# Linear Models

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To gain a better understanding of the factors that predict recovery time from COVID-19 illness, a study was designed to combine three existing cohort studies that have been tracking participants for several years. The study collects recovery information through questionnaires and medical records, and leverages existing data on personal characteristics prior to the pandemic.

In this project, we predict the recovery time based on important risk factors. The training data is in "training\_df", and the test data is in "training\_df". The response is in the column "Time to recovery (tt\_recovery\_time)", and other variables can be used as predictors. The variable definitions can be found in "dictionary.txt".

First, we import the data and adjust the variable type. As we want to compare different models afterwards, we use caret.

```
# data splitting
data.split = initial_split(dat, prop = 0.8)
training_data = training(data.split)
testing_data = testing(data.split)

# check missing values
training_data |> is.na() |> sum()
```

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

```
testing_data |> is.na() |> sum()
```

#### ## [1] 0

```
# summary of training and testing data
training_data |> summary()
```

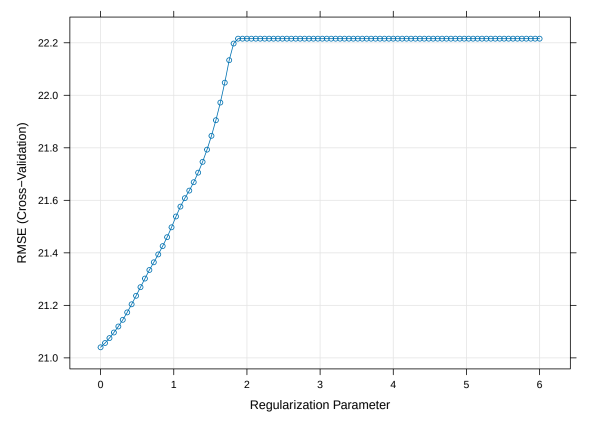
```
##
                                                  height
                                                                  weight
                   gender
                            race
                                     smoking
        age
          :42.00
                   0:1235
                                     0:1442
                                                                     : 57.80
##
  \mathtt{Min}.
                            1:1569
                                              Min.
                                                    :147.8
                                                              Min.
   1st Qu.:57.00
                                                              1st Qu.: 75.30
                   1:1165
                            2: 134
                                     1: 700
                                              1st Qu.:166.1
  Median :60.00
                                              Median :170.0
                            3: 475
                                     2: 258
                                                              Median: 79.80
   Mean :60.13
                            4: 222
                                              Mean :170.0
                                                              Mean
                                                                     : 80.01
##
   3rd Qu.:63.00
                                              3rd Qu.:174.0
                                                              3rd Qu.: 84.83
   Max.
         :79.00
                                                    :188.6
                                                              Max. :103.70
                                              Max.
                                              SBP
                                                              LDL
##
        bmi
                   hypertension diabetes
                                                                         vaccine
                                                         Min.
##
   Min.
          :18.80
                   0:1223
                                0:2032
                                         Min.
                                                :105.0
                                                                : 28.0
                                                                         0: 961
##
   1st Qu.:25.80
                                1: 368
                                                         1st Qu.: 98.0
                                                                         1:1439
                   1:1177
                                         1st Qu.:125.0
  Median :27.60
                                         Median :130.0
                                                         Median :111.0
## Mean :27.76
                                         Mean
                                                :130.3
                                                         Mean
                                                               :110.8
   3rd Qu.:29.52
                                         3rd Qu.:136.0
                                                         3rd Qu.:125.0
## Max.
          :36.70
                                                :156.0
                                         Max.
                                                         Max. :171.0
  severity
               study
                               recovery_time
##
   0:2145
            Length:2400
                               Min. : 2.00
##
   1: 255
                               1st Qu.: 30.00
            Class : character
##
            Mode :character
                               Median: 39.00
##
                               Mean : 42.03
                               3rd Qu.: 49.00
##
##
                               Max. :365.00
testing_data |> summary()
##
                                               height
                                                               weight
                   gender race
                                   smoking
         age
   Min.
         :47.00
                   0:309
                           1:398
                                   0:380
                                                                  : 55.90
                                           Min.
                                                  :148.1
                                                           Min.
   1st Qu.:57.00
                   1:291
                           2: 24
                                   1:159
                                           1st Qu.:165.8
                                                           1st Qu.: 75.10
  Median :60.00
                           3:129
                                   2: 61
                                           Median :169.7
                                                           Median: 79.55
##
  Mean
         :60.47
                           4: 49
                                           Mean
                                                  :169.6
                                                           Mean
                                                                 : 79.75
   3rd Qu.:64.00
                                           3rd Qu.:173.7
                                                           3rd Qu.: 84.50
##
   Max. :74.00
                                           Max.
                                                  :186.1
                                                           Max.
                                                                  :100.00
        bmi
                                              SBP
                                                            LDL
##
                   hypertension diabetes
                                                                       vaccine
##
  Min.
          :19.20
                   0:285
                                0:505
                                         Min.
                                                :105
                                                       Min.
                                                              : 57.0
                                                                       0:251
   1st Qu.:25.90
                   1:315
                                1: 95
                                         1st Qu.:126
                                                       1st Qu.: 95.0
                                                                      1:349
                                         Median:131
## Median :27.80
                                                       Median :108.0
## Mean
          :27.78
                                         Mean
                                                :131
                                                       Mean :108.9
## 3rd Qu.:29.50
                                                       3rd Qu.:123.0
                                         3rd Qu.:136
          :38.90
## Max.
                                         Max.
                                                :154
                                                       Max. :178.0
##
   severity
                               recovery_time
               study
##
  0:534
            Length:600
                               Min. : 2.00
  1: 66
##
            Class : character
                               1st Qu.: 31.00
##
                               Median: 39.00
            Mode :character
##
                               Mean : 42.73
##
                               3rd Qu.: 49.00
##
                               Max.
                                      :330.00
# training data
train.x = model.matrix(recovery_time ~ ., training_data)[, -1]
train.y = training_data$recovery_time
# test data
test.x = model.matrix(recovery_time ~ ., testing_data)[, -1]
```

