Midterm project

Tingyi Li

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<pre>library(ISLR) library(caret) library(tidymodels) library(pls) library(earth) library(ggplot2) library(patchwork) library(corrplot) library(gtsummary)</pre>	
<pre>load("./recovery.RData") data <- subset(dat, select = -id) data\$study <- ifelse(data\$study == "A", 0, 1) data <- data > mutate(gender = as.factor(gender), race = as.factor(race), smoking = as.factor(smoking), hypertension = as.factor(hypertension), diabetes = as.factor(diabetes),</pre>	
<pre>vaccine = as.factor(vaccine), severity = as.factor(severity),</pre>	

```
study = as.factor(study)
)
```

Data Partition

Divide data into training data (80%) and testing data (20%)

```
set.seed(666)
data_split <- initial_split(data, prop = 0.8)

# training_data
training_data <- training(data_split)
training_data <- na.omit(training_data)

# testing_data <- testing(data_split)
testing_data <- na.omit(testing_data)

# matrix of predictors
x <- model.matrix(recovery_time ~ ., training_data) [ ,-1]
y <- training_data$recovery_time

x2 <- model.matrix(recovery_time ~ ., testing_data) [ ,-1]
y2 <- testing_data$recovery_time

# 10-fold_cross_validation
ctrl1 <- trainControl(method = "cv", number = 10)</pre>
```

EDA

Summary Statistics

```
summ_dat<-dat%>%
  select(-id)%>%
  mutate(
    gender = factor(case_when(
      gender == "1" ~ "male",
      gender == "0" ~ "female"),
      levels = c("female", "male")
   ),
   race= factor(case_when(
     race == "1" ~ "White",
     race == "2" ~ "Asian",
     race == "3" ~ "Black",
     race == "4" ~ "Hispanic"),
     levels = c("White", "Asian", "Black", "Hispanic")
    smoking=factor(case_when(
     smoking == "0" ~ "Never smoked",
     smoking == "1" ~ "Former smoker",
      smoking == "2" ~ "Current smoker"),
```

```
levels = c("Never smoked", "Former smoker", "Current smoker")
   ),
   hypertension=factor(case_when(
     hypertension == "0" ~ "No",
     hypertension == "1" ~ "Yes"),
     levels = c("No", "Yes")
   ),
   diabetes=factor(case when(
     diabetes == "0" ~ "No",
     diabetes == "1" ~ "Yes"),
     levels = c("No", "Yes")
   ),
   vaccine=factor(case when(
     vaccine == "0" ~ "Not vaccinated",
     vaccine == "1" ~ "Vaccinated"),
     levels = c("Not vaccinated", "Vaccinated")
   ),
   severity=factor(case_when(
     severity == "0" ~ "Not severe",
     severity == "1" ~ "Severe"),
     levels = c("Not severe", "Severe")
   )
    )
summ_dat %>%
 tbl_summary() %>%
 bold_labels()%>%
  as_gt(include = everything()) %>%
 gt::tab_header("Table 1: Summary of Dataset")
```

Table 1: Summary of Dataset

Characteristic	$\mathbf{N}=3{,}000^{1}$
age	60.0 (57.0, 63.0)
\mathbf{gender}	
female	1,544 (51%)
male	1,456 (49%)
race	
White	1,967~(66%)
Asian	158 (5.3%)
Black	604 (20%)
Hispanic	271 (9.0%)
$\mathbf{smoking}$, ,
Never smoked	1,822 (61%)
Former smoker	859 (29%)
Current smoker	319 (11%)
height	169.9 (166.0, 173.9)
\mathbf{weight}	80 (75, 85)
bmi	$27.65\ (25.80,\ 29.50)$
hypertension	1,492 (50%)
diabetes	463 (15%)
SBP	$130\ (125,\ 136)$
LDL	110 (97, 124)

