Analysis

Arthur Starodynov

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Load in the data and explore the data.

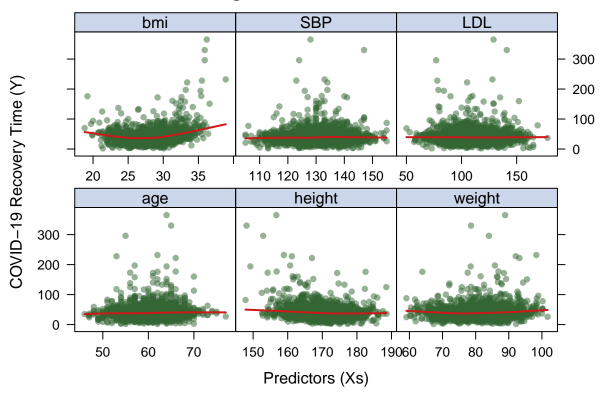
```
load("data/recovery.Rdata")
head(dat)
##
     id age gender race smoking height weight bmi hypertension diabetes SBP LDL
                              2 170.2
                                         78.7 27.2
## 1 1 56
                 0
                                                               0
                                                                        0 120 97
                      1
        70
## 2 2
                                         73.1 25.4
                                 169.6
                                                               1
                                                                        0 134 112
                 1
                      1
## 3 3 57
                                         77.4 27.3
                                                              1
                 1
                      1
                              0 168.4
                                                                        0 131 88
                                                              0
## 4 4 53
                 0
                    1
                              0 166.7
                                         76.1 27.4
                                                                        0 115 87
## 5 5 59
                      1
                              2 173.6
                                         70.2 23.3
                                                              0
                                                                        0 127 118
                 1
## 6 6 60
                 1
                      3
                                         75.1 28.4
                                                              0
                                                                        0 129 104
                              1 162.8
     vaccine severity study recovery_time
## 1
           0
                    0
                          Α
## 2
           0
                    0
                                       44
## 3
           1
                    0
                                       29
## 4
           0
                    1
                          Α
                                       47
## 5
                    0
                          Α
                                       40
## 6
                    0
                                       34
set.seed(2024)
#library(summarytools)
#st_options(plain.ascii = FALSE,
           #style = "rmarkdown",
           #dfSummary.silent = TRUE,
           #footnote = NA,
           #subtitle.emphasis = FALSE)
#dfSummary(data[,-1])
data <- dat %>%
 select(-id)
                # removing the id variable from the data
set.seed(2024)
tRows <- createDataPartition(dat$recovery_time, p = 0.7, list = FALSE)
# training data
data_train <- data[tRows, ]</pre>
x <- model.matrix(recovery_time~.,data)[tRows,-1]</pre>
y <- data$recovery_time[tRows]</pre>
#Test data
```

```
data_test <- data[-tRows, ]
x2 <- model.matrix(recovery_time~.,data)[-tRows,-1]
y2 <- data$recovery_time[-tRows]</pre>
```

Exploring the data set:

```
data_train_viz <- data_train %>%
  mutate(study = case_when( # turn study (character variable) into a numeric variable
    study == "A" ~ 1,
    study == "B" ~ 2,
    study == "C" ~ 3))
non_numeric_cols <- sapply(data_train_viz, function(x) !is.numeric(x))</pre>
# Convert non-numeric columns to numeric
data_train_viz[, non_numeric_cols] <- lapply(data_train_viz[, non_numeric_cols], as.numeric)</pre>
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)
# for continous variables:
featurePlot(x = data_train_viz %>% select(age, height, weight, bmi, SBP, LDL),
            y = data_train_viz[ ,15],
            plot = "scatter",
            span = .5,
            labels = c("Predictors (Xs)", "COVID-19 Recovery Time (Y)"),
            main = "Figure 1. Lattice Plot",
            type = c("p", "smooth"))
```

Figure 1. Lattice Plot



«««< HEAD

======»»»> 8d60aef84a87065beb8632d98d5e0094c4d03360 # Training models

Training Various models to see which will perform the best.