test.y = testing\_data\$recovery\_time

```
# cross validation
ctrl = trainControl(method = "cv", number = 10)
```

There is no missing data in both datasets. The training dataset has 2400 observation and 16 variables, and the test dataset has 600 samples and 17 variables.

# Lasso Regression Model

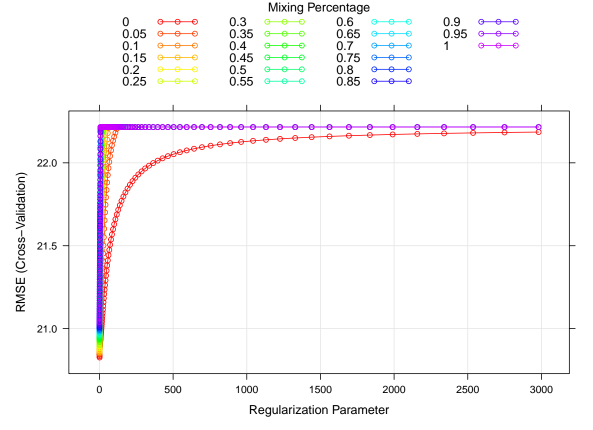


The flattening in the curve occurs because the lasso regression model has reached a stable solution where no more coefficients become zero as the regularization parameter increases. At this point, further increasing the regularization parameter does not change the set of non-zero coefficients or their values, resulting in a flat region where the model's performance remains constant. This behavior indicates that the model has achieved the optimal level of sparsity, with the regularization parameter value at the start of the flat region corresponding to the desired sparse solution.

```
lasso.pred = predict(lasso.fit, newdata = testing_data)
mse.lasso = mean((test.y - lasso.pred) ^ 2)
```

The selected best tuning parameter is  $\lambda = 1$   $\alpha = 1$ , and the test error (MSE) is 562.263383.