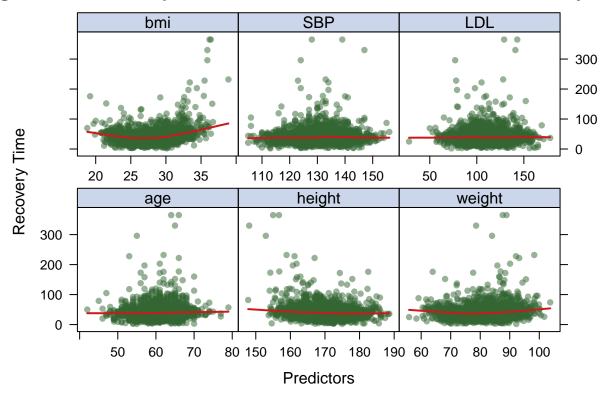
vaccine	
Not vaccinated	1,212 (40%)
Vaccinated	1,788 (60%)
severity	
Not severe	2,679 (89%)
Severe	321 (11%)
study	
A	2,000 (67%)
В	1,000 (33%)
${\bf recovery_time}$	39 (31, 49)

 $^{^{1}}$ Median (IQR); n (%)

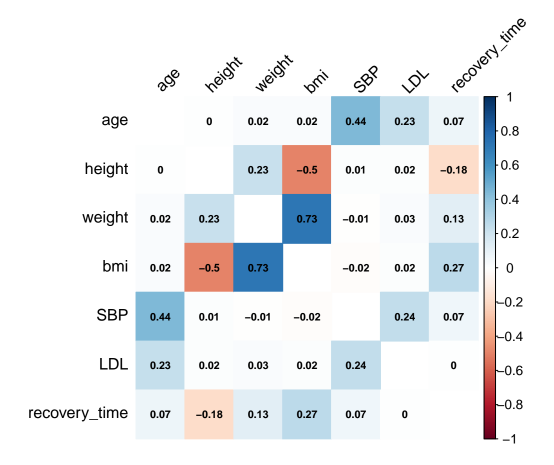
Feature plot for continuous variables

```
theme1 <- trellis.par.get()</pre>
theme1$plot.symbol$col <- rgb(.2, .4, .2, .5)
theme1$plot.symbol$pch <- 16</pre>
theme1$plot.line$col <- rgb(.8, .1, .1, 1)
theme1$plot.line$lwd <- 2</pre>
theme1$strip.background$col <- rgb(.0, .2, .6, .2)
trellis.par.set(theme1)
trainData <- training_data|>
  dplyr::select('age', 'height', 'weight', 'bmi', 'SBP', 'LDL', 'recovery_time')
featurePlot(x = trainData[ , 1:6],
            y = trainData[ , 7],
            plot = "scatter",
            span = .5,
            labels = c("Predictors", "Recovery Time"),
            main = "Figure 1:Relationship between Continuous Predictors and Recovery Time",
            type = c("p", "smooth"),
            layout = c(3, 2)
```

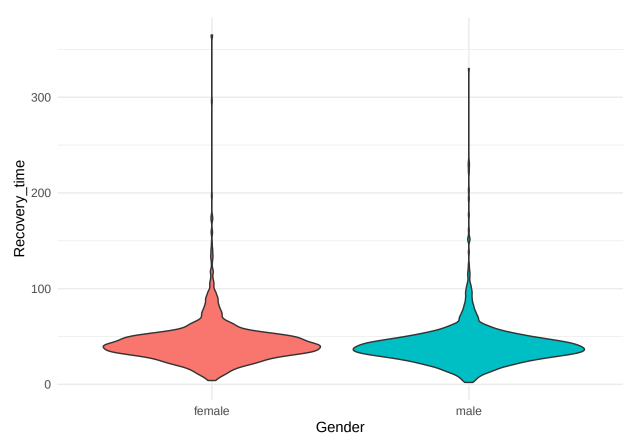
igure 1:Relationship between Continuous Predictors and Recovery Tim

