STAT 331 Final Project

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1 Summary

2 Descriptive Statistics

First, take a look at summary statistics of the fhsd dataset.

Table 1: Summary Statistics

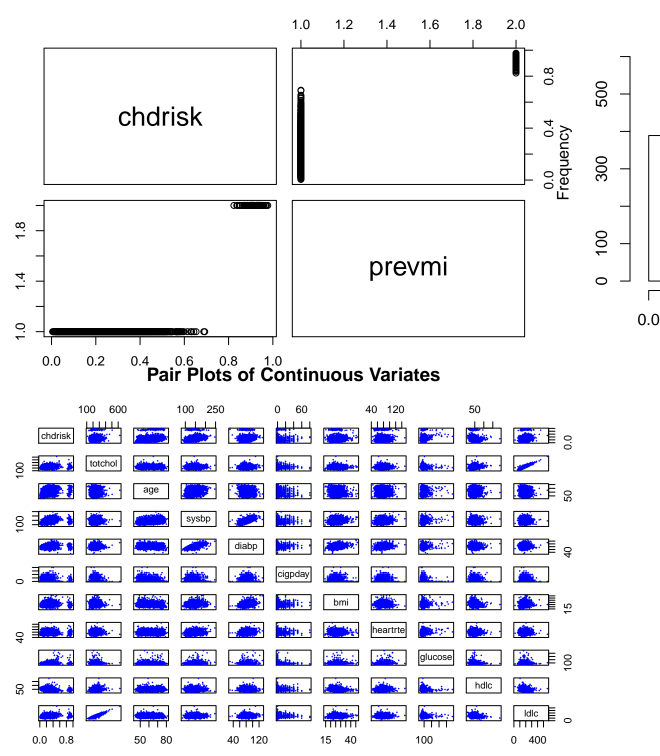
chdrisk	sex	totchol	age	sysbp	diabp	cursmoke	cigpday	bmi	diabetes	bpmeds	heartrte	glucose	prevmi	prevstrk	prevhyp	hdle	ldle
Min. :0.0050	Female:1305										Min.: 44.00						Min. : 20.0
1st Qu.:0.1320	Male :1001	1st Qu.:207.0	1st Qu.:53.00	1st Qu.:122.5	1st Qu.: 73.00	Yes: 802	1st Qu.: 0.00	1st Qu.:23.22	Yes: 164	Yes: 333	1st Qu.: 70.00	1st Qu.: 75.00	Yes: 117	Yes: 46	Yes:1349	1st Qu.: 38.00	1st Qu.:152.0
Median :0.2240	NA	Median :235.5	Median :60.00	Median :136.0	Median : 80.00	NA	Median: 0.00	Median :25.40	NA	NA	Median : 76.00	Median : 83.00	NA	NA	NA	Median : 47.00	Median :180.0
Mean :0.2655	NA	Mean :237.8	Mean :60.23	Mean :139.2	Mean: 81.07	NA	Mean : 6.84	Mean :25.78	NA	NA	Mean: 77.61	Mean: 89.07	NA	NA	NA	Mean: 48.89	Mean :183.1
3rd Qu.:0.3448	NA	3rd Qu.:265.0	3rd Qu.:67.00	3rd Qu.:153.0	3rd Qu.: 88.00	NA	3rd Qu.:10.00	3rd Qu.:27.91	NA	NA	3rd Qu.: 85.00	3rd Qu.: 95.00	NA	NA	NA	3rd Qu.: 57.00	3rd Qu.:210.0
Max. :0.9770	NA	Max. :625.0	Max. :81.00	Max. :246.0	Max. :130.00	NA	Max. :80.00	Max. :46.52	NA	NA	Max. :150.00	Max. :478.00	NA	NA	NA	Max. :189.00	Max. :565.0

