations, and that  $\beta_i^{(m)}$  is the i th beta term in the m th imputed database. Then our linear model looks something like this

$$y = \sum_{i=0}^{N} \beta_i x_i$$
$$= \beta_0 + \sum_{i=1}^{N} \beta_i x_i$$

/newpage

# Question 4

#### 4a

First we impute the missing values of y,  $x_1$ , then calculate our coefficients of

$$y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_1 x_2 + \epsilon$$

Code below.

```
load('dataex4.Rdata')
#Again steps 1-4 + subsetting condensed to one line, fast to code but possibly harder to read
summary(pool(with(mice(dataex4, seed = 1, m =50, printFlag = F), lm (y ~ x1 + x2 + x1*x2))), conf.int =
### term estimate 2.5 % 97.5 %
### d (Tutaurumb) 4 5000001 4 400501 1 7011055
```

We can see our estimates from the *impute*, then transform method above. Note the correct values for  $\beta_{1,2,3,4}$  are 1.5, 1, 2 and 1 respectively. This method does not seem to have worked well however, as the correct values for  $\beta_{2,4}$  are not included in the confidence intervals for these values.

### **4**b

Let's try and improve on this result. Next we will calculate  $x_1x_2$  and append this value as a new column to out dataframe, and see if this helps our model. Note as  $x_1$  had missing values,  $x_1x_2$  will also missing values. By predicting the value of  $\beta_4$  without imputing values for  $x_1x_2$  ourselves, we are doing passive imputation.

```
load('dataex4.Rdata')
#Create a new column, x1*x2
dataex4 = mutate(dataex4, x1x2 = x1*x2)
#Calculate the new predictor matrix with our new column, set maxit to 1 as we only need the predictor m
pred = mice(dataex4, seed = 1, m =50, printFlag = F, maxit = 1)$pred
#Set the values of the predictor matrix to 0 for this new column, this means the new value will not be
pred[,4]=0
pred[4,]=0
#Then complete the usual analysis
summary(pool(with(mice(mutate(dataex4, x1x1 = x1*x2), seed = 1, m =50, printFlag = F, predictorMatrix =
```

```
## term estimate 2.5 % 97.5 %

## 1 (Intercept) 1.5222144 1.3478381 1.6965906

## 2 x1 1.2438490 1.0615724 1.4261256

## 3 x2 2.0024539 1.9059183 2.0989894

## 4 x1x2 0.8656623 0.7687889 0.9625358
```

x1x2 1.017793 0.9303479 1.105238

Unfortunately the correct values for  $\beta_{2,4}$  are still not included in the confidence intervals for their values, although we have seen an improvement in the accuracy of both over the *impute*, then transform method.

#### 4c

## 4

Finally, lets try and impute the values for  $x_1x_2$  also, and see if our results improve any further.

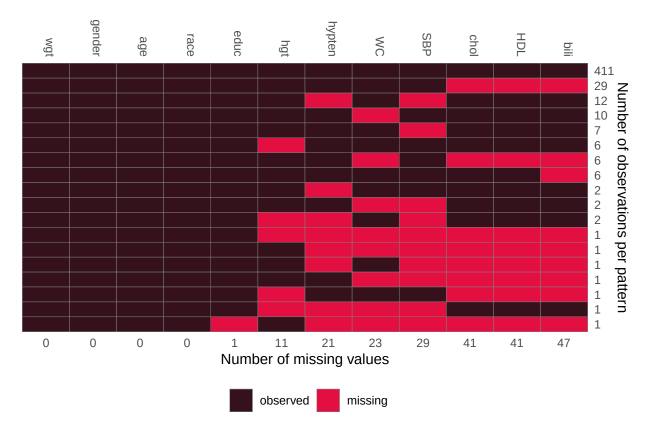
Now we have our strongest model. All values for the  $\beta$ 's are in their confidence intervals, and the point estimates are within 3% of the true value for each. Why does treating  $x_1x_2$  as just another variable improve our results so much?

In each of our three attempts at this model, we have supplied more information to the lm function at each step, the function which attempts to fit a given model to the data. By giving lm this additional information (not just more values to create a model from, but values with a correct variance and means) we inevitably allow it to create a better model.