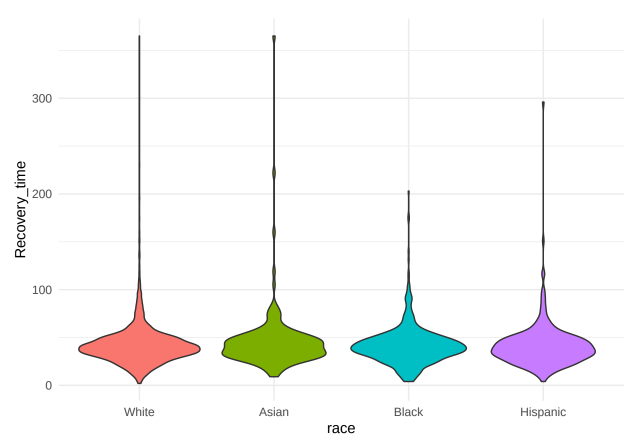


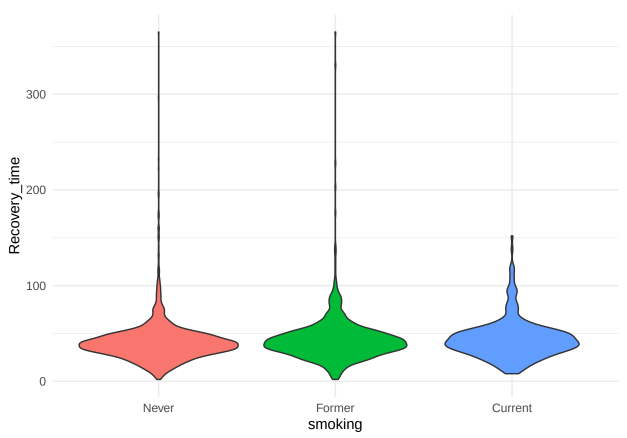
Correlation plot

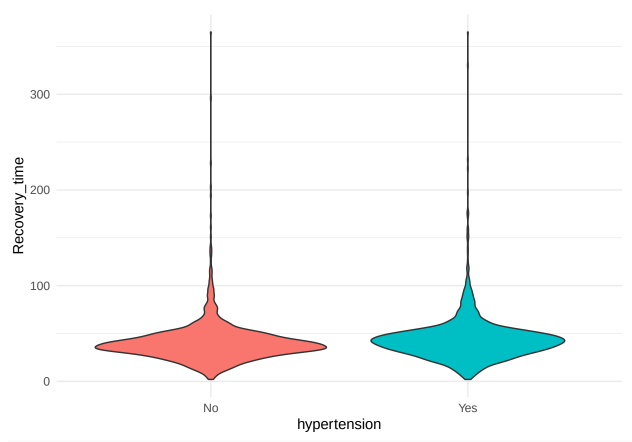


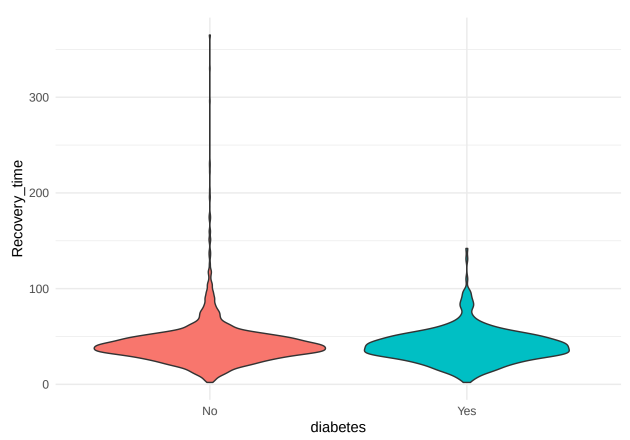
Violin plot for categorical variables

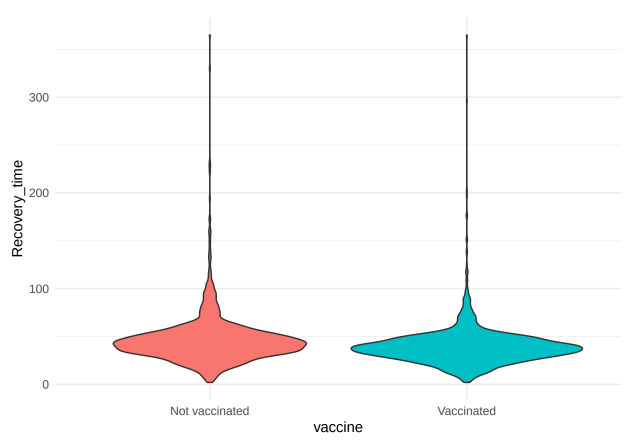




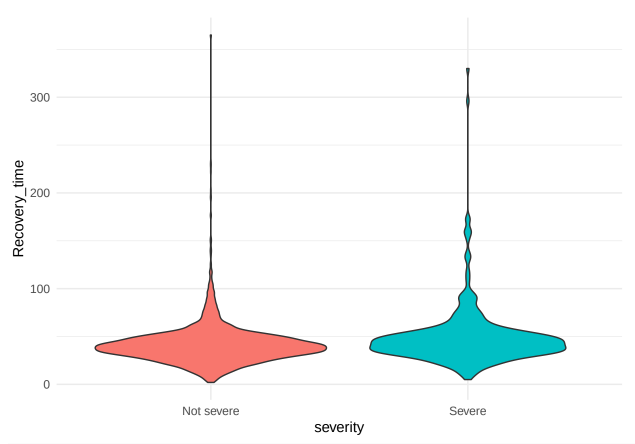


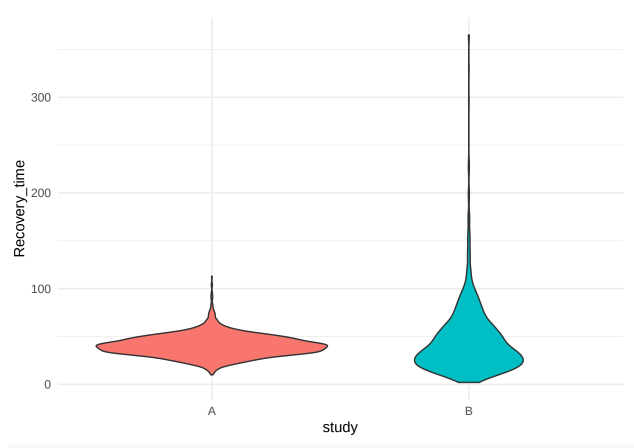






```
severity_plot <- data |>
    ggplot(aes(x = severity, y = recovery_time, fill = severity)) +
    geom_violin() +
    scale_x_discrete(labels = c("Not severe", "Severe")) +
    labs(
        x = "severity",
        y = "Recovery_time") +
    theme_minimal() + theme(legend.position = "none")
severity_plot
```





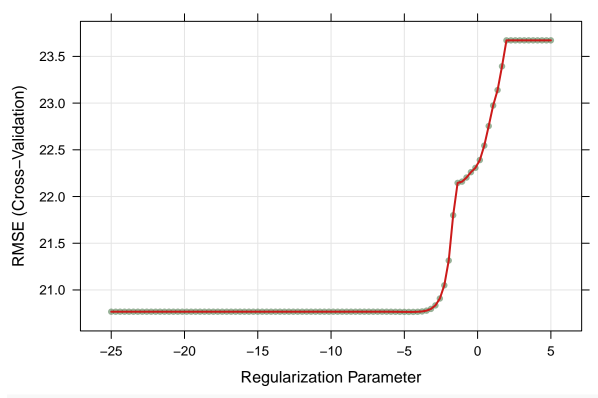
combined <- gender_plot + race_plot + smoking_plot + hypertension_plot + diabetes_plot + vaccine_plot +
combined + plot_annotation(title = "Figure 2: Relationship between Categorical Predictors and Recovery")</pre>

Recovery_time Recovery_time Recovery_time 300 300 300 200 200 200 100 100 100 0 0 0 female White Asian Black-lispanic Never Former Current male Gender smoking race Recovery_time Recovery_time Recovery_time 300 300 300 200 200 200 100 100 100 0 0 0 No Yes No Yes Not vaccinated/accinated hypertension diabetes vaccine Recovery_time Recovery_time 300 300 200 200 100 100 0 Not severe Severe В severity study

Figure 2: Relationship between Categorical Predictors and Recovery Time

Linear models

Fit a lasso model



Select the best tuning parameter lasso.fit\$bestTune

```
## alpha lambda
## 68 1 0.00912288

# coefficients in the final model
coef(lasso.fit$finalModel, lasso.fit$bestTune$lambda)
```

```
## 18 x 1 sparse Matrix of class "dgCMatrix"
##
## (Intercept)
                 -2.229189e+03
                  2.736904e-01
## age
## gender1
                 -3.698081e+00
## race2
                  1.629521e+00
## race3
                 -7.478838e-01
## race4
                 -8.493986e-01
## smoking1
                  2.764637e+00
## smoking2
                  3.624109e+00
## height
                  1.302739e+01
## weight
                 -1.414814e+01
## bmi
                  4.250183e+01
## hypertension1 3.023576e+00
## diabetes1
                 -1.066279e+00
## SBP
                 -1.899186e-02
## LDL
                 -3.709871e-02
## vaccine1
                 -6.014455e+00
## severity1
                  7.390563e+00
                  5.298800e+00
## study1
```