Then take a look at chdrisk grouped by sex as well as chdrisk grouped by cursmoke.

```
## fhsd$sex: Female
##
     Min. 1st Qu. Median
                          Mean 3rd Qu.
    0.005
          0.104 0.179
                          0.215 0.285
                                        0.949
  ______
## fhsd$sex: Male
     Min. 1st Qu. Median
                          Mean 3rd Qu.
   0.0210 0.1860 0.2860 0.3314 0.4060 0.9770
## fhsd$cursmoke: No
     Min. 1st Qu. Median
                         Mean 3rd Qu.
   0.0050 0.1390 0.2350 0.2754 0.3580 0.9770
## fhsd$cursmoke: Yes
##
     Min. 1st Qu. Median
                          Mean 3rd Qu.
   0.0080 0.1220 0.1995 0.2471 0.3140 0.9710
## fhsd$prevstrk: No
##
     Min. 1st Qu. Median
                          Mean 3rd Qu.
   0.0050 0.1300 0.2200 0.2611 0.3392 0.9770
## fhsd$prevstrk: Yes
     Min. 1st Qu. Median
##
                          Mean 3rd Qu.
   0.2020 0.3412 0.4410 0.4820 0.5060 0.9660
## fhsd$prevhyp: No
##
     Min. 1st Qu. Median
                          Mean 3rd Qu.
                                         Max.
          0.077
                   0.140
                          0.176 0.216
                                        0.944
## fhsd$prevhyp: Yes
##
     Min. 1st Qu. Median
                          Mean 3rd Qu.
   0.0320 0.1980 0.2890 0.3291 0.4010 0.9770
## fhsd$prevmi: No
##
     Min. 1st Qu. Median
                          Mean 3rd Qu.
                                         Max.
   0.0050 0.1270 0.2120 0.2309 0.3220
```

[ADD SOME COMMENTS HERE REGARDING SUMMARY]

Now take a look at pair plots of all numeric explanatory variates i.e. variates excluding response variate chdrisk and logical variates such as cursmoke.



From the pair plots, we can observe a strong correlation between low density lipoprotein cholesterol and serum total cholestrol. This correlation could be explained by the fact that there could be a relationship between the amount [TO BE CONTINUED]

Now take a look at the VIFs of these variates.

##	sexMale	totchol	age	sysbp	diabp	cursmokeYes
##	1.225191	10.634882	1.489926	2.918660	2.406411	2.978609
##	cigpday	bmi	${\tt diabetesYes}$	bpmedsYes	heartrte	glucose
##	2.973594	1.181865	1.286401	1.214744	1.105902	1.308923
##	prevmiYes	${\tt prevstrkYes}$	prevhypYes	hdlc	ldlc	
##	1.067134	1.045746	1.823014	2.287571	10.367649	

[ADD COMMENTS]

3 Candidate Models

3.1 Automated Model Selection

```
library(gtools)
load_calcs = TRUE
# model with only intercept
MO <- lm(I(logit(chdrisk)) ~ 1, data = fhsd)</pre>
Mmax <- lm(I(logit(chdrisk)) ~ (.)^2, data = fhsd)</pre>
# starting model for stepwise selection
Mstart <- lm(I(logit(chdrisk)) ~ ., data = fhsd)</pre>
# find model coefficients which are NA
beta.max <- coef(Mmax)</pre>
names(beta.max)[is.na(beta.max)]
## [1] "cursmokeYes:cigpday" "bpmedsYes:prevhypYes"
# find the problem with the NA coeffs
kable(table(fhsd[c("cursmoke", "cigpday")]), "latex")
                      3
                           4
                               5
                                    6
                                            8
                                               9
                                                   10
                                                       12
                                                            14
                                                                15
                                                                     16
                                                                         17
                                                                              18
                                                                                  19
                                                                                        20
                                                                                            23
 No
      1504
                                                                      0
                           0
                               0
                                    0
                                       0
                                            0
                                               0
                                                                                         0
 Yes
         0
             16
                 18
                     34
                          11
                              18
                                   24
                                       9 | 18
                                               5
                                                   76
                                                        3
                                                             3
                                                                 50
                                                                      6
                                                                          1
                                                                               8
                                                                                    1
                                                                                       279
                                                                                             1
kable(table(fhsd[c("bpmeds", "prevhyp")]), "latex")
       No
             Yes
 No
      957
            1016
 Yes
            333
        0
# remove the coeffs with the problem and add quadratic terms for the continuous variables
Mmax <- lm(I(logit(chdrisk)) ~ (.)^2 - cursmoke:cigpday - bpmeds:prevhyp +</pre>
             I(totchol ^ 2) + I(sysbp ^ 2) + I(diabp ^ 2)
           + I(bmi ^ 2) + I(glucose ^ 2)
           + I(hdlc ^ 2) + I(ldlc ^ 2), data = fhsd)
anyNA(coef(Mmax)) # check if there are any remaining NAs
## [1] FALSE
if(!load_calcs){
  #forward model selection
  system.time({
    Mfwd <- step(object = MO,
                  scope = list(lower = MO, upper = Mmax),
                  direction = "forward", trace = FALSE)
 })
  #backward model selection
  system.time({
    Mback <- step(object = Mmax,</pre>
                  scope = list(lower = MO, upper = Mmax),
                  direction = "backward", trace = FALSE)
 })
```

