P8106_midterm_project_v1

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```
library(pacman)
p_load(tidyverse, caret, tidymodels, corrplot, ggplot2, plotmo, ggrepel, patchwork, earth, pdp, mgcv, k
## package 'cli' successfully unpacked and MD5 sums checked

## package 'purrr' successfully unpacked and MD5 sums checked

## package 'rlang' successfully unpacked and MD5 sums checked

## package 'tidymodels' successfully unpacked and MD5 sums checked

## The downloaded binary packages are in

## C:\Users\Persica\AppData\Local\Temp\Rtmpw1Sqj7\downloaded_packages
```

1. Data Processing

```
load("dat1.RData")
load("dat2.RData")
skimr::skim(dat1)
```

Table 1: Data summary

Name	dat1
Number of rows	5000
Number of columns	14
Column type frequency:	
factor	2
numeric	12
Group variables	 None
Group variables	None

Variable type: factor

skim_variable	n_missing	$complete_rate$	ordered	n_unique	top_counts
race	0	1	FALSE	4	1: 3221, 3: 1036, 4: 465, 2: 278
smoking	0	1	FALSE	3	0: 3010, 1: 1504, 2: 486

Variable type: numeric

skim_variablen_	_missing complet	e_r	ate mean	sd	p0	p25	p50	p75	p100	hist
id	0	1	2500.50	1443.52	1.00	1250.75	2500.50	3750.25	5000.00	
age	0	1	59.97	4.50	44.00	57.00	60.00	63.00	75.00	
gender	0	1	0.49	0.50	0.00	0.00	0.00	1.00	1.00	
height	0	1	170.13	5.94	150.20	166.10	170.10	174.22	192.90	
weight	0	1	80.11	7.06	56.70	75.40	80.10	84.90	106.00	
bmi	0	1	27.74	2.76	18.20	25.80	27.60	29.50	38.80	
diabetes	0	1	0.15	0.36	0.00	0.00	0.00	0.00	1.00	
hypertension	0	1	0.46	0.50	0.00	0.00	0.00	1.00	1.00	
SBP	0	1	129.90	8.00	101.00	124.00	130.00	135.00	155.00	
LDL	0	1	109.91	20.15	43.00	96.00	110.00	124.00	185.00	
time	0	1	108.86	43.42	30.00	76.00	106.00	138.00	270.00	
\log _antibody	0	1	10.06	0.60	7.77	9.68	10.09	10.48	11.96	

skimr::skim(dat2)

Table 4: Data summary

Name	dat2
Number of rows	1000
Number of columns	14
Column type frequency:	
factor	2
numeric	12
Group variables	None

Variable type: factor

skim_variable	n_missing	$complete_rate$	ordered	n_unique	top_counts
race smoking	0 0	_	FALSE FALSE		1: 663, 3: 199, 4: 83, 2: 55 0: 601, 1: 296, 2: 103

Variable type: numeric

skim_variablen	_missing comp	lete_rat	te mean	sd	p0	p25	p50	p75	p100	hist
id	0	1	5500.50	288.82	5001.00	5250.75	5500.50	5750.25	6000.00	
age	0	1	60.02	4.45	46.00	57.00	60.00	63.00	75.00	
gender	0	1	0.49	0.50	0.00	0.00	0.00	1.00	1.00	
height	0	1	170.22	6.02	149.40	166.10	170.20	174.20	190.60	

skim_variablen_	_missing comp	olete_ra	te mean	sd	p0	p25	p50	p75	p100	hist
weight	0	1	80.13	7.05	58.80	75.30	80.20	84.40	101.60	
bmi	0	1	27.72	2.82	19.80	25.80	27.60	29.60	35.80	
diabetes	0	1	0.16	0.36	0.00	0.00	0.00	0.00	1.00	
hypertension	0	1	0.46	0.50	0.00	0.00	0.00	1.00	1.00	
SBP	0	1	129.61	8.20	106.00	124.00	130.00	135.00	156.00	
LDL	0	1	110.25	20.32	46.00	96.00	112.00	124.00	174.00	
time	0	1	173.77	46.78	61.00	140.00	171.00	205.00	330.00	
\log _antibody	0	1	9.90	0.59	8.05	9.50	9.93	10.31	11.85	

There is no missing value in this dataset.

