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Midterm Project Analysis

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Data Wrangling 2

```
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)
```

```
##
                                                               weight
        age
                  gender
                           race
                                   smoking
                                                height
## Min.
         :42.0
                  0:1544
                           1:1967
                                   0:1822
                                            Min. :147.8
                                                           Min. : 55.90
                  1:1456
                           2: 158
                                            1st Qu.:166.0
  1st Qu.:57.0
                                   1: 859
                                                            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
                                            3rd Qu.:173.9
                                                            3rd Qu.: 84.80
##
                                                 :188.6
## Max.
          :79.0
                                            Max.
                                                           Max.
                                                                 :103.70
##
        bmi
                   hypertension diabetes
                                             SBP
                                                            LDL
                                                                       vaccine
                                                        Min. : 28.0
## Min. :18.80
                   0:1508
                               0:2537
                                                                       0:1212
                                        Min.
                                               :105.0
##
   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
                                                             :110.5
                                                        Mean
##
   3rd Qu.:29.50
                                        3rd Qu.:136.0
                                                        3rd Qu.:124.0
                                        Max. :156.0
## Max.
          :38.90
                                                        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	nunique	top_counts
gender	0	1	FALSE	2	0: 1544, 1: 1456
race	0	1	FALSE	4	1: 1967, 3: 604, 4: 271, 2: 158
$\operatorname{smoking}$	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_rat	e 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	
$_{ m bmi}$	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	
${\tt recovery_time}$	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

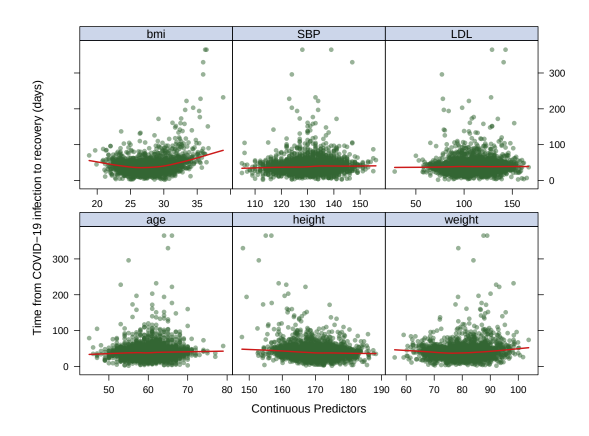
```
# data partition
set.seed(2358)
indexTrain <- createDataPartition(y = dat$recovery_time, p = 0.8, list = FALSE)
train <- dat[indexTrain, ]
test <- dat[-indexTrain, ]

# 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</pre>
```

Expoloratory Data Analysis

```
# Remove all the categorical predictors out from x before plotting
x_continuous <-</pre>
 x[, !(colnames(x) %in%
          c("gender1", "race2", "race3", "race4",
             "smoking1", "smoking2", "hypertension1",
            "diabetes1", "vaccine1", "severity1", "studyB"))]
# For Continuous Predictors
theme1 <- trellis.par.get()</pre>
theme1$plot.symbol$col \leftarrow 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)
featurePlot(x_continuous,
            plot = "scatter",
            span = .5,
            labels = c("Continuous Predictors", "Time from COVID-19 infection to recovery (days)"),
            type = c("p", "smooth"),
            layout = c(3, 2))
```

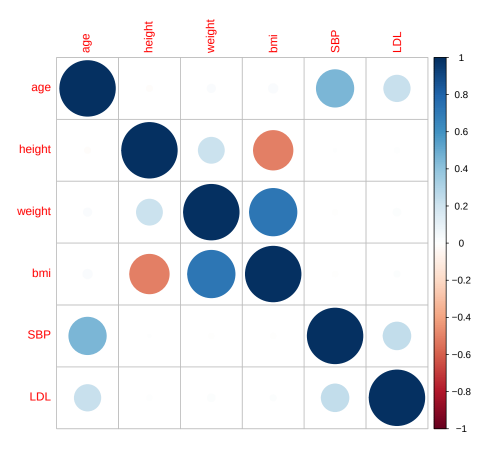