25

14

26

1

27

0

1

```
#stepwise model selection
  system.time({
   Mstep <- step(object = Mstart,</pre>
                  scope = list(lower = MO, upper = Mmax),
                  direction = "both", trace = FALSE)
 })
}
# the caching/loading block
if(!load_calcs) {
  saveRDS(list(Mfwd = Mfwd, Mback = Mback, Mstep = Mstep), file = "models_automated.rds")
} else {
  # just load the calculations
  tmp <- readRDS("models_automated.rds")</pre>
  Mfwd <- tmp$Mfwd
 Mback <- tmp$Mback
 Mstep <- tmp$Mstep</pre>
 rm(tmp) # optionally remove tmp from workspace
# Stepwise model selection
Mstep$call
## lm(formula = I(logit(chdrisk)) ~ sex + totchol + age + sysbp +
       diabp + cursmoke + cigpday + bmi + diabetes + bpmeds + heartrte +
##
       glucose + prevmi + prevstrk + prevhyp + hdlc + ldlc + I(hdlc^2) +
##
       I(bmi^2) + I(diabp^2) + I(sysbp^2) + sysbp:prevmi + totchol:prevhyp +
##
       diabetes:prevmi + prevhyp:ldlc + sysbp:prevhyp + totchol:heartrte +
##
       sysbp:diabetes + diabp:bmi + diabp:hdlc + prevmi:hdlc + prevmi:prevhyp +
       sex:glucose + age:ldlc + age:heartrte + cigpday:hdlc + bmi:ldlc +
##
##
       totchol:hdlc + totchol:prevmi + sysbp:heartrte + sysbp:bpmeds +
##
       cursmoke:hdlc + prevmi:prevstrk + diabetes:hdlc + sex:sysbp +
##
       cigpday:glucose + heartrte:glucose + diabp:glucose + cursmoke:ldlc +
##
       age:cigpday + age:hdlc + hdlc:ldlc + age:prevhyp + diabp:prevhyp +
       diabp:cursmoke + diabp:cigpday + bmi:bpmeds + bpmeds:glucose +
##
##
       age:prevmi + sex:ldlc + cigpday:heartrte + cigpday:prevmi +
       glucose:prevmi + heartrte:prevmi + bpmeds:prevstrk, data = fhsd)
##
# Forward model selection
Mfwd$call
## lm(formula = I(logit(chdrisk)) ~ prevmi + sysbp + sex + age +
       ldlc + prevhyp + diabetes + hdlc + I(hdlc^2) + cigpday +
##
##
       I(bmi^2) + bmi + totchol + I(glucose^2) + I(sysbp^2) + bpmeds +
##
       heartrte + cursmoke + prevstrk + prevmi:sysbp + sysbp:age +
       prevhyp:hdlc + prevmi:diabetes + sysbp:prevhyp + prevhyp:totchol +
##
       sysbp:diabetes + prevmi:hdlc + prevmi:prevhyp + age:ldlc +
##
       age:cigpday + hdlc:cigpday + prevhyp:bmi + ldlc:bmi + prevmi:totchol +
##
##
       ldlc:prevhyp + sysbp:bpmeds + sysbp:hdlc + hdlc:totchol +
##
       totchol:heartrte + age:heartrte + diabetes:hdlc + sysbp:heartrte +
       bmi:bpmeds + sysbp:sex + ldlc:hdlc + prevmi:bmi + age:bmi +
##
```

```
prevmi:age + sysbp:cursmoke + hdlc:cursmoke + ldlc:cursmoke +
       prevmi:cigpday + sex:diabetes + prevmi:prevstrk, data = fhsd)
# Backward model selection
Mback$call
## lm(formula = I(logit(chdrisk)) ~ sex + totchol + age + sysbp +
##
       diabp + cursmoke + cigpday + bmi + diabetes + bpmeds + heartrte +
       glucose + prevmi + prevstrk + prevhyp + hdlc + ldlc + I(totchol^2) +
##
##
       I(sysbp^2) + I(diabp^2) + I(bmi^2) + I(hdlc^2) + I(ldlc^2) +
##
       sex:totchol + sex:sysbp + sex:glucose + sex:prevstrk + sex:prevhyp +
##
       totchol:age + totchol:bpmeds + totchol:heartrte + totchol:prevmi +
       totchol:prevstrk + totchol:prevhyp + totchol:hdlc + totchol:ldlc +
##
       age:cursmoke + age:bmi + age:heartrte + age:prevmi + age:prevhyp +
##
##
       age:hdlc + sysbp:diabetes + sysbp:bpmeds + sysbp:heartrte +
       sysbp:prevmi + sysbp:prevhyp + diabp:cursmoke + diabp:cigpday +
##
##
       diabp:bmi + diabp:glucose + diabp:prevhyp + diabp:hdlc +
##
       cursmoke:bmi + cursmoke:hdlc + cursmoke:ldlc + cigpday:bmi +
       cigpday:heartrte + cigpday:glucose + cigpday:prevmi + cigpday:hdlc +