```
colnames(dat1)
   [1] "id"
                        "age"
                                        "gender"
                                                        "race"
                                                                        "smoking"
    [6] "height"
                        "weight"
                                        "bmi"
                                                        "diabetes"
                                                                        "hypertension"
## [11] "SBP"
                        "LDL"
                                        "time"
                                                        "log_antibody"
# Clean variable names
dat1 = dat1 |> janitor::clean_names()
dat2 = dat2 |> janitor::clean_names()
```

We can see there is no NA values in the dataset.

```
# Convert variables to factors
dat1$gender <- factor(dat1$gender, levels = c(0, 1), labels = c("Female", "Male"))</pre>
dat1$race <- factor(dat1$race, levels = c(1, 2, 3, 4), labels = c("White", "Asian", "Black", "Hispanic"
dat1$smoking <- factor(dat1$smoking, levels = c(0, 1, 2), labels = c("Never", "Former", "Current"))
dat1$diabetes <- factor(dat1$diabetes, levels = c(0, 1), labels = c("No", "Yes"))</pre>
dat1$hypertension <- factor(dat1$hypertension, levels = c(0, 1), labels = c("No", "Yes"))</pre>
dat2$gender <- factor(dat2$gender, levels = c(0, 1), labels = c("Female", "Male"))</pre>
dat2$race <- factor(dat2$race, levels = c(1, 2, 3, 4), labels = c("White", "Asian", "Black", "Hispanic"</pre>
dat2$smoking <- factor(dat2$smoking, levels = c(0, 1, 2), labels = c("Never", "Former", "Current"))
dat2$diabetes <- factor(dat2$diabetes, levels = c(0, 1), labels = c("No", "Yes"))</pre>
dat2$hypertension <- factor(dat2$hypertension, levels = c(0, 1), labels = c("No", "Yes"))</pre>
# Summary statistics
summary_stats <- summary(dat1)</pre>
print(summary_stats)
          id
                         age
                                        gender
                                                          race
                                                                       smoking
```

```
##
## Min.
         :
                  Min.
                         :44.00
                                  Female:2573
                                                        :3221
                                                               Never :3010
              1
                                                White
  1st Qu.:1251
                  1st Qu.:57.00
                                  Male :2427
                                                Asian
                                                        : 278
                                                               Former: 1504
                                                               Current: 486
## Median :2500
                  Median :60.00
                                                        :1036
                                                Black
           :2500
                         :59.97
                                                Hispanic: 465
## Mean
                  Mean
                  3rd Qu.:63.00
## 3rd Qu.:3750
## Max.
          :5000
                  Max.
                         :75.00
##
       height
                       weight
                                                    diabetes
                                                               hypertension
                                         bmi
          :150.2
                  Min. : 56.70
                                    Min.
                                           :18.20
                                                               No :2702
## Min.
                                                    No :4228
```