#### Elastic Net Model



The vertical line pattern in the graph is a characteristic behavior of the elastic net model. It occurs because the elastic net combines lasso and ridge regularization, which can lead to sparse solutions with some coefficients becoming exactly zero. As the regularization parameter increases, the model transitions between different sparse solutions, causing sudden drops or jumps in the cross-validation error curve.

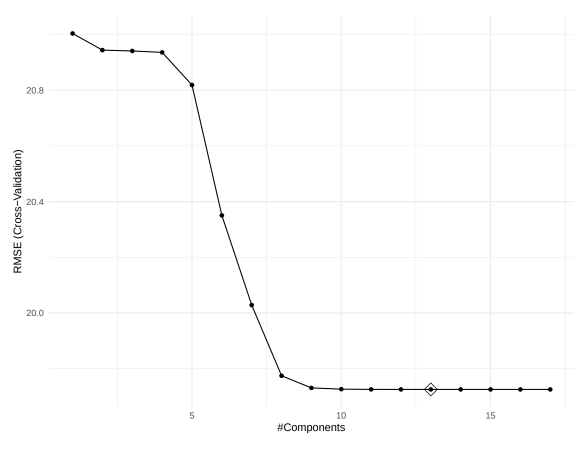
```
enet.pred = predict(enet.fit, newdata = testing_data)
mse.enet = mean((test.y - enet.pred) ^ 2)
```

The selected tuning parameter is  $\alpha = 0$ ,  $\lambda = 1$ , and the test error (MSE) is 536.2385884

# Partial Least Square

```
set.seed(2716)

pls.fit = train(train.x, train.y, method = "pls",
   tuneGrid = data.frame(ncomp = 1:17),
   trControl = ctrl, preProcess = c("center", "scale"))
ggplot(pls.fit, highlight = T)
```



```
pls.pred = predict(pls.fit, newdata = test.x)
mse.pls = mean((test.y - pls.pred) ^ 2)
```

As illustrated in the plot, 11 components are included in my model, and the test error (MSE) is 464.8643889. The coefficients are as follows:

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

```
## , , 13 comps
##
##
                     .outcome
                   1.04306479
## age
## gender1
                  -1.79478232
## race2
                   0.67651655
## race3
                   0.09607768
## race4
                  -0.14876310
## smoking1
                  0.98529363
## smoking2
                  1.02530399
## height
                  74.39909797
## weight
                 -96.19301259
## bmi
                 112.57056811
## hypertension1
                 0.68131672
## diabetes1
                  -0.28468608
## SBP
                   0.46391355
## LDL
                  -0.95602747
## vaccine1
                  -2.97123020
## severity1
                  1.62625094
## studyB
                   2.02187127
```

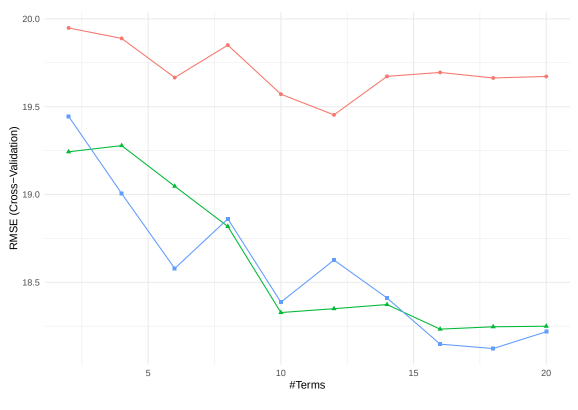
### **MARS**

Now, train a multivariate adaptive regression spline (MARS) model to predict the response variable.

