

Midterm Project Analysis

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```
library(tidyverse)
library(patchwork)
library(caret)
library(mgcv)
library(AppliedPredictiveModeling)
library(pdp)
library(corrplot)
library(plotmo)
library(ggrepel)
```

Data Wrangling

```
load("recovery.RData")
dat = dat %>%
  select(-id) %>%
  mutate(
    gender = as.factor(gender),
    race = as.factor(race),
    smoking = as.factor(smoking),
    hypertension = as.factor(hypertension),
    diabetes = as.factor(diabetes),
    vaccine = as.factor(vaccine),
    severity = as.factor(severity),
    study = as.factor(study))
```

Summary Statistics

```
summary(dat)
```

```
##      age      gender  race  smoking      height      weight
##  Min.   :42.0    0:1544  1:1967  0:1822  Min.   :147.8  Min.   : 55.90
##  1st Qu.:57.0    1:1456  2: 158  1: 859  1st Qu.:166.0  1st Qu.: 75.20
##  Median :60.0              3: 604  2: 319  Median :169.9  Median : 79.80
##  Mean   :60.2              4: 271      Mean   :169.9  Mean   : 79.96
##  3rd Qu.:63.0              Mean   :169.9  Mean   : 79.96
##  Max.   :79.0              3rd Qu.:173.9  3rd Qu.: 84.80
##                               Max.   :188.6  Max.   :103.70
##      bmi      hypertension diabetes      SBP      LDL      vaccine
##  Min.   :18.80    0:1508      0:2537  Min.   :105.0  Min.   : 28.0  0:1212
##  1st Qu.:25.80    1:1492      1: 463  1st Qu.:125.0  1st Qu.: 97.0  1:1788
##  Median :27.65              Median :130.0  Median :110.0
##  Mean   :27.76              Mean   :130.5  Mean   :110.5
##  3rd Qu.:29.50              3rd Qu.:136.0  3rd Qu.:124.0
##  Max.   :38.90              Max.   :156.0  Max.   :178.0
##  severity study  recovery_time
##  0:2679  A:2000  Min.   : 2.00
##  1: 321   B:1000  1st Qu.: 31.00
##                               Median : 39.00
##                               Mean   : 42.17
##                               3rd Qu.: 49.00
```

```
##                               Max.      :365.00
```

```
sum(is.na(dat))
```

```
## [1] 0
```

```
skimr::skim(dat)
```

Table 1: Data summary

Name	dat
Number of rows	3000
Number of columns	15
Column type frequency:	
factor	8
numeric	7
Group variables	None

Variable type: factor

skim_variable	n_missing	complete_rate	ordered	n_unique	top_counts
gender	0	1	FALSE	2	0: 1544, 1: 1456
race	0	1	FALSE	4	1: 1967, 3: 604, 4: 271, 2: 158
smoking	0	1	FALSE	3	0: 1822, 1: 859, 2: 319
hypertension	0	1	FALSE	2	0: 1508, 1: 1492
diabetes	0	1	FALSE	2	0: 2537, 1: 463
vaccine	0	1	FALSE	2	1: 1788, 0: 1212
severity	0	1	FALSE	2	0: 2679, 1: 321
study	0	1	FALSE	2	A: 2000, B: 1000

Variable type: numeric

skim_variable	n_missing	complete_rate	mean	sd	p0	p25	p50	p75	p100	hist
age	0	1	60.20	4.48	42.0	57.0	60.00	63.0	79.0	
height	0	1	169.90	5.97	147.8	166.0	169.90	173.9	188.6	
weight	0	1	79.96	7.14	55.9	75.2	79.80	84.8	103.7	
bmi	0	1	27.76	2.79	18.8	25.8	27.65	29.5	38.9	
SBP	0	1	130.47	7.97	105.0	125.0	130.00	136.0	156.0	
LDL	0	1	110.45	19.76	28.0	97.0	110.00	124.0	178.0	
recovery_time	0	1	42.17	23.15	2.0	31.0	39.00	49.0	365.0	

The “recovery” dataset contains 15 columns and 3000 observations without any missing values after omitting the `id` variable. We have 14 predictors (6 numeric and 8 factor(character) variables) in the dataset. Then we partition the dataset into two parts: training data (80%) and test data (20%).

Model training Preparation

```

# data partition
set.seed(2358)
indexTrain <- createDataPartition(y = dat$recovery_time, p = 0.8, list = FALSE)
train <- dat[indexTrain, ]
test <- dat[-indexTrain, ]

# matrix of predictors
x <- model.matrix(recovery_time~.,train)[,-1]
x2 <- model.matrix(recovery_time~.,test)[,-1]

# vector of response
y <- train$recovery_time
y2 <- test$recovery_time

```

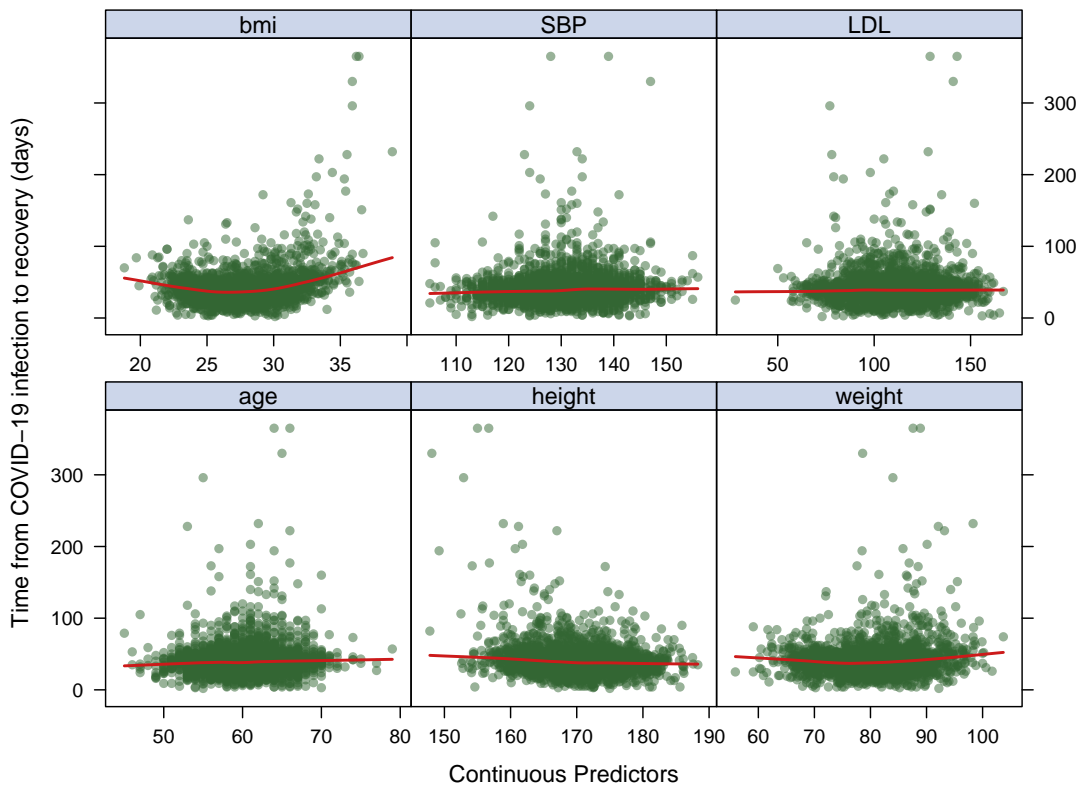
Expoloratory Data Analysis