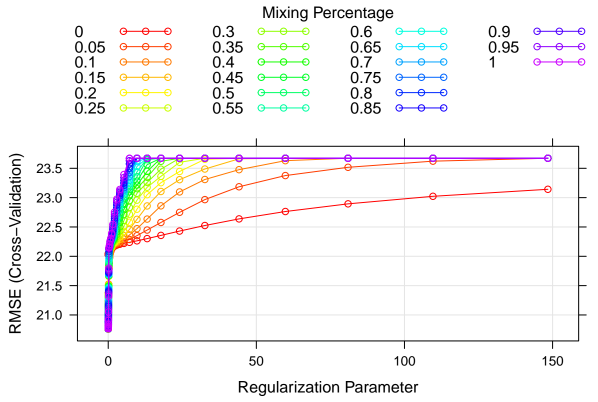
The selected tuning parameter is 0.00912288.

Test errors

```
set.seed(666)
lasso.pred <- predict(lasso.fit, newdata = testing_data)
mean((lasso.pred - testing_data[, "recovery_time"])^2)
## [1] 298.3016
The test error is 298.3016.</pre>
```

Fit an elastic net model

```
set.seed(666)
enet.fit <- train(recovery_time ~ .,</pre>
                  data = training_data,
                  method = "glmnet",
                  tuneGrid = expand.grid(alpha = seq(0, 1, length = 21),
                                          lambda = exp(seq(-25, 5, length = 100))),
                  trControl = ctrl1)
## Warning in nominalTrainWorkflow(x = x, y = y, wts = weights, info = trainInfo,
## : There were missing values in resampled performance measures.
## Select the best tuning parameter
enet.fit$bestTune
                 lambda
##
       alpha
## 365 0.15 0.00367552
#Plot
myCol <- rainbow(25)</pre>
myPar <- list(superpose.symbol = list(col = myCol),</pre>
              superpose.line = list(col = myCol))
plot(enet.fit, par.settings = myPar)
```



coefficients in the final model
coef(enet.fit\$finalModel, enet.fit\$bestTune\$lambda)

```
## 18 x 1 sparse Matrix of class "dgCMatrix"
##
                             s1
                 -2.157409e+03
## (Intercept)
                  2.786152e-01
## age
## gender1
                  -3.708843e+00
## race2
                   1.666442e+00
## race3
                 -7.696907e-01
## race4
                 -9.113345e-01
## smoking1
                  2.780175e+00
## smoking2
                  3.645642e+00
## height
                   1.260703e+01
## weight
                  -1.370319e+01
## bmi
                   4.122283e+01
## hypertension1
                  3.105223e+00
## diabetes1
                 -1.101528e+00
## SBP
                 -2.443694e-02
## LDL
                 -3.746249e-02
## vaccine1
                 -6.040658e+00
## severity1
                  7.430536e+00
## study1
                  5.315665e+00
# test error
enet.pred <- predict(enet.fit, newdata = testing_data)</pre>
mean((enet.pred - testing_data[, "recovery_time"])^2)
```

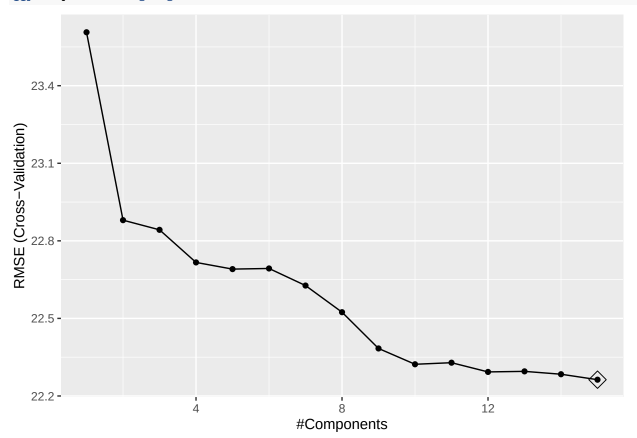
[1] 297.1642

The selected tuning parameter is 0.00367552, and the test error is 297.1642.