# Question 5

Before imputing our data and creating our model we should make some common-sense checks to our imputation. First lets take a look at what's missing from the data

```
load('NHANES2.Rdata')
data2 = NHANES2
#Use the JointAI package missing data visualisation function
md_pattern(data2, pattern = FALSE, color =c('#34111b','#e30f41'))
```



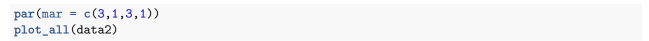
Note that out of 500 observations, 411 are fully observed, and in particular only 32 are missing information required for the model. It's fair to say then that we are missing a relatively small amount of data, and therefor do not have to set our value. For the purposes of this question m = 25 should be reasonable.

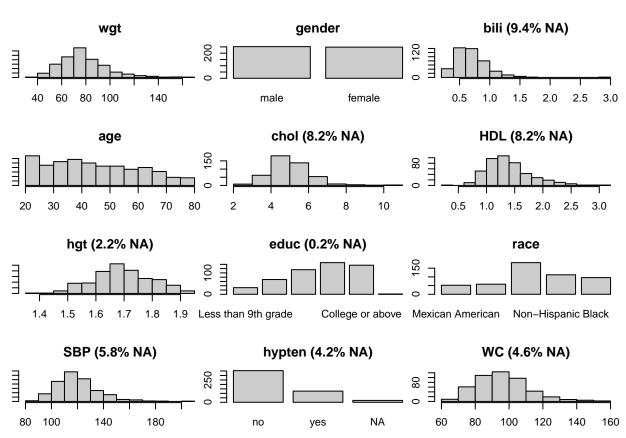
We should also take a look at how mice will impute this data, to ensure it is using sensible values

```
#Dry run mice with 0 it to see which methods it will use
M = mice(data2, maxit = 0)
M$method
##
                                                           HDL
                                                                               educ
         wgt
               gender
                            bili
                                       age
                                                chol
                                                                      hgt
                                        11 11
          11 11
##
                           "pmm"
                                               "pmm"
                                                         "pmm"
                                                                    "pmm"
                                                                             "polr"
##
                   SBP
                         hypten
                                        WC
       race
                 "pmm" "logreg"
##
                                     "pmm"
```

We can see mice will mainly use predictive mean matching to impute values any numerical values which need imputing, ensuring values in the same range as the observations, meaning no ranges need to be set.

We could change the predictive models used if we wished, for instance height is normally distributed in the population, and indeed a visualisation of the data backs this up.





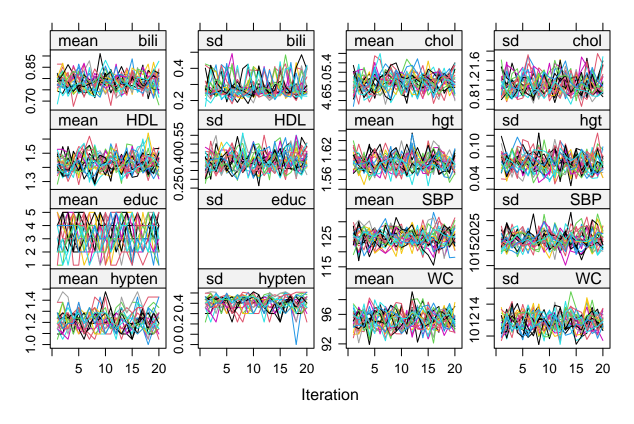
In fact we could argue that weight, cholesterol, HDL, height, SBP and waist circumference are all normally distributed, in which case we could switch our model from pmm to norm. However, whilst these values may be normally distributed in the population as whole, we should note that we do not know that these observations come from a random sample of the population. As discussed above, using norm instead of pmm could also lead to unlikely or impossible imputations, and over a large enough sample pmm should have enough observations to be able to give a full coverage of possible values. It could also be pointed out that several of the continous variables do seem to have a slight skew, again this would favour pmm for imputation.

Whilst it is not something we can check for here, we should also note that one of the main requirements of mice is that our missing data is MAR. Here we do not know the missingness mechanism, we would need to discuss it with the data collector, or some other relevant expert. If this data were to be MNAR it would potentially undermine our model and any other conclusions based on our imputations.