```

# Remove all the categorical predictors out from x before plotting
x_continuous <-
  x[, !(colnames(x) %in%
        c("gender1", "race2","race3", "race4",
          "smoking1","smoking2", "hypertension1",
          "diabetes1", "vaccine1", "severity1", "studyB"))]

# For Continuous Predictors
theme1 <- trellis.par.get()
theme1$plot.symbol$col <- rgb(.2, .4, .2, .5)
theme1$plot.symbol$pch <- 16
theme1$plot.line$col <- rgb(.8, .1, .1, 1)
theme1$plot.line$lwd <- 2
theme1$strip.background$col <- rgb(.0, .2, .6, .2)
trellis.par.set(theme1)
featurePlot(x_continuous,
            y,
            plot = "scatter",
            span = .5,
            labels = c("Continuous Predictors","Time from COVID-19 infection to recovery (days)"),
            type = c("p", "smooth"),
            layout = c(3, 2))

```



```
# For Categorical Predictors
dis_gender = train %>%
  ggplot(aes(x = gender, y = recovery_time)) +
  geom_violin(fill = "orange", color = "blue", alpha = .5) +
  scale_x_discrete(labels = c('Female', 'Male'))

dis_race = train %>%
  ggplot(aes(x = race, y = recovery_time)) +
  geom_violin(fill = "orange", color = "blue", alpha = .5) +
  scale_x_discrete(labels = c('White', 'Asian', 'Black', 'Hispanic'))

dis_smoking = train %>%
  ggplot(aes(x = smoking, y = recovery_time)) +
  geom_violin(fill = "orange", color = "blue", alpha = .5) +
  scale_x_discrete(labels = c('Never smoked', 'Former smoker', 'Current smoker'))

dis_hyper = train %>%
  ggplot(aes(x = hypertension, y = recovery_time)) +
  geom_violin(fill = "orange", color = "blue", alpha = .5) +
  scale_x_discrete(labels = c('No', 'Yes'))

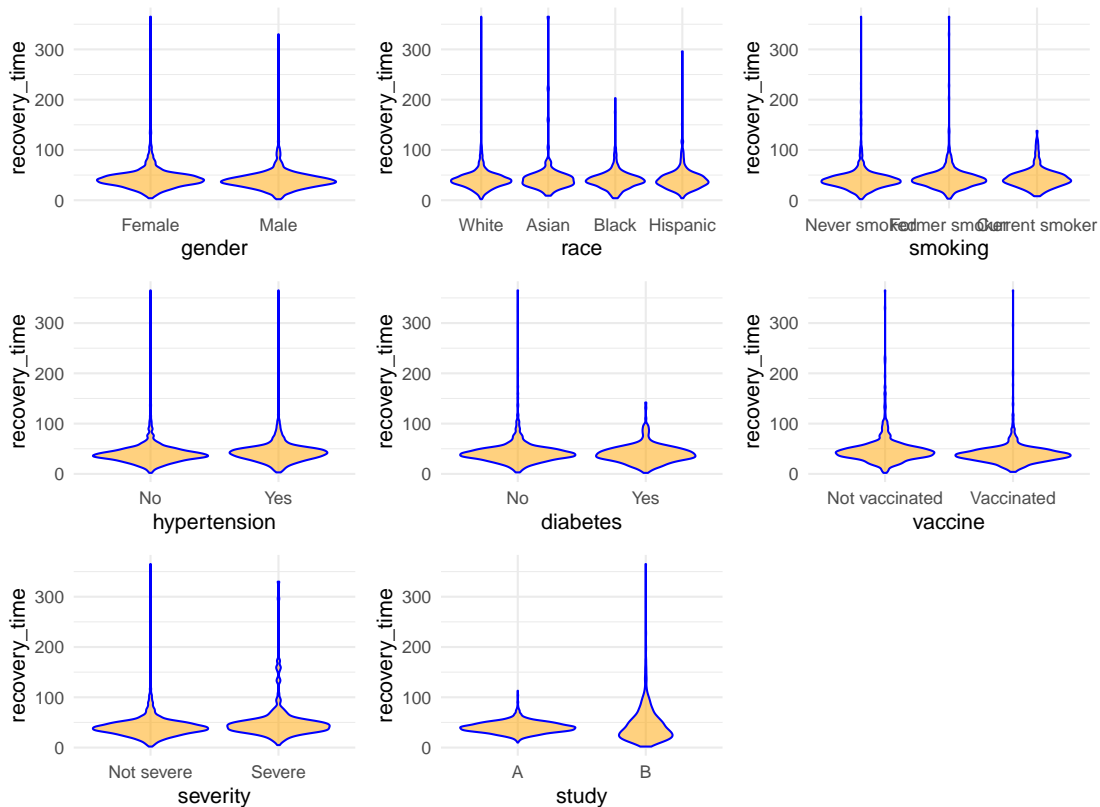
dis_diabetes = train %>%
  ggplot(aes(x = diabetes, y = recovery_time)) +
  geom_violin(fill = "orange", color = "blue", alpha = .5) +
  scale_x_discrete(labels = c('No', 'Yes'))
```

```
dis_vac = train %>%
  ggplot(aes(x = vaccine, y = recovery_time)) +
  geom_violin(fill = "orange", color = "blue", alpha = .5) +
  scale_x_discrete(labels = c('Not vaccinated', 'Vaccinated'))
```