##
       bmi:prevmi + bmi:prevhyp + bmi:ldlc + diabetes:prevmi + diabetes:hdlc +
##
##
       bpmeds:glucose + bpmeds:prevstrk + bpmeds:ldlc + heartrte:glucose +
##
       heartrte:prevmi + glucose:prevmi + prevmi:prevhyp + prevmi:hdlc +
##
       prevhyp:ldlc, data = fhsd)
beta.fwd = coef(Mfwd)
beta.back = coef(Mback)
beta.step = coef(Mstep)
identical(names(beta.fwd) [names(beta.fwd) %in% names(beta.back)], names(beta.fwd))
## [1] FALSE
identical(names(beta.fwd) [names(beta.fwd) %in% names(beta.step)], names(beta.fwd))
## [1] FALSE
identical(names(beta.back) [names(beta.back) %in% names(beta.step)], names(beta.back))
## [1] FALSE
3.2
     Manual Model Selection
library(stringr) # For string operations
## Warning: package 'stringr' was built under R version 3.5.2
table <- c() # Initialize empty vector
names.table <- names(beta.step)</pre>
                                                   # Obtain variate names in stepwise model
names.table <- str_remove_all(names.table, "Yes") # Remove "Yes" from interactions</pre>
names.table <- str_remove_all(names.table, "Male") # Remove "Male"</pre>
 # Perform F-tests by removing one variate at a time
 for(i in names.table){
    # Obtain model without variate i
   mdl <- lm(as.formula(paste0("update(Mstep, . ~ . -", i,")")),data = fhsd)</pre>
  test <- anova(Mstep,mdl)</pre>
                                           \# F-Test between Stepwise and reduced model
```

```
table <- cbind(table,test$`Pr(>F)`[2]) # Add corresponding p-value to the table
 }
table <- as.data.frame(table)</pre>
 colnames(table) <- names.table # Add appropriate column names to the table
sort(table, decreasing = TRUE)  # Look at the variates that show least significance
##
     cigpday:heartrte bpmeds:prevstrk bpmeds:glucose diabp:cigpday
## 1
           0.1506282
                            0.1492283
                                           0.1189197
                                                         0.1155989 0.1151079
      sex:ldlc age:prevmi cigpday:prevmi hdlc:ldlc bmi:bpmeds prevmi:prevstrk
##
## 1 0.1141483 0.1097987
                               0.1051865 0.0923568 0.0855445
     heartrte:prevmi glucose:prevmi I(sysbp^2) cursmoke:hdlc age:heartrte
                         0.05883116 0.0585469
          0.06451949
                                                  0.05660935
                                                               0.05562064
## 1
      age:hdlc cursmoke:ldlc sex:sysbp sysbp:bpmeds
                                                       age:ldlc
##
                                           0.0300776 0.02915113
## 1 0.0510796
                   0.0417893 0.03623249
     cigpday:glucose prevmi:prevhyp
                                          hdlc sex:glucose diabetes:hdlc
## 1
           0.0291137
                         0.02242217 0.01880445 0.01702301
                                                               0.01394662
##
    diabp:glucose
                      bmi:ldlc totchol:hdlc
                                               bpmeds age:cigpday
        0.01362058 0.009985489 0.009840662 0.0077735 0.006735591
## 1
##
    heartrte:glucose
                         cursmoke totchol:prevmi sysbp:heartrte diabp:prevhyp
## 1
          0.004772297 0.004188557
                                     0.003609581
                                                    0.002926201
                                                                  0.001409115
##
    diabp:cursmoke prevhyp:ldlc
                                          bmi age:prevhyp sysbp:diabetes