Since there are two tuning parameters associated with the MARS model: the degree of interactions and the number of retained terms, we need to perform a grid search to identify the optimal combination of these hyperparameters that minimize prediction error.

```
ctrl = trainControl(method = "cv", number = 10)
mars.grid = expand.grid(degree = 1 : 3, nprune = seq(2, 20, by = 2))
set.seed(2716)
mars.fit = train(train.x, train.y, method = "earth", tuneGrid = mars.grid, trControl = ctrl)
## Loading required package: earth
## Loading required package: Formula
## Loading required package: plotmo
## Loading required package: plotrix
##
## Attaching package: 'plotrix'
## The following object is masked from 'package:scales':
##
## rescale
## Loading required package: TeachingDemos
```

### ggplot(mars.fit)



Product Degree - 1 - 2 - 3

### summary(mars.fit\$finalModel)

```
## Call: earth(x=matrix[2400,17], y=c(69,9,7,43,36,...), keepxy=TRUE, degree=3,
##
               nprune=18)
##
##
                                          coefficients
## (Intercept)
                                             17.425888
## gender1
                                             -3.838226
## vaccine1
                                             -5.523433
## severity1
                                              5.146386
## h(bmi-25.7)
                                              6.207509
## h(31.1-bmi)
                                              4.065175
## h(bmi-35.4)
                                             27.540176
## h(bmi-31.1) * studyB
                                           -305.070403
## h(bmi-25.7) * h(140-SBP)
                                             -0.064119
## smoking1 * h(bmi-31.1) * studyB
                                             15.366973
## smoking2 * h(bmi-31.1) * studyB
                                             28.176098
## h(age-61) * h(bmi-31.1) * studyB
                                             5.249927
## h(weight-78.7) * h(bmi-31.1) * studyB
                                             23.765436
## h(89.3-weight) * h(bmi-31.1) * studyB
                                             24.335697
## h(weight-89.3) * h(bmi-31.1) * studyB
                                            -25.433893
## h(bmi-31.1) * h(LDL-94) * studyB
                                             -5.448260
## h(bmi-31.1) * h(94-LDL) * studyB
                                             5.533714
## h(bmi-31.1) * h(LDL-84) * studyB
                                             5.716112
```

```
##
## Selected 18 of 25 terms, and 11 of 17 predictors (nprune=18)
## Termination condition: Reached nk 35
## Importance: bmi, studyB, LDL, age, vaccine1, weight, smoking1, gender1, ...
## Number of terms at each degree of interaction: 1 6 2 9
## GCV 261.8665 RSS 605903.8 GRSq 0.4893553 RSq 0.5072879

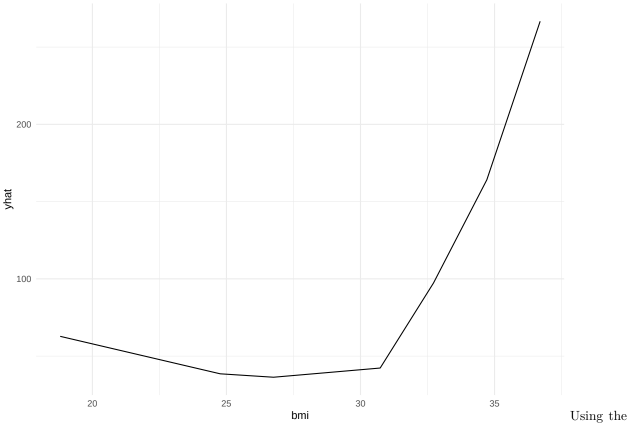
## Coefficient of the MARS model
coef(mars.fit$finalModel)
```

```
##
                              (Intercept)
                                                                      h(31.1-bmi)
##
                              17.42588791
                                                                       4.06517510
                    h(bmi-31.1) * studyB
##
                                                h(age-61) * h(bmi-31.1) * studyB
##
                            -305.07040296
                                                                       5.24992718
##
                              h(bmi-25.7)
                                                                         vaccine1
##
                               6.20750882
                                                                      -5.52343301
##
        h(bmi-31.1) * h(LDL-94) * studyB
                                                h(bmi-31.1) * h(94-LDL) * studyB
##
                              -5.44825972
                                                                       5.53371358
   h(weight-89.3) * h(bmi-31.1) * studyB h(89.3-weight) * h(bmi-31.1) * studyB
##
                             -25.43389302
                                                                      24.33569724
##
        h(bmi-31.1) * h(LDL-84) * studyB
                                                 smoking1 * h(bmi-31.1) * studyB
                               5.71611207
##
                                                                      15.36697296
##
                                  gender1
                                                                        severity1
##
                              -3.83822562
                                                                       5.14638556
         smoking2 * h(bmi-31.1) * studyB
                                                        h(bmi-25.7) * h(140-SBP)
##
##
                              28.17609759
                                                                      -0.06411943
  h(weight-78.7) * h(bmi-31.1) * studyB
                                                                      h(bmi-35.4)
                              23.76543599
                                                                      27.54017572
```