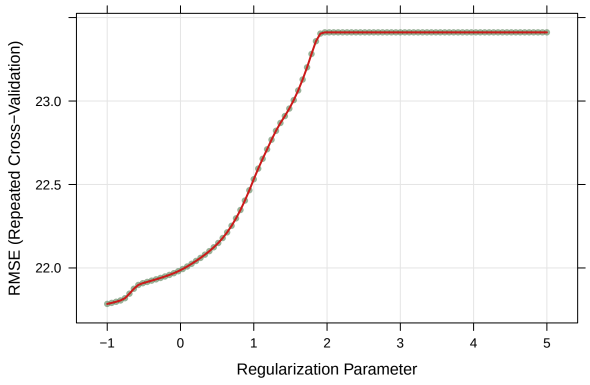
LGM:

```
set.seed(2024)
ctrl <- trainControl(method = "repeatedcv", number = 10, repeats = 5) # Using the best rule
linear_model <- train(recovery_time ~ .,</pre>
               data = data_train,
               method = "lm",
               trControl = ctrl)
summary(linear_model)
##
## Call:
## lm(formula = .outcome ~ ., data = dat)
##
## Residuals:
##
       Min
                1Q Median
                                3Q
                                       Max
  -57.256 -11.308
                   -0.031
                             8.772 248.452
##
##
## Coefficients:
                  Estimate Std. Error t value Pr(>|t|)
## (Intercept) -2.327e+03 1.235e+02 -18.849 < 2e-16 ***
                 2.290e-01 1.131e-01
                                        2.026 0.042941 *
## age
## gender
                -3.220e+00 8.879e-01 -3.627 0.000294 ***
## race2
                 4.179e+00 2.097e+00
                                        1.992 0.046451 *
```

```
## race3
              -7.288e-01 1.132e+00 -0.644 0.519609
## race4
              -4.931e-01 1.562e+00 -0.316 0.752287
                                     2.393 0.016815 *
## smoking1
              2.409e+00 1.007e+00
               3.094e+00 1.470e+00
                                      2.105 0.035408 *
## smoking2
## height
                1.360e+01 7.263e-01 18.721 < 2e-16 ***
## weight
               -1.482e+01 7.659e-01 -19.356 < 2e-16 ***
## bmi
               4.430e+01 2.199e+00 20.147 < 2e-16 ***
## hypertension 1.917e+00 1.457e+00
                                     1.316 0.188388
## diabetes
             -2.118e+00 1.220e+00 -1.737 0.082585 .
## SBP
               6.747e-02 9.462e-02
                                    0.713 0.475889
## LDL
               -6.237e-02 2.354e-02 -2.650 0.008120 **
               -6.935e+00 9.061e-01 -7.654 2.96e-14 ***
## vaccine
               8.689e+00 1.424e+00
                                     6.102 1.24e-09 ***
## severity
## studyB
               5.139e+00 9.427e-01 5.452 5.58e-08 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 20.29 on 2084 degrees of freedom
## Multiple R-squared: 0.2817, Adjusted R-squared: 0.2758
## F-statistic: 48.07 on 17 and 2084 DF, p-value: < 2.2e-16
Finding the RMSE:
## [1] 19.85811
```

We can see that the RMSE is 19.858 for the Generalized linear model.

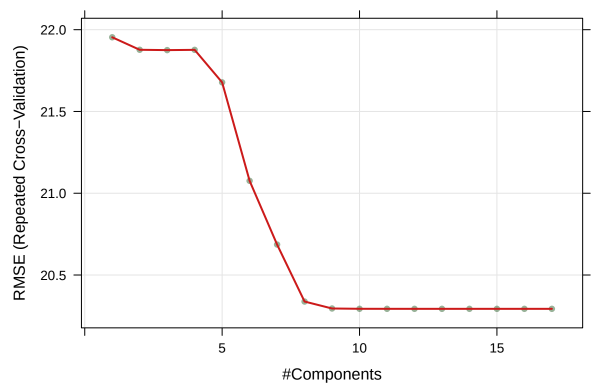
Lasso Model



```
tuning_param <- lasso_model$bestTune
coef(lasso_model$finalModel, lasso_model$bestTune$lambda)</pre>
```

```
## 18 x 1 sparse Matrix of class "dgCMatrix"
## (Intercept)
                -25.43825107
                   0.19265246
## age
                  -1.93903073
## gender
                   3.10474378
## race2
## race3
## race4
                   0.94047609
## smoking1
## smoking2
                   1.41274621
## height
## weight
                  -0.40581611
## bmi
                   2.90669760
## hypertension
                   0.23950557
## diabetes
                  -1.66338038
## SBP
                   0.10053663
## LDL
                  -0.03119877
## vaccine
                  -6.45367070
## severity
                   7.96723577
## studyB
                   5.11509535
Finding RMSE of the Lasso model:
lasso_pred <- predict(lasso_model, newdata = x2)</pre>
lasso_rmse <- sqrt(mean((lasso_pred - data_test$recovery_time)^2))</pre>
lasso_rmse
```