Let's assume however that our data is MAR, now we can start imputation and checking for issues.

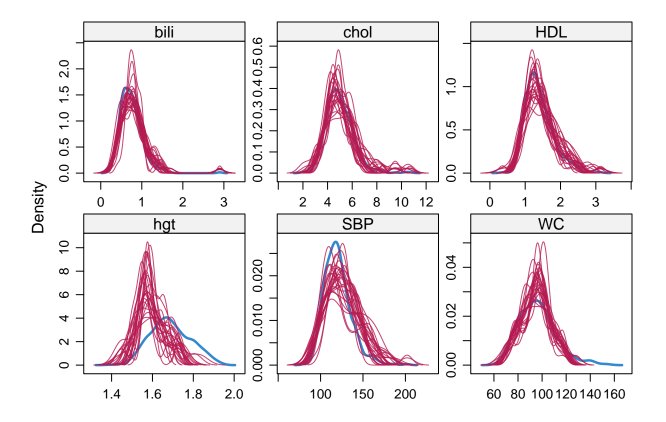
```
#Impute the data
impdata = mice(data2, m = 25, maxit = 20, seed = 1, printFlag = F)
#Check for issues, should be null
impdata$loggedEvents
## NULL
```

```
#Plot convergence
plot(impdata, layout= c(4,4))
```



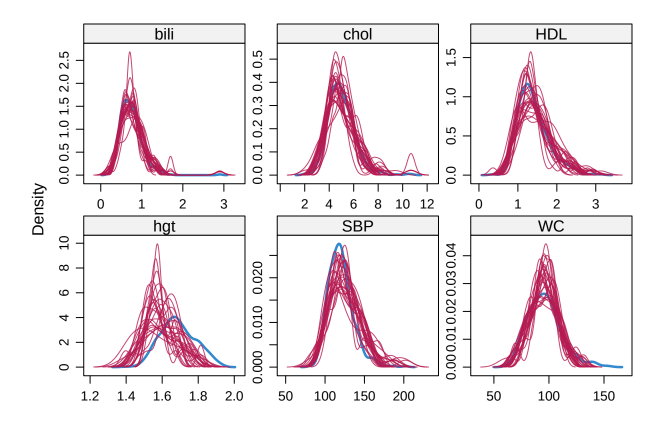
We have no logged events, so mice has had no issues when running the code, and see no obvious problems with convergance. We can also check density plots to double check the distribution of the imputed variables agrres with the observed ones.

densityplot(impdata)



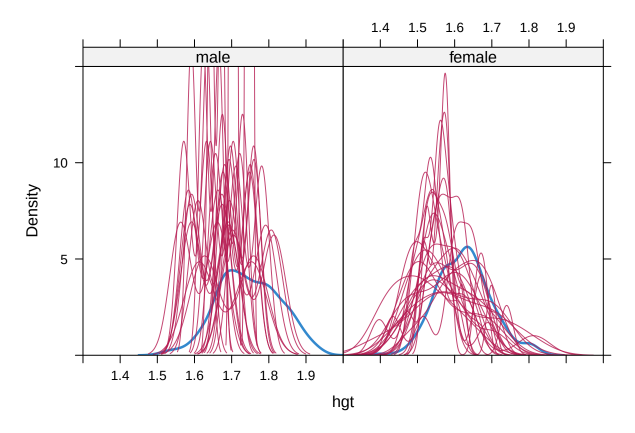
Here we do see an issue, the imputed values of height are well clearly skewed from the observed values. Given these results we should go back to our predictive model and consider using norm instead of pmm. Despite my initial hesitancy our results clearly show that pmm isn't working.

```
#Load the predictive matrix
meth = impdata$meth
#Edit predictor for height
meth['hgt'] = 'norm'
#Also changing for waist circumference to help accuracy
meth['WC'] = 'norm'
#Now rerun imputation and plot again
impdata2 = mice(data2, m = 25, maxit = 20, seed = 1, printFlag = F, method = meth)
densityplot(impdata2)
```



Hmm, this model is still showing issues. It should be noted we were only missing 11 values for height, perhaps they were outliers (or even missing MNAR), we should look a bit closer at what is missing. Perhaps we're missing a correlation, let's look at height and gender.

densityplot(impdata2, ~hgt|gender, xlim = c(1.3, 2), ylim =c(0,15))



Here, when accounting for gender, we see much more consistent density plots, suggesting our model robust enough to proceed. We will now move on to creating the and finishing and examining the model.