```
dis_serverity = train %>%
  ggplot(aes(x = severity, y = recovery_time)) +
  geom_violin(fill = "orange", color = "blue", alpha = .5) +
  scale_x_discrete(labels = c('Not severe', 'Severe'))
```

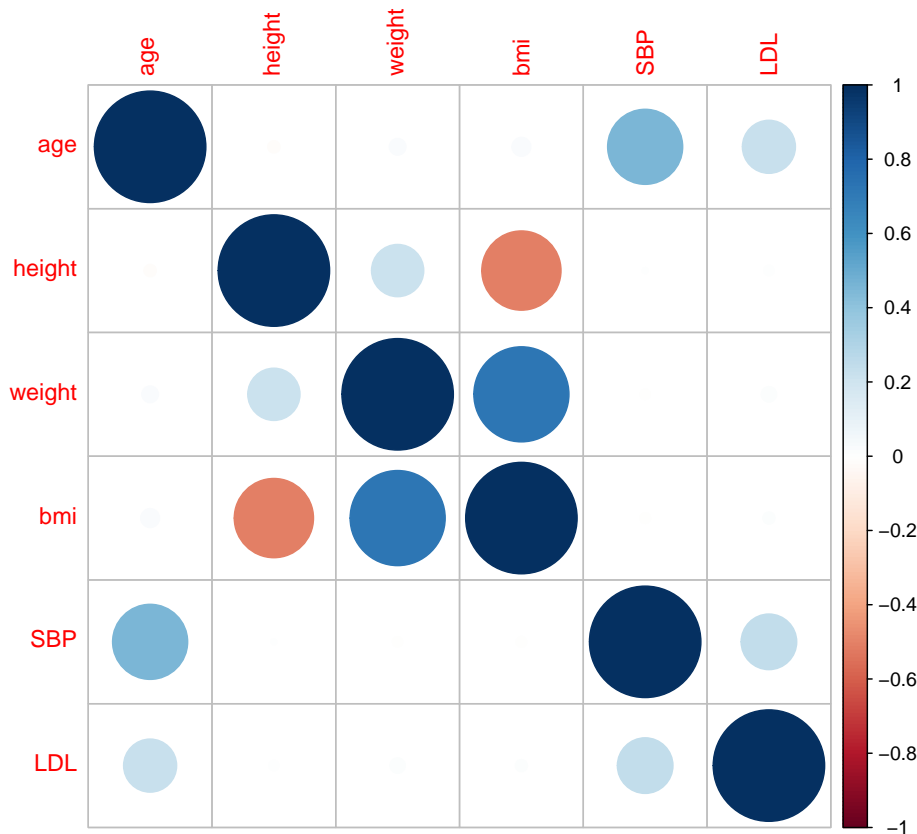
```
dis_study = train %>%
  ggplot(aes(x = study, y = recovery_time)) +
  geom_violin(fill = "orange", color = "blue", alpha = .5)
```

```
dis_gender + dis_race + dis_smoking + dis_hyper + dis_diabetes + dis_vac + dis_serverity + dis_study +
```



```
# Correlation plots
```

```
corrplot(cor(x_continuous), method = "circle", type = "full")
```



From the correlation matrix, some multicollinearities are severe shown in the continuous predictors of training data, cross-validation will be applied in the next steps.

Next, we will fit 6 linear models(Linear, Lasso, ridge, elastic net, PCR, PLS) and 2 non-linear models(GAM, MARS) using caret, and conduct the model comparison to choose the best fitted one.

Linear

```
# 10-fold cv on best
ctrl1 <- trainControl(method = "cv", number = 10)

# Fitting the model with Cross-validation
set.seed(2358)
lm.fit <- train(x, y, method = "lm", trControl = ctrl1)

# Final model coefficients
coef(lm.fit$finalModel)
```

```
## (Intercept)      age      gender1      race2      race3
## -2.253974e+03  2.085591e-01 -2.727477e+00  2.089360e+00 -1.155809e+00
##      race4      smoking1      smoking2      height      weight
## -2.959891e-01  2.280249e+00  3.185637e+00  1.310556e+01 -1.426157e+01
##      bmi hypertension1      diabetes1      SBP      LDL
##  4.285697e+01  1.752483e+00 -2.040537e+00  9.762877e-02 -3.373236e-02
##      vaccine1      severity1      studyB
## -7.247882e+00  7.938166e+00  5.629688e+00
```

```
# Make prediction on test data
lm.pred <- predict(lm.fit, newdata = x2)

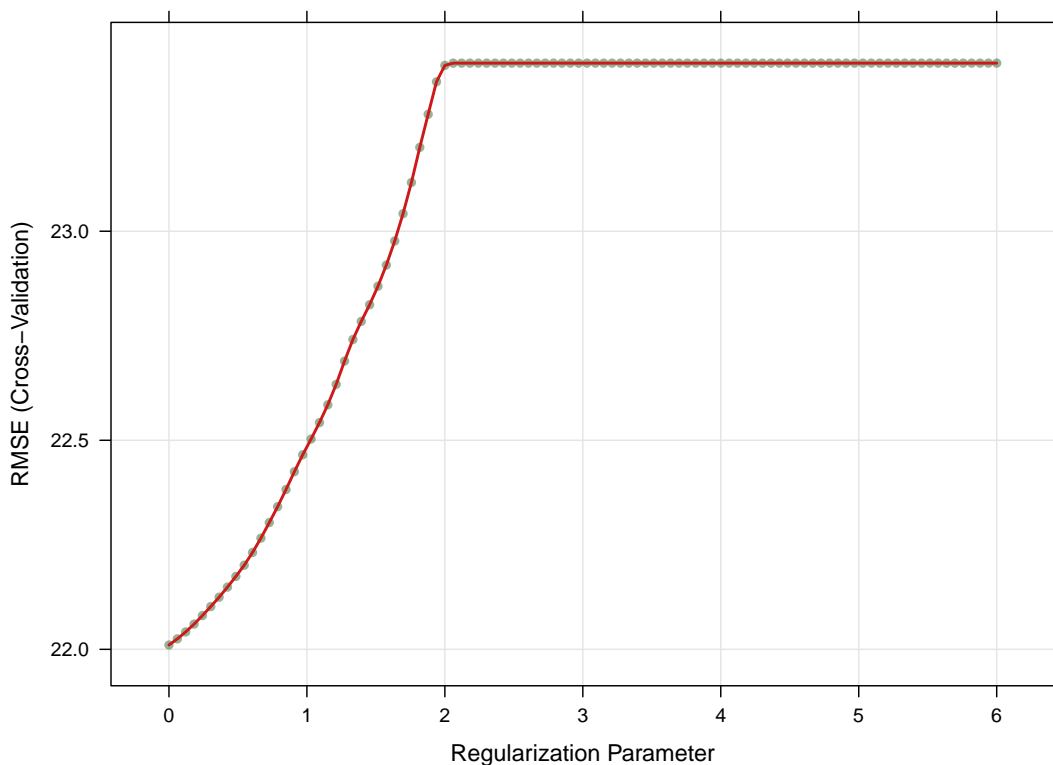
# Test error
lm.mse <- mean((lm.pred - y2)^2)
lm.mse
```