```
# For Categorical Predictors
dis_gender = train %>%
  ggplot(aes(x = gender, y = recovery_time)) +
  geom_violin(fill = "orange", color = "blue", alpha = .5) +
  scale_x_discrete(labels = c('Female','Male'))
dis_race = train %>%
  ggplot(aes(x = race, y = recovery_time)) +
  geom_violin(fill = "orange", 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 +
                                recovery_time
 recovery_time
                                                               recovery_time
     0
                                    0
                                             Asian Black Hispanic
           Female
                       Male
                                       White
                                                                     Never smokedmer smoker
                gender
                                                race
                                                                              smoking
 recovery_time
                                recovery_time
                                                               recovery_time
                                    0
                                                                   0
     0
                                           No
                                                      Yes
                                                                       Not vaccinated Vaccinated
              hypertension
                                               diabetes
                                                                              vaccine
 a000 1000 1000
                                recovery_time
     0
                                    0
                                                       В
                      Severe
          Not severe
                severity
                                                study
```

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

Linear 7



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)</pre>
# Fitting the model with Cross-validation
set.seed(2358)
lm.fit <- train(x, y, method = "lm", trControl = ctrl1)</pre>
# Final model coefficients
coef(lm.fit$finalModel)
##
     (Intercept)
                                      gender1
                                                       race2
                            age
##
  -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
                                               9.762877e-02 -3.373236e-02
##
    4.285697e+01
                  1.752483e+00 -2.040537e+00
        vaccine1
                     severity1
                                       studyB
## -7.247882e+00 7.938166e+00 5.629688e+00
```

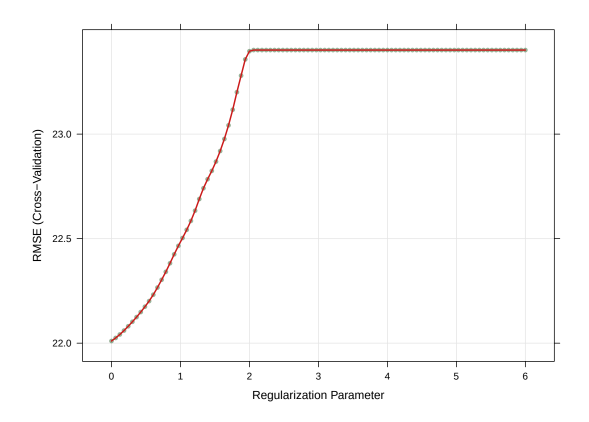
Lasso 8

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

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

[1] 314.7791

Lasso



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

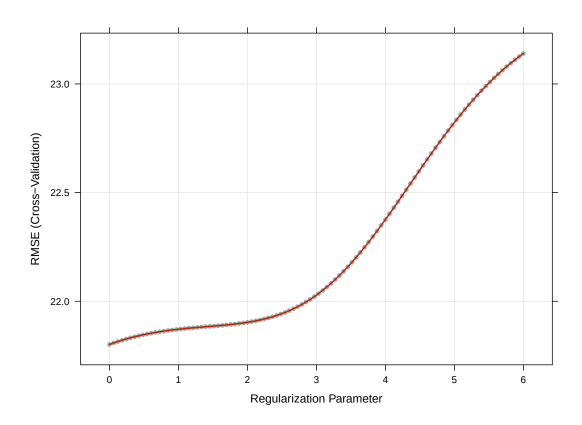
Ridge 9

```
## alpha lambda
## 1
       1
# coefficients in the final model
coef(lasso.fit$finalModel, lasso.fit$bestTune$lambda)
## 18 x 1 sparse Matrix of class "dgCMatrix"
## (Intercept)
               14.07004988
                 0.07075850
## age
## 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)</pre>
# Test error
lasso.mse <- mean((lasso.pred - y2)^2)</pre>
lasso.mse
```

[1] 352.0247

Ridge

Ridge 10



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
```

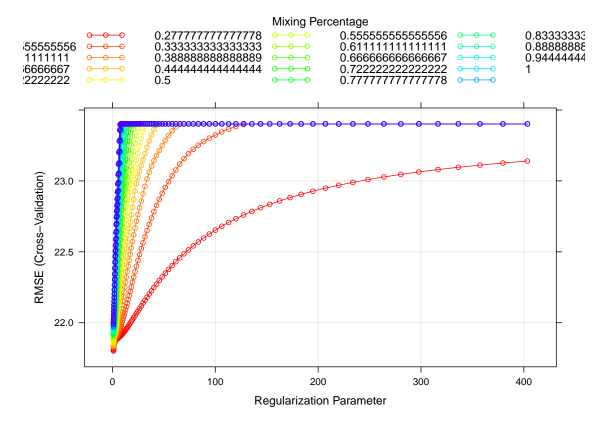
Elastic net 11