PLS model

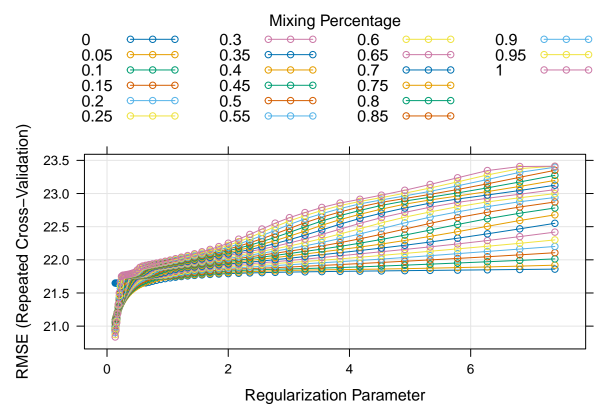


Finding the RMSE

```
pls_pred <- predict(pls_model, newdata = x2)
pls_rmse <- sqrt(mean((pls_pred - data_test$recovery_time)^2))
pls_rmse</pre>
```

[1] 19.85812

Elastic net model



Finding the RMSE

```
enet_pred <- predict(enet_model, newdata = x2)
enet_rmse <- sqrt(mean((enet_pred - data_test$recovery_time)^2))
enet_rmse</pre>
```

[1] 19.95906

MARS model

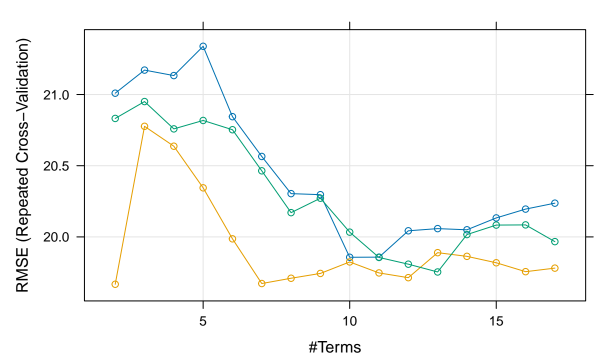
Resampling the data by encoding dummy variables to be able to use a MARS model.

```
data_test_mars <- data_mars[-tRows, ]
## matrix of predictors
mars_x2 <- model.matrix(recovery_time~.,data_mars)[-tRows,-1]
## vector of response
mars_y2 <- data_mars$recovery_time[-tRows]</pre>
```

Using newly created data to train a MARS model.

Product Degree





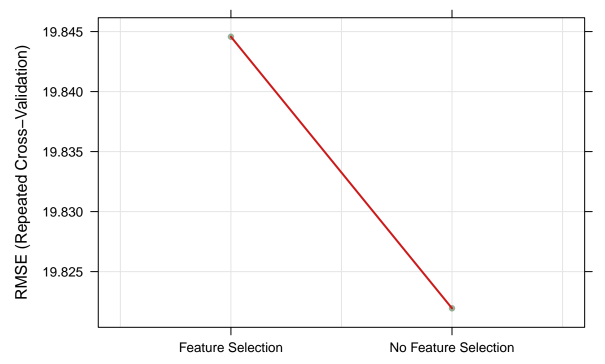
Find the RMSE

```
mars_pred <- predict(mars_model, newdata = mars_x2)
mars_rmse <- sqrt(mean((mars_pred - data_test_mars$recovery_time)^2))
mars_rmse</pre>
```