```
## [1] 314.7791
```

Lasso

```
# Fitting the model with Cross-validation
set.seed(2358)
lasso.fit <- train(x, y, method = "glmnet",
                  tuneGrid = expand.grid(alpha = 1,
                                         lambda = exp(seq(6, 0, length=100))),
                  trControl = ctrl1)

plot(lasso.fit, xTrans = log)
```



```
# Tuning parameters
lasso.fit$bestTune
```



```
##    alpha lambda
## 1      1      1

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

```
## 18 x 1 sparse Matrix of class "dgCMatrix"
##              s1
## (Intercept) 14.07004988
## age         0.07075850
## gender1     -0.66417312
## race2        .
## race3        .
## race4        .
## smoking1     .
## smoking2     .
## height      -0.20378959
## weight       .
## bmi          1.92393220
## hypertension1 0.61066535
## diabetes1    .
## SBP          0.05151048
## LDL          .
## vaccine1     -5.55415366
## severity1    4.79327540
## studyB       3.63357454
```

```
# Make prediction on test data
lasso.pred <- predict(lasso.fit, newdata = x2)

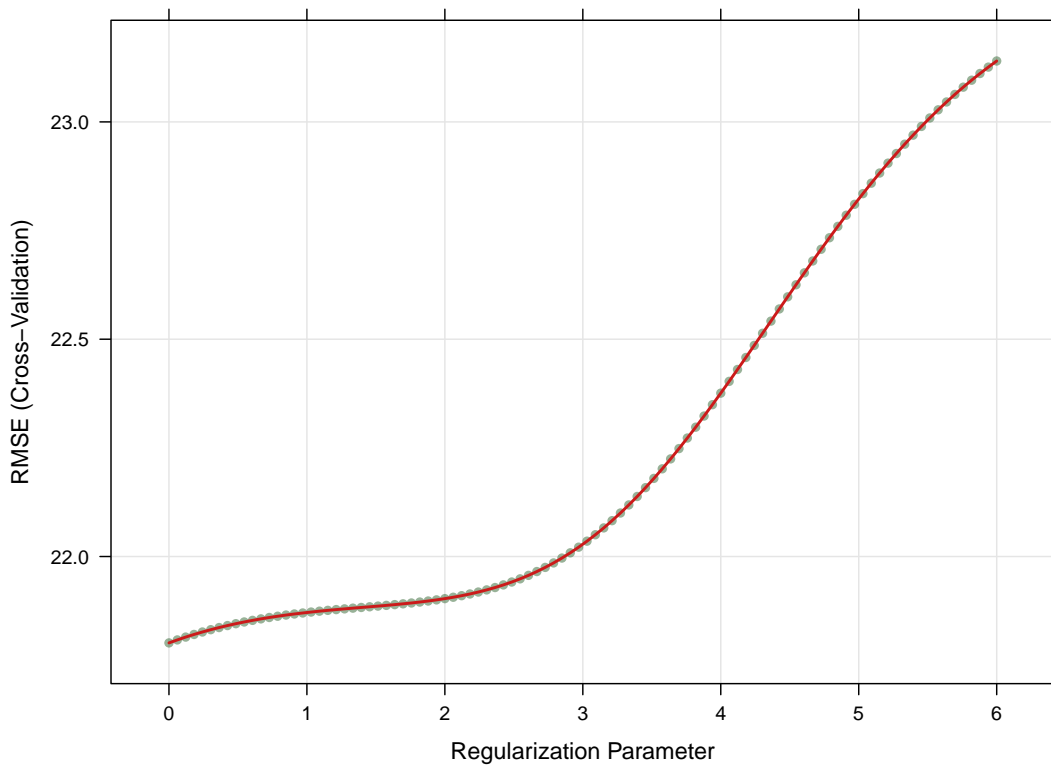
# Test error
lasso.mse <- mean((lasso.pred - y2)^2)
lasso.mse
```

```
## [1] 352.0247
```

Ridge

```
# Fitting the model with Cross-validation
set.seed(2358)
ridge.fit <- train(x, y, method = "glmnet",
                  tuneGrid = expand.grid(alpha = 0,
                                         lambda = exp(seq(6, 0, length=100))),
                  trControl = ctrl1)

plot(ridge.fit, xTrans = log)
```



```
# Tuning parameters
```

```
ridge.fit$bestTune
```

```
##   alpha lambda
```

```
## 1      0      1
```

```
# coefficients in the final model
```

```
coef(ridge.fit$finalModel, s = ridge.fit$bestTune$lambda)
```

```
## 18 x 1 sparse Matrix of class "dgCMatrix"
```

```
##              s1
## (Intercept) -52.70721475
## age         0.22021302
## gender1     -2.48744811
## race2        2.46037359
## race3       -1.08502802
## race4       -0.90753700
## smoking1     2.11825022
## smoking2     2.35768222
## height       0.11928447
## weight      -0.52054391
## bmi          3.42880724
## hypertension1 1.62856332
## diabetes1    -2.15217999
## SBP          0.09818922
```

```
## LDL          -0.02968295
## vaccine1     -7.25775052
## severity1    7.44593749
## studyB       5.66750779

# Make prediction on test data
ridge.pred <- predict(ridge.fit, newdata = x2)

# Test error
ridge.mse <- mean((ridge.pred - y2)^2)
ridge.mse
```

```
## [1] 353.4649
```

Elastic net

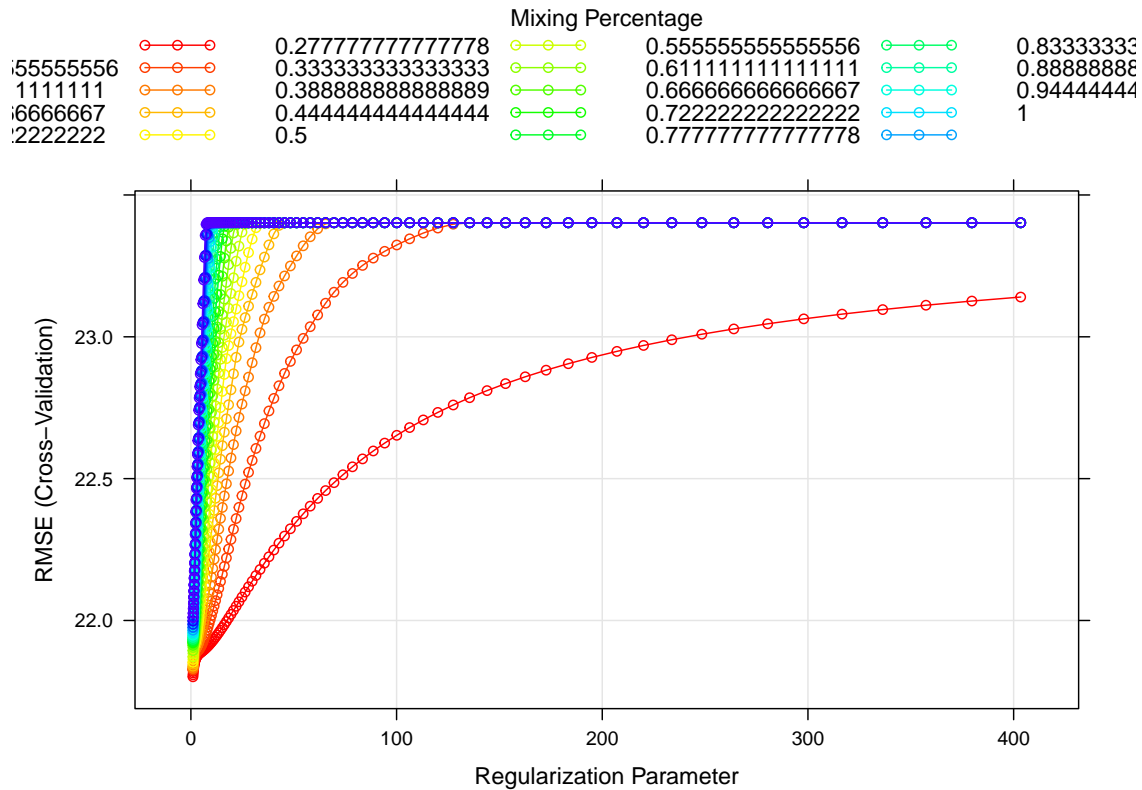
```
# Fitting the model with Cross-validation
set.seed(2358)
enet.fit <- train(x, y, method = "glmnet",
                  tuneGrid = expand.grid(alpha = seq(0, 1, length = 19),
                                          lambda = exp(seq(6, 0, length = 100))),
                  trControl = ctrl1)

# Tuning parameters
enet.fit$bestTune

##      alpha lambda
## 1      0      1

# 25 kinds of Plot colors
myCol <- rainbow(25)
myPar <- list(superpose.symbol = list(col = myCol),
              superpose.line = list(col = myCol))

# Plot CV RMSE-alpha&lambda
plot(enet.fit, par.settings = myPar)
```



```
# Coefficients
coef(enet.fit$finalModel, enet.fit$bestTune$lambda)
```

```
## 18 x 1 sparse Matrix of class "dgCMatrix"
##              s1
## (Intercept) -52.70721475
## age         0.22021302
## gender1     -2.48744811
## race2       2.46037359
## race3      -1.08502802
## race4      -0.90753700
## smoking1    2.11825022
## smoking2    2.35768222
## height     0.11928447
## weight     -0.52054391
## bmi        3.42880724
## hypertension1 1.62856332
## diabetes1   -2.15217999
## SBP         0.09818922
## LDL        -0.02968295
## vaccine1    -7.25775052
## severity1    7.44593749
## studyB      5.66750779
```

```
# Make prediction on test data
enet.pred <- predict(enet.fit, newdata = x2)

# Test error
enet.mse <- mean((enet.pred - y2)^2)
enet.mse
```

```
## [1] 353.4649
```