The MARS model selects 12 of 27 terms, and 8 of 17 predictors. The most important variables are bmi (Body Mass Index; BMI = weight (in kilograms) / height (in meters) squared) and studyB(The study (A/B) that the participant belongs to).

To better understand the relationship between these features and outstate, we can create partial dependence plots (PDPs) for each feature individually and also an interaction PDP. To simplify, here we only present the PDP for number of full-time undergraduates sbp.

```
pdp::partial(mars.fit, pred.var = c("bmi"), grid.resolution = 10) |> autoplot()
```



final model, we can predict on the test data.

```
pred.mars = predict(mars.fit, newdata = test.x)
mse.mars = mean((pred.mars - test.y) ^ 2)
```

The test error measured by MSE using the final MARS model is 392.3426264

## **GAM**

```
set.seed(2716)
gam.fit = train(train.x, train.y, method = "gam", trControl = ctrl)

## Loading required package: mgcv

## Loading required package: nlme

## ## Attaching package: 'nlme'

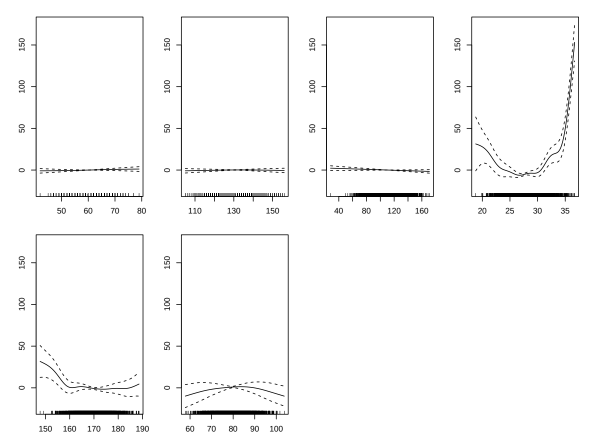
## The following object is masked from 'package:dplyr':

## collapse

## This is mgcv 1.9-0. For overview type 'help("mgcv-package")'.
```

### summary(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)
##
## Parametric coefficients:
               Estimate Std. Error t value Pr(>|t|)
                         1.0022 43.437 < 2e-16 ***
## (Intercept)
                43.5319
## gender1
               -3.9286
                          0.7579 -5.184 2.36e-07 ***
## race2
                         1.6763 1.485 0.13761
                2.4897
## race3
                0.2184
                          0.9730 0.224 0.82239
## race4
               -1.0791
                           1.3330 -0.810 0.41831
## smoking1
                ## smoking2
                ## hypertension1 2.2691
                         1.0394 2.183 0.02913 *
## diabetes1
               -0.9912
                          1.0520 -0.942 0.34617
                         0.7773 -8.051 1.29e-15 ***
## vaccine1
               -6.2577
## severity1
               5.6267
                           1.2306 4.572 5.07e-06 ***
                           0.8026 5.197 2.20e-07 ***
## studyB
                4.1707
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Approximate significance of smooth terms:
##
              edf Ref.df
                             F p-value
           0.6260
                   9 0.153 0.115489
## s(age)
## s(SBP)
          0.5946
                      9 0.079 0.293041
                      9 0.277 0.068772 .
## s(LDL)
           0.7589
## s(bmi)
           8.8974
                      9 67.421 < 2e-16 ***
## s(height) 6.6451
                      9 3.220 2.26e-05 ***
                    9 1.614 0.000227 ***
## s(weight) 2.3895
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## R-sq.(adj) = 0.334 Deviance explained = 34.3\%
## GCV = 345.74 Scale est. = 341.14 n = 2400
par(mar = c(2, 2, 2, 2), mfrow = c(2, 4))
plot(gam.fit$finalModel)
```



It could be observed that certain variables (age, sbp, ldl) have no relationship with the recovery\_time, bmi, and height both have a positive relationship with recovery\_time.

Using the final model, we can predict on the test data.