```
1st Qu.: 75.40
## 1st Qu.:166.1
                                     1st Qu.:25.80
                                                     Yes: 772
                                                               Yes:2298
##
  Median :170.1
                   Median : 80.10
                                    Median :27.60
  Mean :170.1
                   Mean : 80.11
                                    Mean
                                          :27.74
   3rd Qu.:174.2
                   3rd Qu.: 84.90
                                     3rd Qu.:29.50
##
##
   Max.
          :192.9
                   Max.
                         :106.00
                                    Max.
                                            :38.80
##
                        ldl
        sbp
                                        time
                                                     log_antibody
          :101.0
                          : 43.0
                                          : 30.0
                                                    Min.
                                                          : 7.765
   Min.
                   Min.
                                   Min.
   1st Qu.:124.0
                   1st Qu.: 96.0
                                   1st Qu.: 76.0
                                                    1st Qu.: 9.682
##
## Median :130.0
                   Median :110.0
                                   Median :106.0
                                                    Median :10.089
## Mean
         :129.9
                   Mean
                         :109.9
                                   Mean
                                         :108.9
                                                    Mean
                                                          :10.064
## 3rd Qu.:135.0
                   3rd Qu.:124.0
                                    3rd Qu.:138.0
                                                    3rd Qu.:10.478
## Max. :155.0
                   Max.
                          :185.0
                                   Max.
                                         :270.0
                                                    Max.
                                                          :11.961
# Check for missing values
missing_values <- colSums(is.na(dat1))</pre>
print(missing_values)
##
                                                                          height
            id
                         age
                                   gender
                                                  race
                                                            smoking
##
             0
                                                     0
                          0
                                       0
                                                                 0
                                                                              0
##
         weight
                         bmi
                                 diabetes hypertension
                                                                sbp
                                                                             ldl
##
             0
                           0
                                       0
                                                                  0
                                                                               0
```

There is no missing value in the dataset.

0

##

##

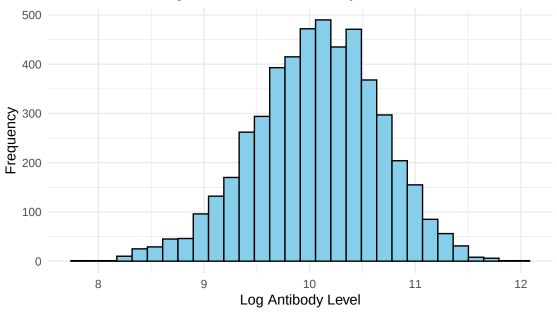
2. Exploratory Data Analysis (EDA)

time log_antibody

0

2.1 Distribution of log antibody



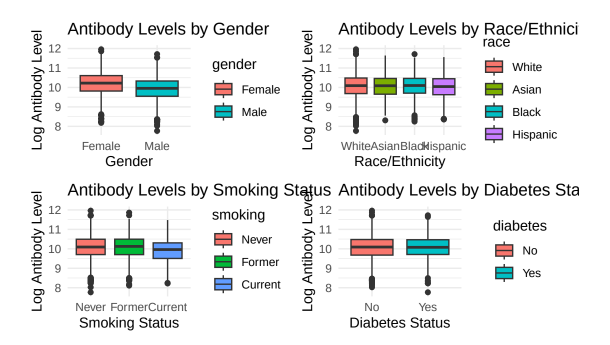


2.2 Boxplot of log antibody

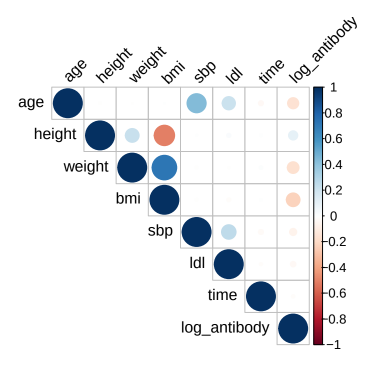
```
# Boxplot of log_antibody by gender
p2 <- ggplot(dat1, aes(x = gender, y = log_antibody, fill = gender)) +
  geom_boxplot() +
  labs(title = "Antibody Levels by Gender",
       x = "Gender",
      y = "Log Antibody Level") +
 theme_minimal()
# Boxplot of log_antibody by race
p3 <- ggplot(dat1, aes(x = race, y = log_antibody, fill = race)) +
 geom_boxplot() +
  labs(title = "Antibody Levels by Race/Ethnicity",
      x = "Race/Ethnicity",
      y = "Log Antibody Level") +
  theme_minimal()
# Boxplot of log_antibody by smoking status
p4 <- ggplot(dat1, aes(x = smoking, y = log_antibody, fill = smoking)) +
  geom_boxplot() +
  labs(title = "Antibody Levels by Smoking Status",
      x = "Smoking Status",
      y = "Log Antibody Level") +
  theme_minimal()
# Boxplot of log_antibody by diabetes status
p5 <- ggplot(dat1, aes(x = diabetes, y = log_antibody, fill = diabetes)) +
  geom boxplot() +
 labs(title = "Antibody Levels by Diabetes Status",
```