Principal Component Regression(PCR)

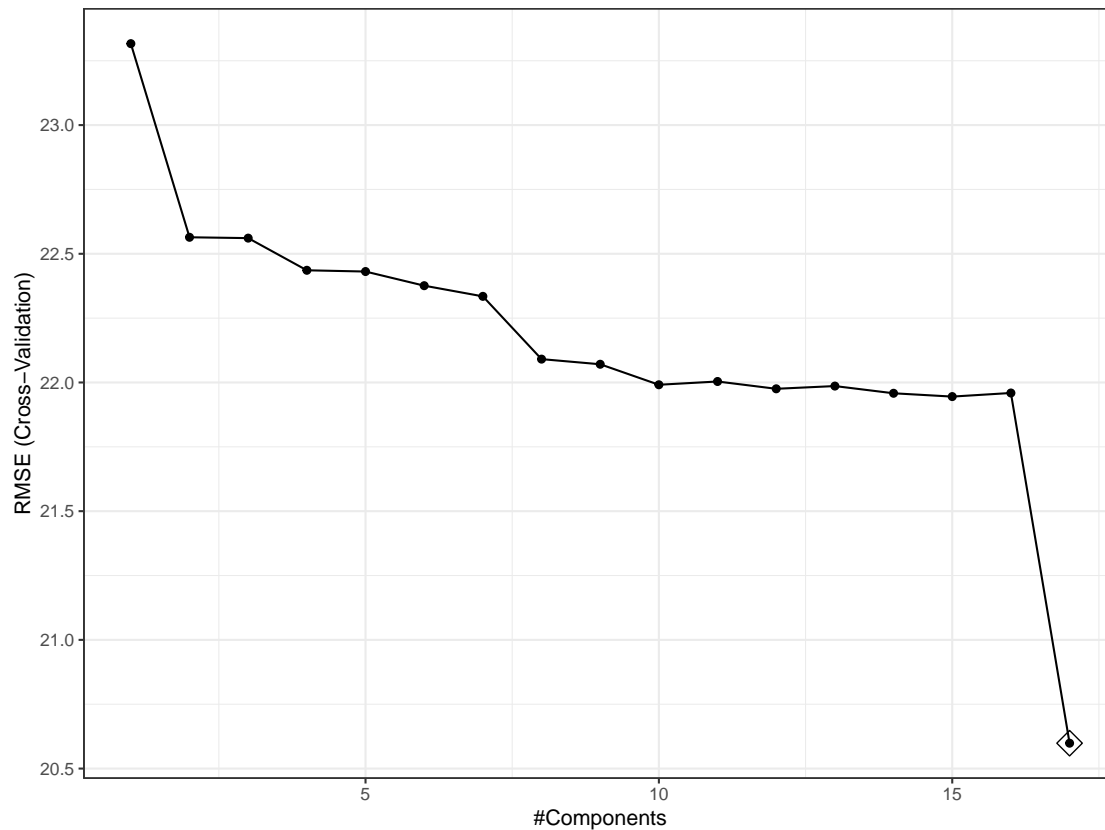
```
# Fitting the model with Cross-validation
set.seed(2358)
pcr.fit <- train(x, y,
                 method = "pcr",
                 tuneGrid = data.frame(ncomp = 1:17),
                 trControl = ctrl1,
                 preProcess = c("center", "scale"))

# Make prediction on test data
pcr.pred <- predict(pcr.fit, newdata = x2)

# Test error
pcr.mse <- mean((pcr.pred - y2)^2)
pcr.mse
```

```
## [1] 314.7791
```

```
# Plot cv RMSE-components
ggplot(pcr.fit, highlight = TRUE) + theme_bw()
```



Partial least squares(PLS)