[1] 18.05067

GAM Model

```
set.seed(2024)
```



Feature Selection

summary(GAM_model\$finalModel)

```
##
## Family: gaussian
## Link function: identity
##
## Formula:
  .outcome ~ gender + race3 + race4 + smoking1 + smoking2 + hypertension +
      diabetes + vaccine + severity + studyB + s(age) + s(SBP) +
##
##
      s(LDL) + s(bmi) + s(height) + s(weight)
##
## Parametric coefficients:
               Estimate Std. Error t value Pr(>|t|)
##
                            1.1752 37.560 < 2e-16 ***
## (Intercept)
                44.1392
## gender
                -3.5459
                            0.8381 -4.231 2.43e-05 ***
## race3
                -0.6689
                            1.0599 -0.631 0.52803
## race4
                -1.0546
                            1.4686 -0.718 0.47276
## smoking1
                 2.6229
                            0.9504
                                    2.760 0.00583 **
## smoking2
                 3.6774
                            1.3880
                                    2.649 0.00813 **
## hypertension
                1.9348
                            1.4111
                                    1.371 0.17048
## diabetes
                -2.0844
                            1.1518 -1.810 0.07050 .
## vaccine
                -7.0273
                            0.8545 -8.224 3.43e-16 ***
                            1.3413 6.974 4.13e-12 ***
## severity
                 9.3538
```

```
## studyB
                 4.6053
                            0.8908
                                   5.170 2.57e-07 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Approximate significance of smooth terms:
              edf Ref.df
##
                              F p-value
            1.000 1.000 1.677 0.195457
## s(age)
            1.478 1.813 0.786 0.543422
## s(SBP)
## s(LDL)
            1.219
                   1.408 3.834 0.027315 *
## s(bmi)
            7.199 8.167 57.524 < 2e-16 ***
## s(height) 6.908 7.956 9.740 < 2e-16 ***
## s(weight) 1.670 2.194 6.925 0.000453 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## R-sq.(adj) = 0.358
                        Deviance explained = 36.7%
## GCV = 370.41 Scale est. = 365.04
Finding the RMSE:
GAM_pred <- predict(GAM_model, newdata = x2)</pre>
GAM_rmse <- sqrt(mean((GAM_pred - data_test$recovery_time)^2))</pre>
GAM_rmse
```

Model Selection

[1] 18.48635

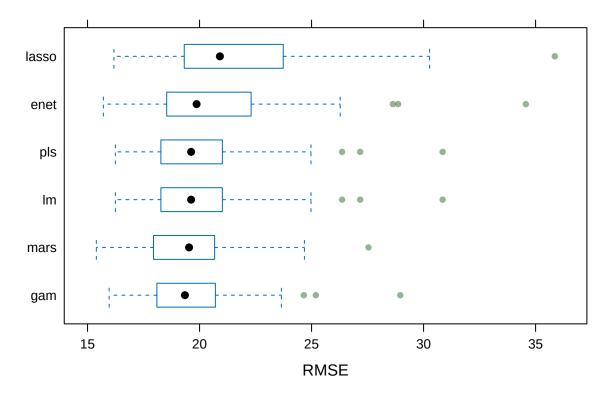
For choosing the best model, we assessed the RMSE and checked which model had the lowest RMSE. This indicates that the model is the best performing due to a low prediction error. Models with high RMSE has high prediction error meaning they are worse models.

```
resample_data <- resamples(list(
  lm = linear_model,
  lasso = lasso_model,
  enet = enet_model,
  pls = pls_model,
  gam = GAM_model,
  mars = mars_model
))</pre>
```

```
##
## Call:
## summary.resamples(object = resample_data)
##
## Models: lm, lasso, enet, pls, gam, mars
## Number of resamples: 50
##
## MAE
## MAE
## Min. 1st Qu. Median Mean 3rd Qu. Max. NA's
## lm 12.01315 13.18428 13.42575 13.55674 13.93071 15.25681 0
```

```
## lasso 11.68470 13.14360 13.58178 13.65305 14.00670 16.14233
         11.64707 12.77613 13.24593 13.30991 13.69025 15.49942
         12.01308 13.18429 13.42575 13.55674 13.93070 15.25680
         11.37981 12.28795 12.91906 12.99219 13.49834 14.52340
                                                                   0
  gam
##
         11.46416 12.40770 13.04918 13.02956 13.47370 15.26110
##
## RMSE
##
             Min.
                   1st Qu.
                             Median
                                        Mean 3rd Qu.
## lm
         16.24262 18.34594 19.62327 20.29317 21.01705 30.85055
  lasso 16.17754 19.31831 20.90705 21.78457 23.72520 35.85732
        15.70800 18.53689 19.86654 20.83649 22.29719 34.56517
         16.24261 18.34592 19.62328 20.29317 21.01705 30.85054
                                                                   0
  pls
   gam
##
         15.96439 18.12753 19.34727 19.82194 20.70164 28.95957
                                                                   0
         15.39199 18.03713 19.53205 19.66747 20.66927 27.54021
##
## Rsquared
##
                 Min.
                        1st Qu.
                                   Median
                                                       3rd Qu.
                                                Mean
## lm
         8.905049e-02 0.1802640 0.2591439 0.2617221 0.3401823 0.4626231
## lasso 1.426456e-02 0.1158527 0.1460068 0.1482118 0.1843823 0.2711260
        7.122077e-02 0.1692736 0.2205285 0.2234573 0.2806290 0.3810741
## pls
         8.905086e-02 0.1802631 0.2591458 0.2617223 0.3401832 0.4626238
         9.561505e-02 0.1879885 0.2928082 0.3091493 0.4603588 0.5413315
## gam
        4.678139e-05 0.1377927 0.2789584 0.3029493 0.4813938 0.6480672
bwplot(resample_data,
       metric = "RMSE",
       main = "Figure 2. Model Comparison Plot Using RMSE")
```