```
x = "Diabetes Status",
y = "Log Antibody Level") +
theme_minimal()
```

p2+p3+p4+p5



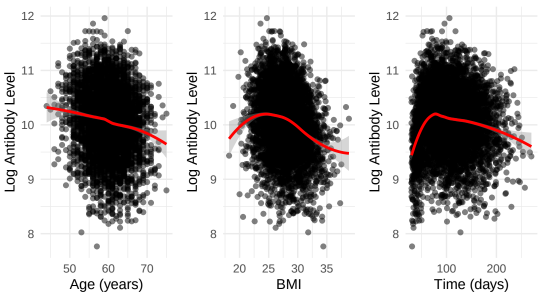
2.3 Correlation Plot



2.4 Scatterplots

```
# Create scatterplots for continuous variables vs response
p6 <- ggplot(dat1, aes(x = age, y = log_antibody)) +
  geom_point(alpha = 0.5) +
  geom_smooth(method = "loess", color = "red") +
  labs(title = "Age vs Log Antibody Level",
       x = "Age (years)", y = "Log Antibody Level") +
  theme_minimal()
p7 <- ggplot(dat1, aes(x = bmi, y = log_antibody)) +
  geom point(alpha = 0.5) +
  geom_smooth(method = "loess", color = "red") +
  labs(title = "BMI vs Log Antibody Level",
       x = "BMI", y = "Log Antibody Level") +
  theme_minimal()
p8 <- ggplot(dat1, aes(x = time, y = log_antibody)) +
  geom_point(alpha = 0.5) +
  geom_smooth(method = "loess", color = "red") +
  labs(title = "Time Since Vaccination vs Log Antibody Level",
       x = "Time (days)", y = "Log Antibody Level") +
  theme_minimal()
```





There are

some potential nonlinear trend between log_antibody and bmi, and between log_antibody and time.

3. Model Training with Cross-validation

3.1 Linear models

```
model_formula <- log_antibody ~ . - id

# Set up cross-validation control
ctrl <- trainControl(method = "cv", number = 10)

# 3.1 Linear Regression
set.seed(123)
lm.fit <- train(
    model_formula,
    data = dat1,
    method = "lm",
    trControl = ctrl
)
summary(lm.fit$finalModel)</pre>
```

3.1.1 Linear regression