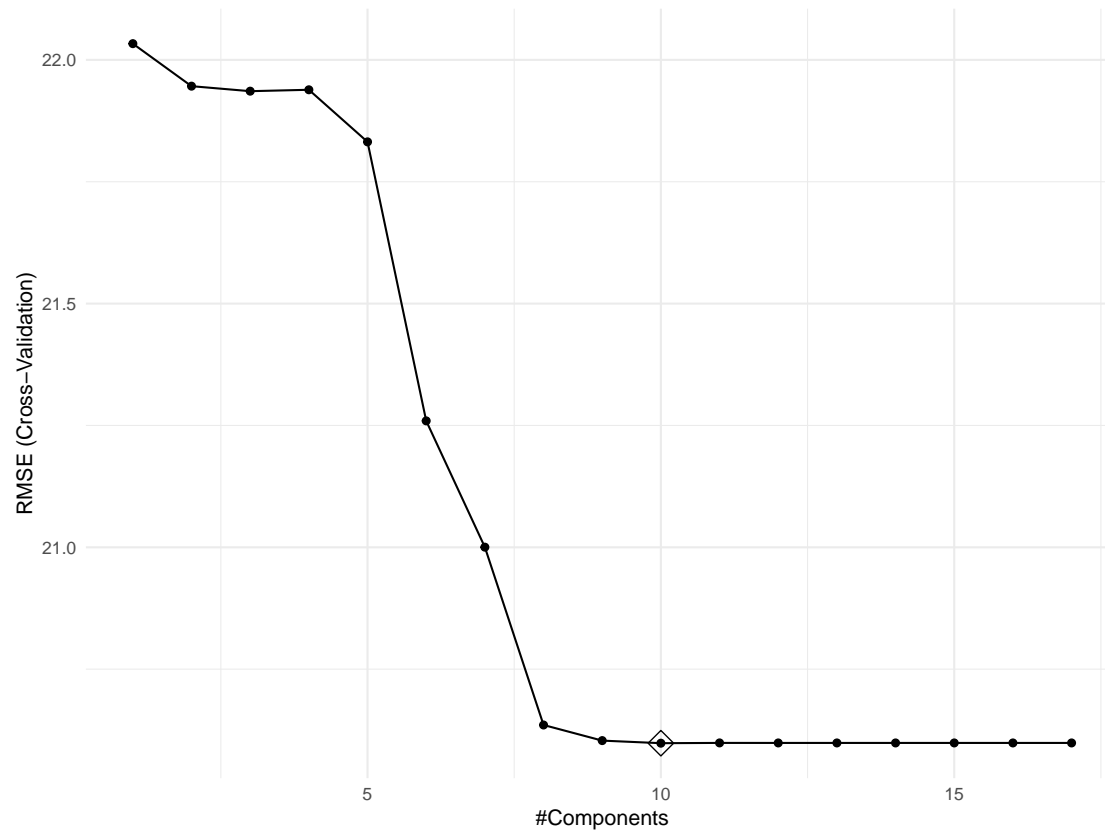
```
# Fitting the model with Cross-validation
set.seed(2358)
pls.fit <- train(x, y,
  method = "pls",
  tuneGrid = data.frame(ncomp = 1:17),
  trControl = ctrl1,
  preProcess = c("center", "scale"))
```

```
# Make prediction on test data
pls.pred <- predict(pls.fit, newdata = x2)
```

```
# Test error
pls.mse <- mean((pls.pred - y2)^2)
pls.mse
```

```
## [1] 314.7983
```

```
# Plot cv RMSE-components
ggplot(pls.fit, highlight = TRUE)
```



Generalized additive model (GAM)

```
# Fit a generalized additive model (GAM) using all the predictors
set.seed(2358)
gam.fit <- train(x, y,
  method = "gam",
  tuneGrid = data.frame(method = "GCV.Cp", select = TRUE),
  trControl = ctrl1)
```

```
gam.fit$bestTune
```

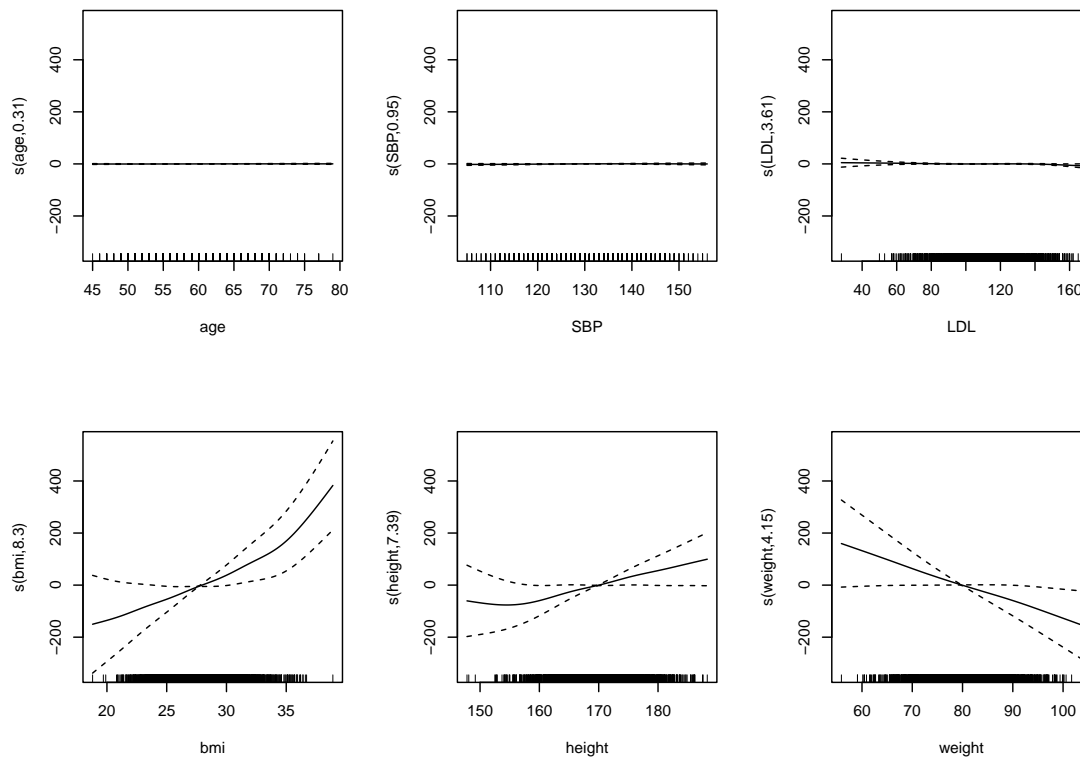
```
## select method
## 1 TRUE GCV.Cp
```

```
gam.fit$finalModel
```

```
##
## Family: gaussian
## Link function: identity
##
## Formula:
## .outcome ~ gender1 + race2 + race3 + race4 + smoking1 + smoking2 +
## hypertension1 + diabetes1 + vaccine1 + severity1 + studyB +
```

```
##      s(age) + s(SBP) + s(LDL) + s(bmi) + s(height) + s(weight)
##
## Estimated degrees of freedom:
## 0.311 0.948 3.615 8.298 7.395 4.146  total = 36.71
##
## GCV score: 380.8098
```

```
# Plot the results
par(mfrow=c(2,3))
plot(gam.fit$finalModel)
```



```
# Make prediction on test data
gam.pred <- predict(gam.fit, newdata = x2)

# test error
gam.test.error <- mean((gam.pred - y2)^2)
gam.test.error
```

```
## [1] 291.3596
```