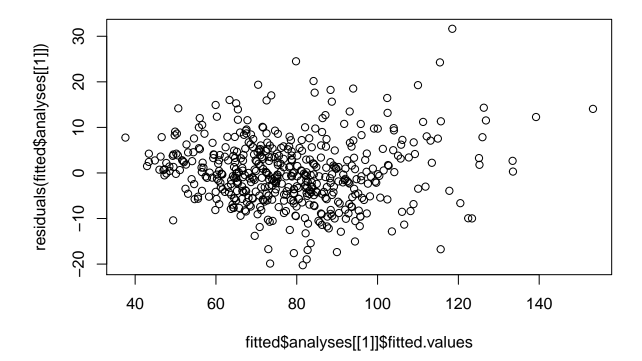
```
fitted = with(impdata2, lm (wgt ~ gender + age + hgt + WC))
summary(fitted$analyses[[1]])
```

```
##
## Call:
##
  lm(formula = wgt ~ gender + age + hgt + WC)
##
## Residuals:
##
       Min
                1Q Median
                                3Q
                                       Max
  -20.252 -4.585
                   -0.341
                             3.955
                                    31.642
##
##
## Coefficients:
                  Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                -101.94901
                              7.45924 -13.667
                                               < 2e-16 ***
## genderfemale
                  -1.36071
                              0.82303
                                       -1.653
                                                 0.0989
## age
                  -0.15216
                              0.02085
                                       -7.297 1.18e-12 ***
## hgt
                  52.69178
                              4.25515
                                       12.383
                                                < 2e-16 ***
## WC
                   1.02972
                              0.02215
                                       46.484
                                                < 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
```

Our model.

Now we'll check it through plotting.

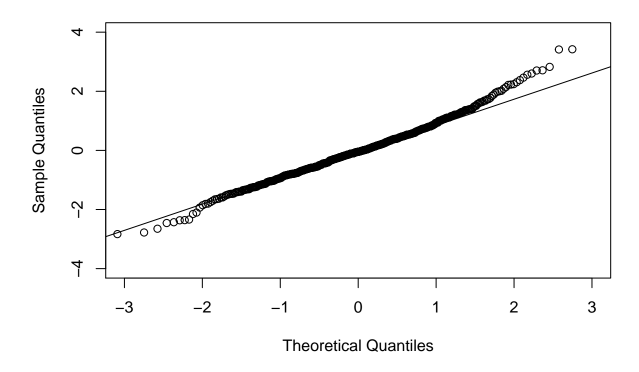
```
plot(fitted$analyses[[1]]$fitted.values, residuals(fitted$analyses[[1]]))
```



In our first plot (fitted vs residuals) we can see (whilst we start around 0) an increasing variance as the we go rightward along the graph. This would indicate a certain amount of heteroskedacity. This isn't ideal, however by also checking the qq plot we that the model holds for the most part at least.

```
qqnorm(rstandard(fitted$analyses[[1]]), xlim = c(-3, 3), ylim = c(-4, 4))
qqline(rstandard(fitted$analyses[[1]], col = 2))
```

# Normal Q-Q Plot



Here our tails only deviate towards the ends, backing up the model, even if it isn't perfect.

Now finally let's give our coefficients

```
summary(pool(fitted), conf.int = T)[c(1,2,7,8)]
```

##		term	estimate	2.5 %	97.5 %
##	1	(Intercept)	-100.8813248	-115.9072584	-85.8553913
##	2	genderfemale	-1.3318448	-2.9717613	0.3080716
##	3	age	-0.1567008	-0.1985983	-0.1148032
##	4	hgt	52.4760861	43.8822630	61.0699091
##	5	WC	1.0247435	0.9807554	1.0687317

Here we see the results of our model after pooling, and that is that.

Have a good Christmas and thank you for an interesting course!