```
##
## Call:
## lm(formula = .outcome ~ ., data = dat)
##
```

```
## Residuals:
##
      Min
                  Median
               10
                               30
                                      Max
## -2.14396 -0.35840 0.02944 0.37802 1.65090
## Coefficients:
##
                  Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                26.6751961 2.3149812 11.523 < 2e-16 ***
                -0.0205979  0.0019385  -10.626  < 2e-16 ***
## age
## genderMale
              -0.2974929  0.0155977  -19.073  < 2e-16 ***
## raceAsian
                -0.0060422 0.0344613 -0.175
                                           0.8608
## raceBlack
               -0.0075295 0.0196815 -0.383
                                            0.7021
## raceHispanic -0.0417571 0.0273309 -1.528
                                             0.1266
## smokingFormer 0.0219907 0.0173992 1.264
                                            0.2063
## smokingCurrent -0.1934834 0.0269576 -7.177 8.15e-13 ***
## height
                ## weight
                0.0859034 0.0143481 5.987 2.29e-09 ***
## bmi
                ## diabetesYes
                 0.0112795 0.0215643 0.523
                                           0.6010
## hypertensionYes -0.0179106 0.0260931 -0.686
                                            0.4925
                 0.0015181 0.0017049
                                    0.890
                                            0.3733
## ldl
                -0.0001645 0.0004028 -0.409
                                            0.6829
## time
                -0.0003011 0.0001795 -1.677
                                            0.0936 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.5503 on 4984 degrees of freedom
## Multiple R-squared: 0.1513, Adjusted R-squared: 0.1488
## F-statistic: 59.25 on 15 and 4984 DF, p-value: < 2.2e-16
```

3.1.2 Ridge Regression

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

## s1

## (Intercept) 12.7384560516

## age -0.0197679168

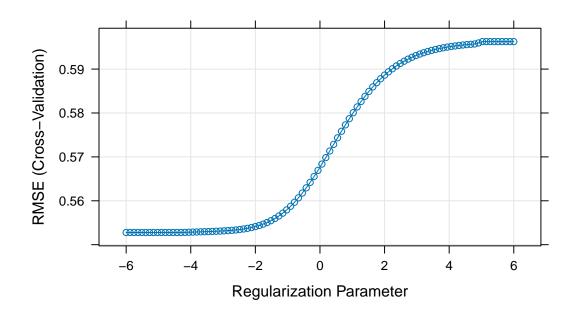
## genderMale -0.2880871615

## raceAsian -0.0038318346

## raceBlack -0.0066654722

## raceHispanic -0.0417690245
```

```
## smokingFormer
                    0.0242172365
## smokingCurrent -0.1847225211
## height
                   -0.0001949552
## weight
                   -0.0009185524
## bmi
                   -0.0473178893
## diabetesYes
                    0.0113069924
## hypertensionYes -0.0166641972
## sbp
                    0.0010847735
## ldl
                   -0.0001614795
## time
                   -0.0002807285
print(ridge.fit$bestTune)
##
      alpha
               lambda
          0 0.0135275
## 15
plot(ridge.fit, xTrans = log)
```



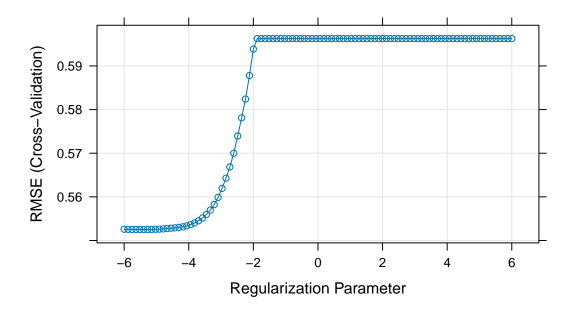
3.1.3 Lasso Regression

```
## 16 x 1 sparse Matrix of class "dgCMatrix"
##
## (Intercept)
                    1.272648e+01
                   -1.915006e-02
## age
## genderMale
                   -2.859715e-01
## raceAsian
## raceBlack
## raceHispanic
                   -2.472809e-02
## smokingFormer
                    1.593301e-02
## smokingCurrent -1.770968e-01
## height
## weight
                   -1.029699e-04
## bmi
                   -4.801089e-02
## diabetesYes
                    4.273525e-05
## hypertensionYes
## sbp
## ldl
## time
                   -1.848766e-04
```

print(lasso.fit\$bestTune)

```
## alpha lambda
## 6 1 0.004544037
```

plot(lasso.fit, xTrans = log)