Multivariate Adaptive Regression Splines (MARS)


```

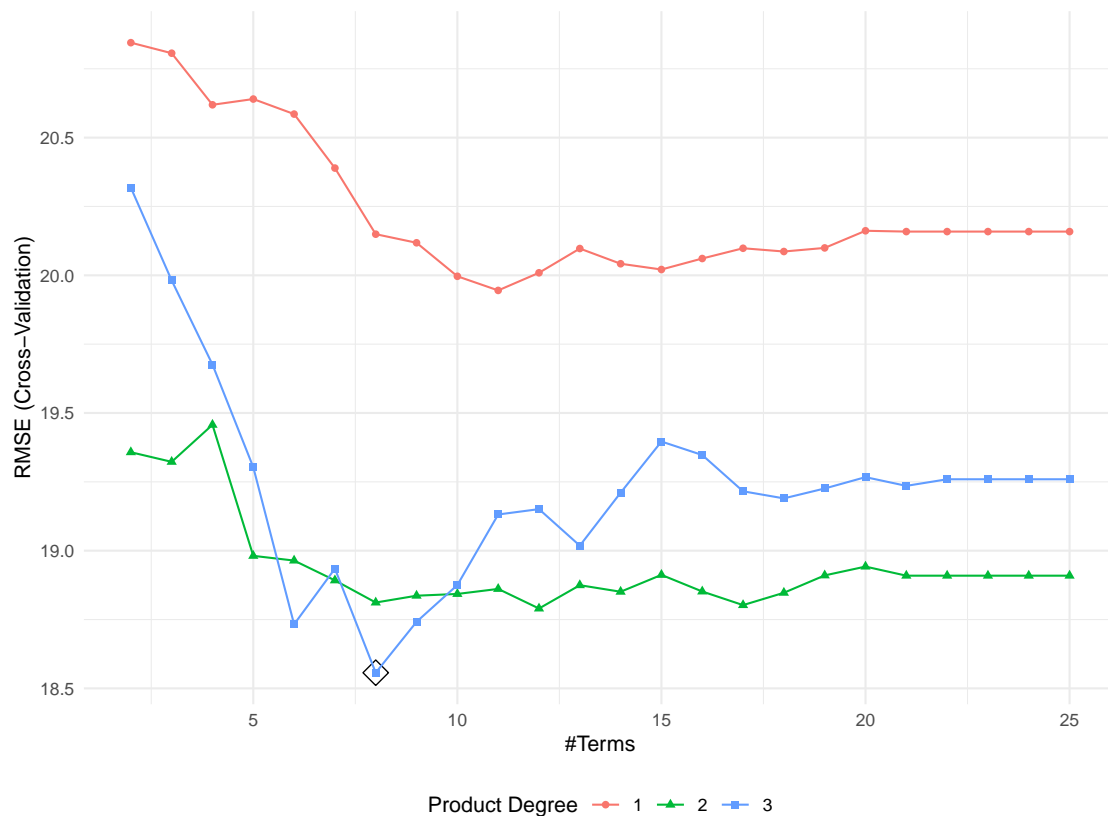
mars_grid <- expand.grid(degree = 1:3,
                        nprune = 2:25)

set.seed(2358)
mars.fit <- train(x, y,
                 method = "earth",
                 tuneGrid = mars_grid,
                 trControl = ctrl1)

```

```
## Loading required package: earth
```

```
ggplot(mars.fit, highlight = T)
```



```
mars.fit$bestTune
```

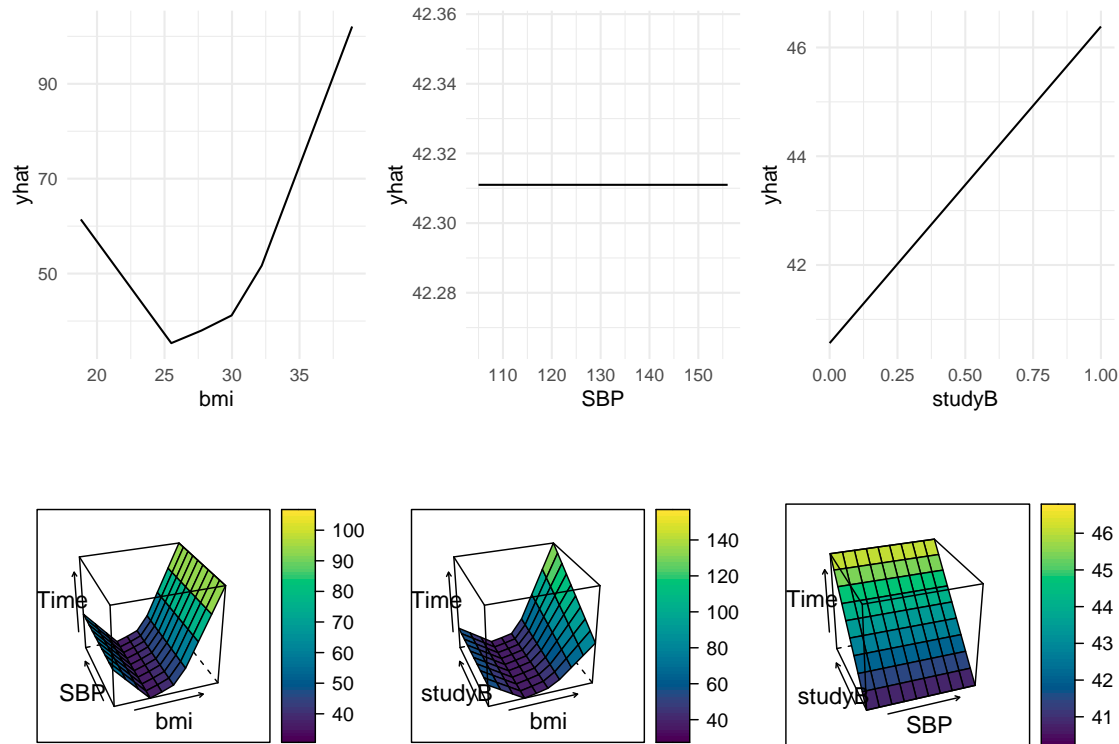
```
##      nprune degree
## 55         8      3
```

```
coef(mars.fit$finalModel)
```

```
##              (Intercept)              h(31-bmi)
##              18.222464              3.892250
##      h(bmi-31) * studyB      h(age-62) * h(bmi-31) * studyB
```

```
##                      32.871468                      26.143670
##                      vaccine1 h(height-161.4) * h(bmi-31) * studyB
##                      -7.204188                      -3.139983
##                      h(bmi-25.6)          h(age-61) * h(bmi-31) * studyB
##                      5.316962                      -17.146877

# Present the partial dependence plot of an arbitrary predictor
p1 <- pdp::partial(mars.fit, pred.var = c("bmi"), grid.resolution = 10) %>% autoplot()
p2 <- pdp::partial(mars.fit, pred.var = c("SBP"), grid.resolution = 10) %>% autoplot()
p3 <- pdp::partial(mars.fit, pred.var = c("studyB"), grid.resolution = 10) %>% autoplot()
p4 <- pdp::partial(mars.fit, pred.var = c("bmi", "SBP"),
  grid.resolution = 10) %>%
  pdp::plotPartial(levelplot = FALSE, zlab = "Time", drape = TRUE,
    screen = list(z = 20, x = -60))
p5 <- pdp::partial(mars.fit, pred.var = c("bmi", "studyB"),
  grid.resolution = 10) %>%
  pdp::plotPartial(levelplot = FALSE, zlab = "Time", drape = TRUE,
    screen = list(z = 20, x = -60))
p6 <- pdp::partial(mars.fit, pred.var = c("SBP", "studyB"),
  grid.resolution = 10) %>%
  pdp::plotPartial(levelplot = FALSE, zlab = "Time", drape = TRUE,
    screen = list(z = 20, x = -60))
gridExtra::grid.arrange(p1, p2,p3, p4, p5, p6, ncol = 3)
```



```
# Make prediction on test data
mars.pred <- predict(mars.fit, newdata = x2)

# test error
mars.test.error <- mean((mars.pred - y2)^2)
mars.test.error

## [1] 405.4464
```