```
## LDL
                  -0.02968295
## vaccine1
                  -7.25775052
## severity1
                  7.44593749
## studyB
                   5.66750779
# Make prediction on test data
ridge.pred <- predict(ridge.fit, newdata = x2)</pre>
# Test error
ridge.mse <- mean((ridge.pred - y2)^2)</pre>
ridge.mse
## [1] 353.4649
Elastic net
# Fitting the model with Cross-validation
set.seed(2358)
enet.fit <- train(x, y,method = "glmnet",</pre>
                  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
# 25 kinds of Plot colors
myCol<- rainbow(25)</pre>
myPar <- list(superpose.symbol = list(col = myCol),</pre>
superpose.line = list(col = myCol))
```

Plot CV RMSE-alpha&lambda

plot(enet.fit, par.settings = myPar)

Elastic net 12



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

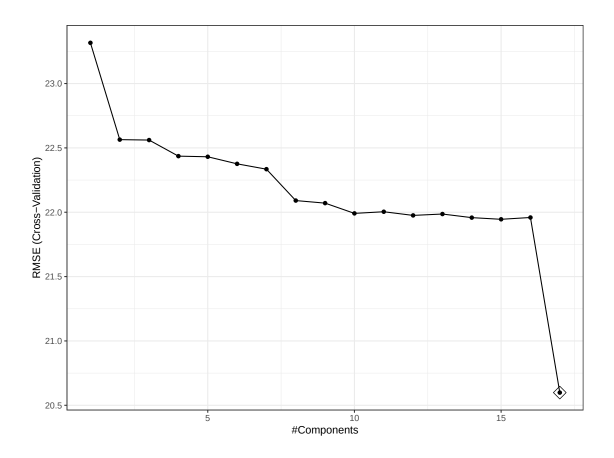
```
## 18 x 1 sparse Matrix of class "dgCMatrix"
##
## (Intercept)
                 -52.70721475
## age
                    0.22021302
## gender1
                   -2.48744811
                    2.46037359
## race2
## race3
                  -1.08502802
## race4
                  -0.90753700
## smoking1
                   2.11825022
## smoking2
                   2.35768222
## height
                   0.11928447
## weight
                   -0.52054391
                   3.42880724
## bmi
## 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</pre>
```

[1] 353.4649

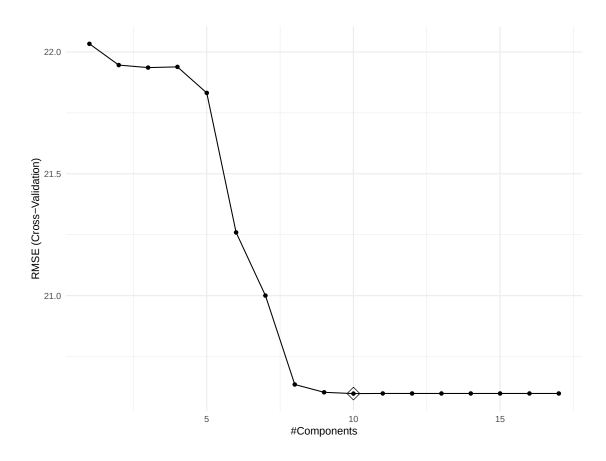
Principal Component Regression(PCR)



Partial least squares(PLS)

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

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



Generalized additive model (GAM)

```
{\it \# Fit \ a \ generalized \ additive \ model \ (\textit{GAM}) \ using \ all \ the \ predictors}
set.seed(2358)
gam.fit <- train(x, y,</pre>
                  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)
    400
                                                                               400
                                          400
s(age,0.31)
                                      s(SBP,0.95)
                                                                           s(LDL,3.61)
                                          200
                                                                               200
    200
                                         0
                                                                               0
    0
    -200
                                          -200
                                                                               -200
                                             Լուդասարտարաարապատուրու
        <u> Լրուդուդուդուդուդուդ</u>
                                               110 120
       45 50 55 60 65 70 75 80
                                                        130 140 150
                                                                                     40 60 80
                                                                                                   120
                                                                                                          160
                                                         SBP
                                                                                               LDL
                   age
    400
                                          400
                                                                               400
                                                                            s(weight, 4.15)
                                      s(height,7.39)
                                          200
    200
                                                                               200
s(bmi, 8.3)
                                                                               0
                                          0
                                          -200
                                                                               -200
    -200
         20
               25
                     30
                           35
                                                    160
                                                          170
                                                                180
                                                                                          70
                                                                                               80
                                                                                                    90
                   bmi
                                                        height
                                                                                              weight
# Make prediction on test data
gam.pred <- predict(gam.fit, newdata = x2)</pre>
```

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

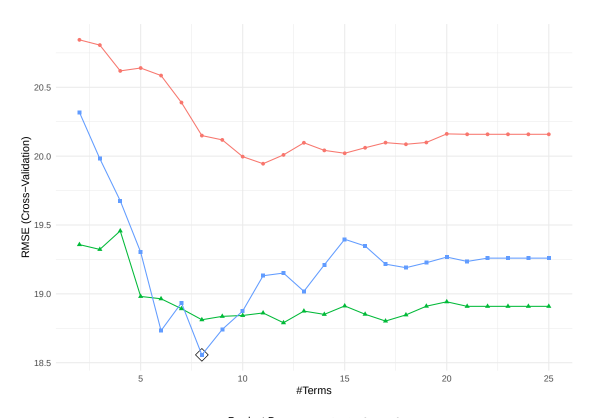
# test error
gam.test.error <- mean((gam.pred - y2)^2)
gam.test.error</pre>
```