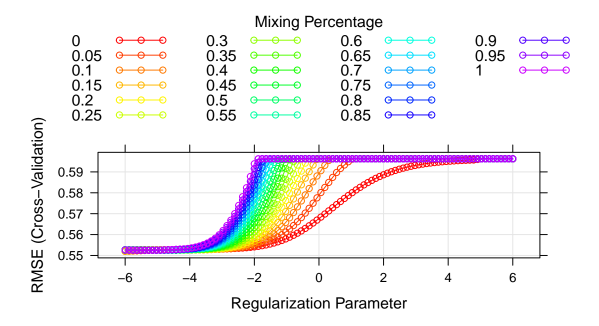


3.1.4 Elastic Net Regression

```
##
              Length Class
                               Mode
              100
## a0
                    -none-
                               numeric
## beta
              1500
                    dgCMatrix
                              S4
## df
              100
                    -none-
                               numeric
## dim
               2
                   -none-
                               numeric
## lambda
              100
                   -none-
                               numeric
                    -none-
## dev.ratio
             100
                               numeric
## nulldev
               1 -none-
                               numeric
## npasses
               1 -none-
                               numeric
## jerr
                1 -none-
                               numeric
## offset
                1 -none-
                               logical
## call
                5 -none-
                               call
## nobs
                1 -none-
                               numeric
## lambdaOpt
                1
                   -none-
                               numeric
## xNames
               15 -none-
                               character
## problemType
                1 -none-
                               character
## tuneValue
                2 data.frame list
## obsLevels
                1
                    -none-
                               logical
## param
                0
                    -none-
                               list
```

```
print(enet.fit$bestTune)
```

```
## alpha lambda
## 101 0.05 0.002478752
```



```
# Show coefficients of the final Elastic Net model
coef_enet <- coef(enet.fit$finalModel, enet.fit$bestTune$lambda)
print(coef_enet)</pre>
```

```
## 16 x 1 sparse Matrix of class "dgCMatrix"
##
## (Intercept)
                   16.0225822334
                   -0.0202906145
## age
## genderMale
                   -0.2942760076
## raceAsian
                   -0.0036856007
## raceBlack
                   -0.0066593257
## raceHispanic
                   -0.0417893824
## smokingFormer
                    0.0235504017
## smokingCurrent
                   -0.1891969582
## height
                   -0.0193618230
## weight
                    0.0193847860
## bmi
                   -0.1063133265
## diabetesYes
                    0.0111516911
## hypertensionYes -0.0167677112
## sbp
                    0.0012484795
## ldl
                   -0.0001495619
## time
                   -0.0002879691
```

```
coef_enet <- coef(enet.fit$finalModel, enet.fit$bestTune$lambda)
print(coef_enet)</pre>
```

```
## 16 x 1 sparse Matrix of class "dgCMatrix"
## s1
## (Intercept) 16.0225822334
## age -0.0202906145
## genderMale -0.2942760076
```

```
-0.0036856007
## raceAsian
                 -0.0066593257
## raceBlack
## raceHispanic
                  -0.0417893824
## smokingFormer
                  0.0235504017
## smokingCurrent -0.1891969582
## height
                  -0.0193618230
## weight
                  0.0193847860
## bmi
                  -0.1063133265
## diabetesYes
                   0.0111516911
## hypertensionYes -0.0167677112
## sbp
                  0.0012484795
## ldl
                  -0.0001495619
## time
                  -0.0002879691
```

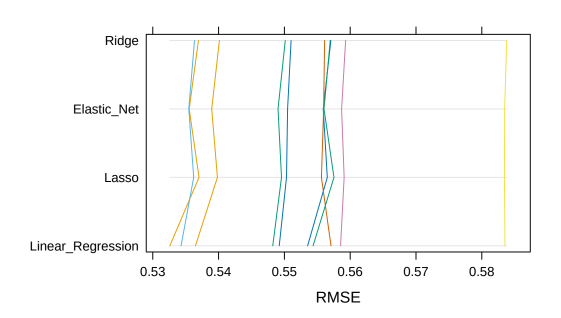
```
resamp <- resamples(list(
   Linear_Regression = lm.fit,
   Ridge = ridge.fit,
   Lasso = lasso.fit,
   Elastic_Net = enet.fit
))
summary(resamp)</pre>
```