Model Comparison

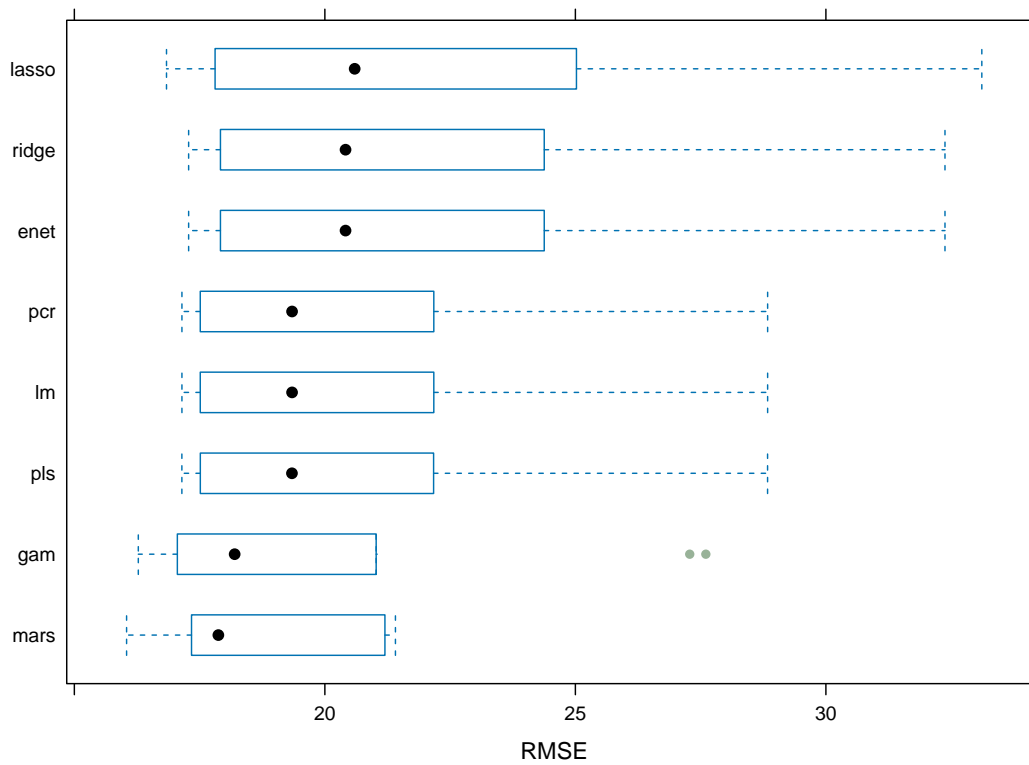
```
# re-samples
set.seed(2358)
resamp <- resamples(list(lm = lm.fit,
                        lasso = lasso.fit,
                        ridge = ridge.fit,
                        enet = enet.fit,
                        pcr = pcr.fit,
                        pls = pls.fit,
                        gam = gam.fit,
                        mars = mars.fit))

summary(resamp)
```

```
##
```

```
## Call:
## summary.resamples(object = resamp)
##
## Models: lm, lasso, ridge, enet, pcr, pls, gam, mars
## Number of resamples: 10
##
## MAE
##           Min.   1st Qu.   Median     Mean   3rd Qu.     Max. NA's
## lm      12.56665 13.01623 13.50877 13.55186 13.86917 14.80595    0
## lasso   12.31484 12.67224 12.92787 13.42197 13.90163 15.49975    0
## ridge   12.42148 12.90384 12.98603 13.55880 14.05688 15.50008    0
## enet    12.42148 12.90384 12.98603 13.55880 14.05688 15.50008    0
## pcr     12.56665 13.01623 13.50877 13.55186 13.86917 14.80595    0
## pls     12.56717 13.01642 13.50925 13.55162 13.86545 14.80890    0
## gam     11.67916 12.35691 12.82913 12.92176 13.03443 15.00257    0
## mars    11.22777 11.51274 11.78872 12.08433 12.73739 13.29310    0
##
## RMSE
##           Min.   1st Qu.   Median     Mean   3rd Qu.     Max. NA's
## lm      17.14718 17.58935 19.34523 20.59863 21.68019 28.83700    0
## lasso   16.83939 17.89280 20.59503 22.01030 24.12462 33.11216    0
## ridge   17.28100 17.91872 20.41250 21.80126 23.60665 32.37707    0
## enet    17.28100 17.91872 20.41250 21.80126 23.60665 32.37707    0
## pcr     17.14718 17.58935 19.34523 20.59863 21.68019 28.83700    0
## pls     17.14588 17.59106 19.34304 20.59811 21.67704 28.83590    0
## gam     16.27655 17.07963 18.19943 19.99675 20.85941 27.60386    0
## mars    16.04255 17.41683 17.87424 18.55684 20.44732 21.40862    0
##
## Rsquared
##           Min.   1st Qu.   Median     Mean   3rd Qu.     Max. NA's
## lm      0.06848279 0.1670135 0.2323908 0.2405915 0.3099305 0.4833811    0
## lasso   0.02068200 0.1122997 0.1342710 0.1326346 0.1495219 0.2400910    0
## ridge   0.02866402 0.1259909 0.1395051 0.1480371 0.1789634 0.2479223    0
## enet    0.02866402 0.1259909 0.1395051 0.1480371 0.1789634 0.2479223    0
## pcr     0.06848279 0.1670135 0.2323908 0.2405915 0.3099305 0.4833811    0
## pls     0.06853575 0.1669119 0.2323691 0.2406330 0.3101671 0.4835539    0
## gam     0.11995974 0.1673279 0.2158768 0.2909863 0.3792122 0.6079131    0
## mars    0.07183623 0.2437275 0.3843057 0.3954056 0.5643194 0.7308590    0

# RMSE box-plot between models
bwplot(resamp, metric = "RMSE")
```



Based on the summary and plot, we would likely use the Multivariate Adaptive Regression Splines (MARS) model to predict the recovery time from COVID-19 illness because it minimizes the mean RMSE over re-samples.