Fit a PCR model

[1] 323.779

ggplot(pcr.fit, highlight = TRUE)



```
summary(pcr.fit)
```

```
## Data:
            X dimension: 2400 17
## Y dimension: 2400 1
## Fit method: svdpc
## Number of components considered: 15
## TRAINING: % variance explained
##
            1 comps 2 comps 3 comps 4 comps 5 comps
                                                         6 comps 7 comps
## X
             13.117
                      23.728
                               31.171
                                        38.302
                                                 45.266
                                                          51.762
                                                                    57.87
## .outcome
              0.602
                       7.376
                                7.832
                                         9.341
                                                  9.353
                                                           9.575
                                                                    10.59
##
            8 comps 9 comps 10 comps 11 comps 12 comps 13 comps 14 comps
```

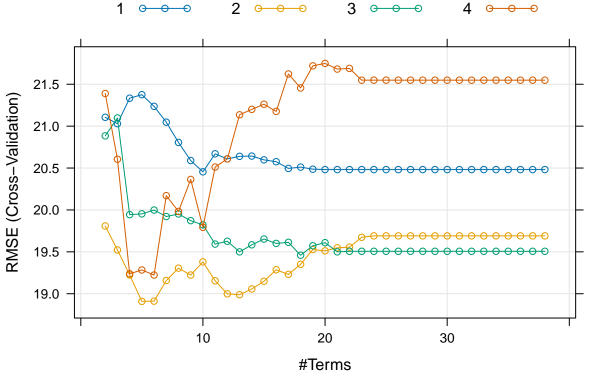
```
## X
                63.91
                         69.84
                                    75.59
                                              81.01
                                                         85.98
                                                                    90.51
                                                                              94.88
                11.16
                         12.02
                                    12.51
                                              12.51
                                                                    12.98
## .outcome
                                                         12.88
                                                                              13.16
##
             15 comps
## X
                 98.82
                 13.32
## .outcome
```

There are 15 components in the model and the test error is 323.779.

Nonlinear models

Multivariate Adaptive Regression Spline (MARS)

Product Degree



```
# best tune
mars.fit$bestTune

## nprune degree
## 41 5 2

coef(mars.fit$finalModel)
```