3.1.5 Model Comparison (resampling)

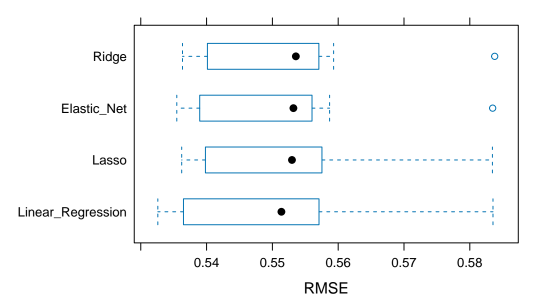
```
##
## Call:
## summary.resamples(object = resamp)
## Models: Linear_Regression, Ridge, Lasso, Elastic_Net
## Number of resamples: 10
##
## MAE
                          Min.
                                  1st Qu.
                                             Median
                                                         Mean
                                                                3rd Qu.
## Linear Regression 0.4261808 0.4320149 0.4340128 0.4390390 0.4425761 0.4726409
                     0.4281731 0.4334230 0.4362590 0.4405051 0.4454071 0.4720434
## Ridge
                     0.4276920 0.4330647 0.4364914 0.4403860 0.4449243 0.4721327
## Lasso
                     0.4274328 0.4330542 0.4354391 0.4399269 0.4448176 0.4720702
## Elastic_Net
##
                     NA's
## Linear_Regression
                        0
                        0
## Ridge
## Lasso
                        0
## Elastic_Net
                        0
##
## RMSE
##
                                  1st Qu.
                                             Median
                                                                3rd Qu.
                          Min.
                                                         Mean
                                                                              Max.
## Linear_Regression 0.5325907 0.5394231 0.5513718 0.5507854 0.5564032 0.5835238
                     0.5363406\ 0.5426274\ 0.5535668\ 0.5527796\ 0.5570390\ 0.5837848
## Ridge
## Lasso
                     0.5362188 0.5422580 0.5529754 0.5525153 0.5572793 0.5834368
                     0.5354741 0.5414764 0.5531955 0.5519547 0.5560081 0.5834749
## Elastic_Net
## Linear_Regression
```

```
## Ridge
                         0
## Lasso
                         0
## Elastic_Net
##
## Rsquared
##
                            Min.
                                   1st Qu.
                                               Median
                                                           Mean
                                                                   3rd Qu.
                                                                                Max.
## Linear_Regression 0.10033968 0.1319832 0.1534930 0.1479018 0.1622355 0.1906302
## Ridge
                      0.09354935\ 0.1329969\ 0.1421825\ 0.1418538\ 0.1502030\ 0.1878098
                      0.09475560 0.1342489 0.1433703 0.1428172 0.1503016 0.1912475
## Lasso
## Elastic_Net
                      0.09585259\ 0.1339416\ 0.1458955\ 0.1443698\ 0.1542137\ 0.1902642
                      NA's
## Linear_Regression
## Ridge
                         0
## Lasso
                         0
## Elastic_Net
                         0
```

parallelplot(resamp, metric = "RMSE")



bwplot(resamp, metric = "RMSE")



The

RMSE from resampling looks similar to each other.

```
# Function to calculate RMSE
rmse <- function(actual, predicted) {
   sqrt(mean((actual - predicted)^2))
}</pre>
```

```
# Make predictions using each model
lm.pred <- predict(lm.fit, newdata = dat2)
ridge.pred <- predict(ridge.fit, newdata = dat2)
lasso.pred <- predict(lasso.fit, newdata = dat2)
enet.pred <- predict(enet.fit, newdata = dat2)</pre>
```

```
# Calculate test MSE for each model
lm.test.error <- mean((lm.pred - dat2[,"log_antibody"])^2)
ridge.test.error <- mean((ridge.pred - dat2[,"log_antibody"])^2)
lasso.test.error <- mean((lasso.pred - dat2[,"log_antibody"])^2)
enet.test.error <- mean((enet.pred - dat2[,"log_antibody"])^2)</pre>
```

```
print(test_performance)
```