[1] 291.3596

Multivariate Adaptive Regression Splines (MARS)

Loading required package: earth

ggplot(mars.fit, highlight = T)



Product Degree → 1 → 2 → 3

```
mars.fit$bestTune
```

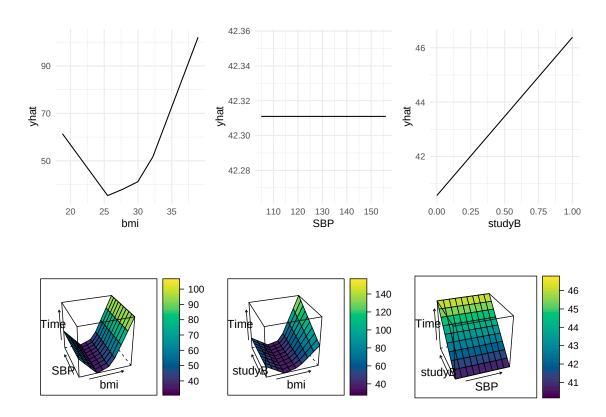
```
## nprune degree
## 55 8 3
```

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"),</pre>
                   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)
```

Model Comparison 19



```
# 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</pre>
```

[1] 405.4464

Model Comparison

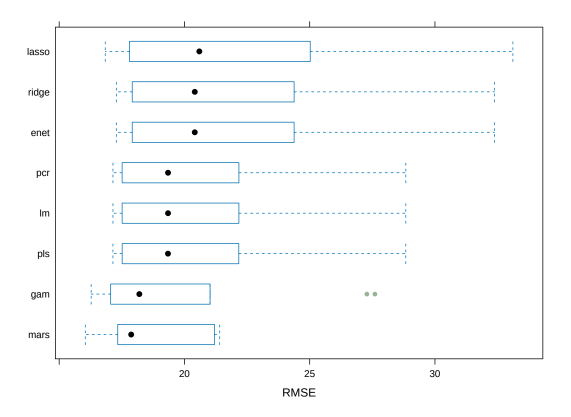
##

Model Comparison 20

```
## 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.
## lm
         12.56665 13.01623 13.50877 13.55186 13.86917 14.80595
## lasso 12.31484 12.67224 12.92787 13.42197 13.90163 15.49975
## ridge 12.42148 12.90384 12.98603 13.55880 14.05688 15.50008
## enet 12.42148 12.90384 12.98603 13.55880 14.05688 15.50008
                                                                  0
         12.56665 13.01623 13.50877 13.55186 13.86917 14.80595
                                                                  0
## pcr
         12.56717 13.01642 13.50925 13.55162 13.86545 14.80890
## pls
         11.67916 12.35691 12.82913 12.92176 13.03443 15.00257
## gam
                                                                  0
## mars 11.22777 11.51274 11.78872 12.08433 12.73739 13.29310
##
## RMSE
##
             Min. 1st Qu.
                             Median
                                        Mean 3rd Qu.
## lm
         17.14718 17.58935 19.34523 20.59863 21.68019 28.83700
## lasso 16.83939 17.89280 20.59503 22.01030 24.12462 33.11216
## ridge 17.28100 17.91872 20.41250 21.80126 23.60665 32.37707
## enet 17.28100 17.91872 20.41250 21.80126 23.60665 32.37707
         17.14718 17.58935 19.34523 20.59863 21.68019 28.83700
## pcr
         17.14588 17.59106 19.34304 20.59811 21.67704 28.83590
## pls
## gam
         16.27655 17.07963 18.19943 19.99675 20.85941 27.60386
## mars 16.04255 17.41683 17.87424 18.55684 20.44732 21.40862
##
## Rsquared
##
                      1st Qu.
                                 Median
                                                    3rd Qu.
               Min.
                                             Mean
## lm
         0.06848279 0.1670135 0.2323908 0.2405915 0.3099305 0.4833811
## 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
         0.06848279 0.1670135 0.2323908 0.2405915 0.3099305 0.4833811
                                                                         0
## pcr
         0.06853575\ 0.1669119\ 0.2323691\ 0.2406330\ 0.3101671\ 0.4835539
                                                                         0
## pls
         0.11995974 0.1673279 0.2158768 0.2909863 0.3792122 0.6079131
## mars 0.07183623 0.2437275 0.3843057 0.3954056 0.5643194 0.7308590
# RMSE box-plot between models
```

bwplot(resamp, metric = "RMSE")

Model Comparison 21



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 resamples.