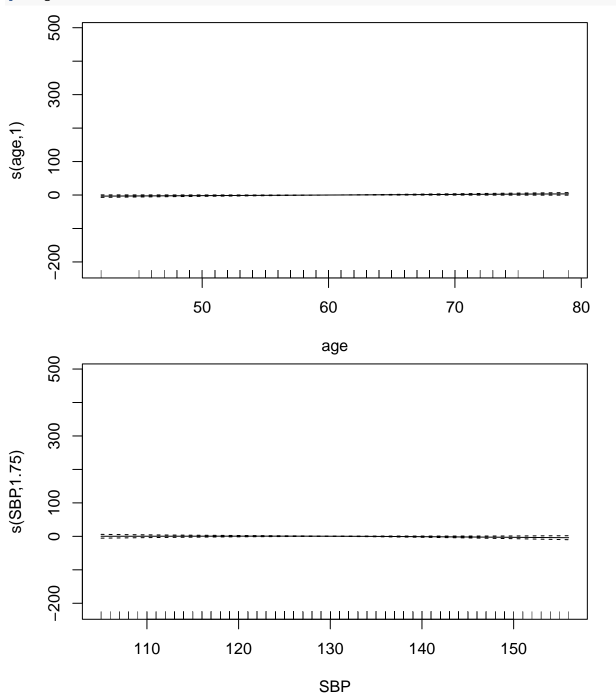
```
##
                   (Intercept)
                                                  h(31-bmi)
                    -3.1983530
                                                  6.3999877
##
            h(bmi-31) * study1
                                               h(bmi-25.2)
##
                                                  7.9260754
##
                    25.6820131
## h(weight-86.4) * h(bmi-31)
                    -0.6277843
Test error for MARS
mars.pred <- predict(mars.fit, newdata = x2)</pre>
mars.test.error <- mean((mars.pred - y2)^2)</pre>
mars.test.error
## [1] 279.0367
The regression function should be: -3.1983530 + 6.3999877 * h(31-bmi) + 25.6820131 * h(bmi-31) * study1
+7.9260754 * h(bmi-25.2) - 0.6277843 * h(weight-86.4) * h(bmi-31)
The test error is 279.0367
GAM
set.seed(666)
gam.fit <- train(x, y,</pre>
                  method = "gam",
                  tuneGrid = data.frame(method = "GCV.Cp", select = c(TRUE,FALSE)),
                  trControl = ctrl1)
## Loading required package: mgcv
## Loading required package: nlme
##
## Attaching package: 'nlme'
## The following object is masked from 'package:dplyr':
##
##
       collapse
## This is mgcv 1.9-0. For overview type 'help("mgcv-package")'.
gam.fit$bestTune
     select method
## 1 FALSE GCV.Cp
gam.fit$finalModel
##
## Family: gaussian
## Link function: identity
##
## Formula:
   .outcome ~ gender1 + race2 + race3 + race4 + smoking1 + smoking2 +
       hypertension1 + diabetes1 + vaccine1 + severity1 + study1 +
##
##
       s(age) + s(SBP) + s(LDL) + s(bmi) + s(height) + s(weight)
##
## Estimated degrees of freedom:
## 1.00 1.75 1.00 8.56 7.24 2.81 total = 34.36
```

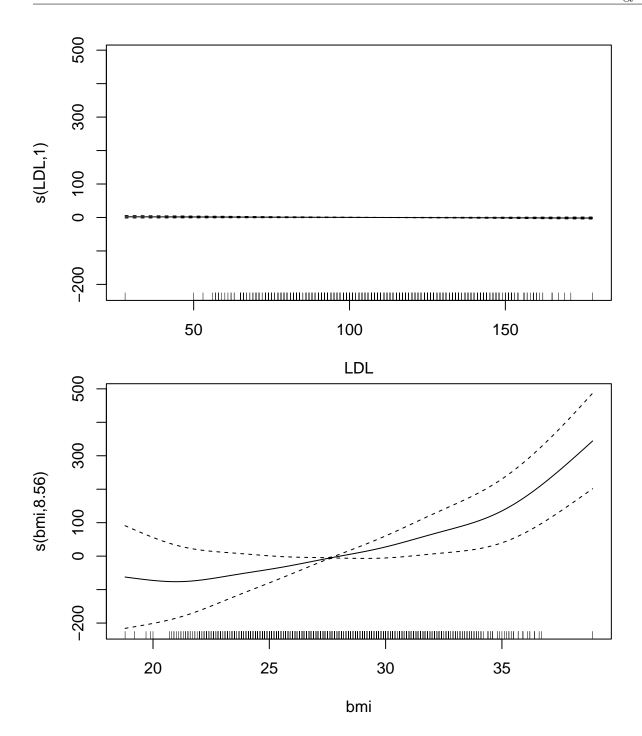
GCV score: 384.8296

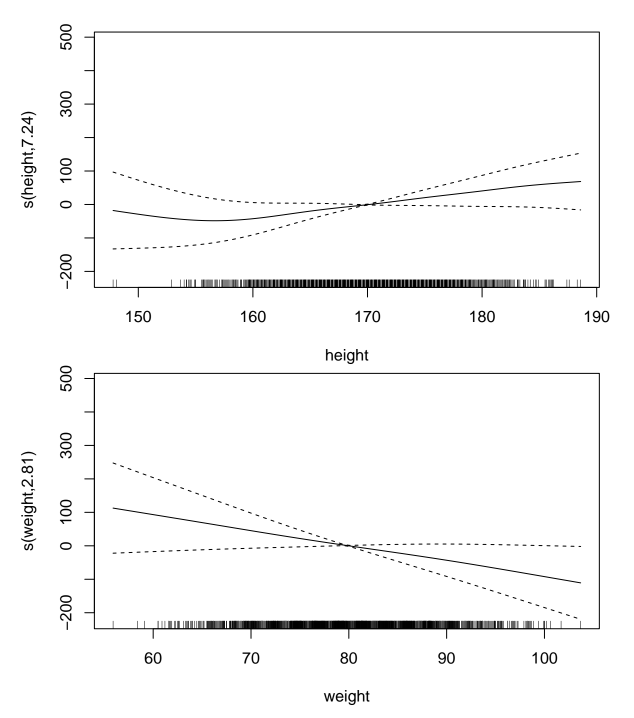
The GAM model includes all the predictors.