Figure 2. Model Comparison Plot Using RMSE



Results

##

```
summary(mars_model$finalModel)
## Call: earth(x=matrix[2102,18], y=c(31,47,40,34,3...), keepxy=TRUE, degree=2,
              nprune=2)
##
##
                     coefficients
## (Intercept)
                         40.14471
## studyB * h(bmi-31)
                         30.05710
##
## Selected 2 of 26 terms, and 2 of 18 predictors (nprune=2)
## Termination condition: Reached nk 37
## Importance: studyB, bmi, gender-unused, race1-unused, race2-unused, ...
## Number of terms at each degree of interaction: 1 0 1
## GCV 383.1087
                  RSS 802615
                                GRSq 0.3266962
                                                  RSq 0.3282976
summary(GAM_model$finalModel)
##
## Family: gaussian
## Link function: identity
##
## Formula:
  .outcome ~ gender + race3 + race4 + smoking1 + smoking2 + hypertension +
##
##
      diabetes + vaccine + severity + studyB + s(age) + s(SBP) +
      s(LDL) + s(bmi) + s(height) + s(weight)
##
##
## Parametric coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                44.1392
                           1.1752 37.560 < 2e-16 ***
## gender
                -3.5459
                            0.8381 -4.231 2.43e-05 ***
## race3
                -0.6689
                            1.0599
                                   -0.631 0.52803
## race4
                -1.0546
                            1.4686 -0.718 0.47276
## smoking1
                 2.6229
                            0.9504
                                    2.760 0.00583 **
                                    2.649 0.00813 **
## smoking2
                 3.6774
                            1.3880
## hypertension
                1.9348
                            1.4111
                                     1.371 0.17048
## diabetes
                -2.0844
                            1.1518 -1.810 0.07050 .
## vaccine
                -7.0273
                            0.8545 -8.224 3.43e-16 ***
## severity
                 9.3538
                            1.3413
                                    6.974 4.13e-12 ***
                 4.6053
                            0.8908
                                    5.170 2.57e-07 ***
## studyB
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Approximate significance of smooth terms:
##
              edf Ref.df
                              F p-value
## s(age)
            1.000 1.000 1.677 0.195457
## s(SBP)
            1.478 1.813 0.786 0.543422
## s(LDL)
            1.219 1.408 3.834 0.027315 *
## s(bmi)
            7.199 8.167 57.524 < 2e-16 ***
## s(height) 6.908 7.956 9.740 < 2e-16 ***
## s(weight) 1.670 2.194 6.925 0.000453 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

```
## R-sq.(adj) = 0.358 Deviance explained = 36.7%
## GCV = 370.41 Scale est. = 365.04 n = 2102
```

Although we see through the RMSE comparison that the mean and median RMSE of the mars model was the smallest when we compare the number of predictors used for the mars model was 2, we know that this would not be a good and accurate model for future use. Hence, we want to use the GAM model for any further comparison.

All of these factors together explain 44.2% of the deviance in COVID-19 recovery time. Additionally, the model's training error (RMSE using the

Appendix / GAM plots

```
gam.m1 <- gam(recovery_time ~ gender + race + smoking + hypertension +
    diabetes + vaccine + severity + study + s(age) + s(SBP) +
    s(LDL) + s(bmi) + s(height) + s(weight), data = data_train)

plot(gam.m1)</pre>
```