```
pred.gam = predict(gam.fit, newdata = test.x)
mse.gam = mean((pred.gam - test.y) ^ 2)
```

The test error measured by MSE using the final GAM is  $454.3607584. \,$ 

# **Model Comparison**

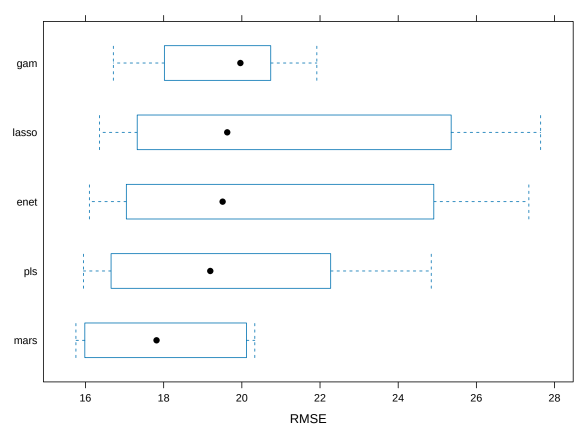
Here, we compare the CV results of different models and choose the model with the smallest median RMSE.

```
##
## Call:
## summary.resamples(object = resamp)
##
```

```
## Models: lasso, gam, enet, pls, mars
## Number of resamples: 10
##
## MAE
            Min. 1st Qu.
                            Median
                                        Mean 3rd Qu.
                                                          Max. NA's
## lasso 11.91749 12.22876 12.78822 13.10307 13.97511 14.75607
        11.71387 12.30531 12.64584 12.60154 13.04721 13.17282
## enet 11.82640 12.24000 12.86273 13.14786 14.09859 14.70406
## pls
        12.00552 12.42079 13.07887 13.06909 13.69203 14.12036
                                                                  0
## mars 10.80694 11.46341 11.88141 11.87063 12.24314 12.74929
## RMSE
##
             Min. 1st Qu.
                             Median
                                        Mean 3rd Qu.
                                                          Max. NA's
## lasso 16.36102 17.35644 19.62606 21.04028 24.59723 27.64478
        16.71229 18.02794 19.96280 19.42835 20.62654 21.91883
## enet 16.09922 17.14833 19.50831 20.82568 24.22164 27.34263
                                                                  0
        15.94665 16.77630 19.19268 19.72570 22.05070 24.84530
                                                                  0
## pls
## mars 15.75520 16.33340 17.81756 18.12274 20.06809 20.33183
##
## Rsquared
##
              Min.
                       1st Qu.
                                  Median
                                              Mean
                                                     3rd Qu.
                                                                  Max. NA's
## lasso 0.07035909 0.08459142 0.1095733 0.1133160 0.1199068 0.1790213
        0.12563865\ 0.16945853\ 0.2130396\ 0.2981010\ 0.4564618\ 0.5962197
## gam
                                                                          0
## enet 0.08321485 0.10886399 0.1216388 0.1284815 0.1374788 0.1954873
                                                                          0
        0.10933283 0.16901995 0.2338977 0.2315719 0.2969252 0.3415088
                                                                          0
## pls
## mars 0.07128380 0.22502531 0.2658819 0.3406511 0.4887175 0.7078823
```

Using bw-plot to compare their RMSE.

```
bwplot(resamp, metric = "RMSE")
```



Hence, we selected MARS model as it has smallest RMSE.