Generate plot for GAM

plot(gam.fit\$finalModel)







Test error for GAM

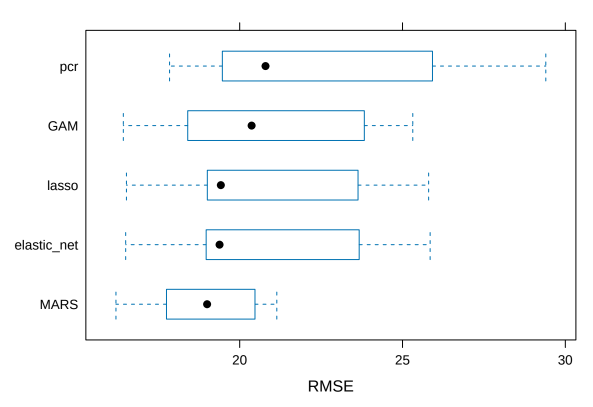
```
gam.pred <- predict(gam.fit, newdata = x2)
gam.test.error <- mean((gam.pred - y2)^2)
gam.test.error</pre>
```

[1] 272.0012

The test error for GAM is 272.0012.

Model comparison

```
resamp <- resamples(list(lasso = lasso.fit,</pre>
                         elastic_net = enet.fit,
                         pcr = pcr.fit,
                         MARS = mars.fit,
                         GAM = gam.fit))
summary(resamp)
##
## Call:
## summary.resamples(object = resamp)
## Models: lasso, elastic_net, pcr, MARS, GAM
## Number of resamples: 10
##
## MAE
##
                        1st Qu.
                                   Median
                                               Mean 3rd Qu.
                   Min.
               12.43437 13.34989 13.55760 13.60117 13.83628 14.80446
## elastic_net 12.39330 13.33858 13.50759 13.56655 13.79648 14.79199
                                                                          0
## pcr
               12.79843 13.31630 13.65714 13.79429 14.32701 15.26141
## MARS
               11.10656 12.33992 12.52758 12.48434 12.95145 13.22857
                                                                          0
## GAM
               11.95625 12.48434 12.90413 13.04720 13.67832 14.43838
##
## RMSE
##
                        1st Qu.
                                   Median
                                               Mean 3rd Qu.
                   Min.
                                                                 Max. NA's
               16.52182 19.01349 19.42116 20.76489 22.92278 25.80217
## lasso
## elastic net 16.49848 18.99457 19.38126 20.76102 22.93865 25.84810
               17.84596 19.61842 20.79266 22.26300 24.74262 29.40059
## pcr
## MARS
               16.19940 17.79482 18.99934 18.90878 20.37825 21.13959
                                                                          0
## GAM
               16.42592 18.51665 20.36608 20.61337 23.26164 25.31540
##
## Rsquared
                             1st Qu.
                                        Median
                                                            3rd Qu.
##
                     Min.
                                                     Mean
               0.12468127 0.17202217 0.2314610 0.2404963 0.2896872 0.3926191
## lasso
## elastic_net 0.12572907 0.17252570 0.2295363 0.2401821 0.2906348 0.3898854
               0.05140385\ 0.09539054\ 0.1072900\ 0.1273979\ 0.1285069\ 0.2390132
                                                                                  0
## pcr
               0.12610733\ 0.18362545\ 0.2804839\ 0.3515976\ 0.5442009\ 0.6258534
## MARS
                                                                                  0
## GAM
               0.11451833 0.18716669 0.2610268 0.2888378 0.3891996 0.4926768
bwplot(resamp, metric = "RMSE")
```



Mars model should be selected as the final model since it has the lowest RMSE value.