        0.001393474
                      0.00066789 0.0006664543 0.0005753017
                                                             0.0004931994
## 1
       I(hdlc^2)
                  diabp:hdlc sysbp:prevhyp cigpday:hdlc prevmi:hdlc
## 1 0.000320732 0.0001422969 0.0001292531 0.0001038006 7.056001e-05
                            diabp totchol:heartrte
                                                      diabp:bmi sysbp:prevmi
##
    diabetes:prevmi
## 1
        6.226049e-05 6.021714e-05
                                      3.512093e-05 2.940165e-05 2.305381e-05
##
              SAY
                      heartite
                                        age totchol:prevhyp
                                                                 I(bmi^2)
## 1 2.396724e-06 9.478088e-07 4.238229e-07
                                               1.203731e-09 2.735937e-11
##
       I(diabp^2)
                        prevmi
                                    prevhyp
## 1 1.257752e-19 1.595006e-22 1.119628e-27
# Remove first three variates from table since starting the 4th variate, reduction becomes significant
anova(Mstep, update(Mstep,. ~ . - cigpday:heartrte - bpmeds:prevstrk - bpmeds:glucose))
## Analysis of Variance Table
##
## Model 1: I(logit(chdrisk)) ~ sex + totchol + age + sysbp + diabp + cursmoke +
##
       cigpday + bmi + diabetes + bpmeds + heartrte + glucose +
##
       prevmi + prevstrk + prevhyp + hdlc + ldlc + I(hdlc^2) + I(bmi^2) +
##
       I(diabp^2) + I(sysbp^2) + sysbp:prevmi + totchol:prevhyp +
       diabetes:prevmi + prevhyp:ldlc + sysbp:prevhyp + totchol:heartrte +
##
       sysbp:diabetes + diabp:bmi + diabp:hdlc + prevmi:hdlc + prevmi:prevhyp +
##
       sex:glucose + age:ldlc + age:heartrte + cigpday:hdlc + bmi:ldlc +
##
##
       totchol:hdlc + totchol:prevmi + sysbp:heartrte + sysbp:bpmeds +
       cursmoke:hdlc + prevmi:prevstrk + diabetes:hdlc + sex:sysbp +
##
##
       cigpday:glucose + heartrte:glucose + diabp:glucose + cursmoke:ldlc +
       age:cigpday + age:hdlc + hdlc:ldlc + age:prevhyp + diabp:prevhyp +
##
       diabp:cursmoke + diabp:cigpday + bmi:bpmeds + bpmeds:glucose +
##
##
       age:prevmi + sex:ldlc + cigpday:heartrte + cigpday:prevmi +
##
       glucose:prevmi + heartrte:prevmi + bpmeds:prevstrk
## Model 2: I(logit(chdrisk)) ~ sex + totchol + age + sysbp + diabp + cursmoke +
##
       cigpday + bmi + diabetes + bpmeds + heartrte + glucose +
       prevmi + prevstrk + prevhyp + hdlc + ldlc + I(hdlc^2) + I(bmi^2) +
##
```

```
I(diabp^2) + I(sysbp^2) + sysbp:prevmi + totchol:prevhyp +
##
##
       diabetes:prevmi + prevhyp:ldlc + sysbp:prevhyp + totchol:heartrte +
       sysbp:diabetes + diabp:bmi + diabp:hdlc + prevmi:hdlc + prevmi:prevhyp +
##
       sex:glucose + age:ldlc + age:heartrte + cigpday:hdlc + bmi:ldlc +
##
       totchol:hdlc + totchol:prevmi + sysbp:heartrte + sysbp:bpmeds +
##
##
       cursmoke:hdlc + prevmi:prevstrk + diabetes:hdlc + sex:sysbp +
       cigpday:glucose + heartrte:glucose + diabp:glucose + cursmoke:ldlc +
##
       age:cigpday + age:hdlc + hdlc:ldlc + age:prevhyp + diabp:prevhyp +
##
##
       diabp:cursmoke + diabp:cigpday + bmi:bpmeds + age:prevmi +
##
       sex:ldlc + cigpday:prevmi + glucose:prevmi + heartrte:prevmi
##
              RSS Df Sum of Sq
                                    F Pr(>F)
## 1
      2240 489.70
      2243 491.12 -3 -1.4219 2.1681 0.08981 .
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

4 Model Diagnostics

- 5 Model Selection
- 6 Discussion