3.1.6 Test Set Performance

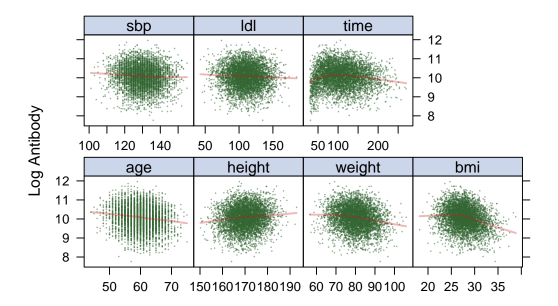
```
## Model Test_ERROR
## 1 Linear Regression 0.3229854
## 2 Ridge Regression 0.3258077
## 3 Lasso Regression 0.3288173
## 4 Elastic Net Regression 0.3247256
```

They are very similar to each other. We probably want to choose the Lasso model as it is the simplest model among those four models.

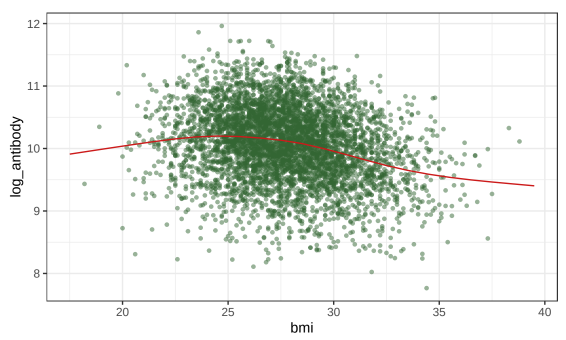
3.2 Non-linear Models

3.2.1 Smoothing Spline We use scatterplot to explore the relationship between the log antibody level and other continuous variables. Time and bmi tend shows potentially nonlinear trend.

```
x = model.matrix(log_antibody ~ . - id, data = dat1)[, -1]
y = dat1[,"log_antibody"]
x_test = model.matrix(log_antibody ~ . - id, data = dat2)[, -1]
y_test = dat2[,"log_antibody"]
```

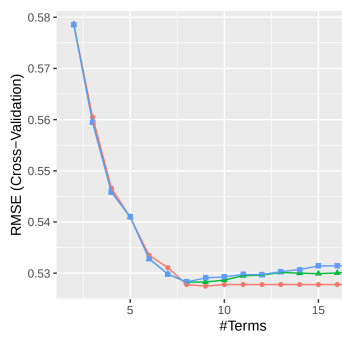


```
# choose the best df
fit.ss = smooth.spline(dat1$bmi, dat1$log_antibody)
fit.ss$df
## [1] 5.769535
print(fit.ss)
## Call:
## smooth.spline(x = dat1$bmi, y = dat1$log_antibody)
## Smoothing Parameter spar= 0.8460071 lambda= 0.05288648 (13 iterations)
## Equivalent Degrees of Freedom (Df): 5.769535
## Penalized Criterion (RSS): 61.11728
## GCV: 0.3330358
# plot optimal fit
bmi.grid = seq(from = 17.5, to = 40, by = 1)
pred.ss = predict(fit.ss, x = bmi.grid)
pred.ss.df = data.frame(pred = pred.ss$y, bmi = bmi.grid)
p = ggplot(dat1, aes(x = bmi, y = log_antibody)) +
  geom_point(color = rgb(.2, .4, .2, .5), size = 1) + theme_bw()
geom_line(aes(x = bmi, y = pred), data = pred.ss.df,
color = rgb(.8, .1, .1, 1)) + theme_bw()
```



The scatter plot shows a weak non-linear relationship between BMI and log-transformed antibody levels. A smooth spline with an effective degree of freedom of approximately 5.77 was fitted. The trend suggests that antibody levels tend to be higher for individuals with BMI in the range of 25–28 and slightly decrease for higher BMI.

```
set.seed(123)
ctrl1 = trainControl(method = "cv", number = 10)
mars_grid = expand.grid(degree = 1:3, nprune = 2:20)
mars.fit = train(x, y, method = "earth", tuneGrid = mars_grid,
trControl = ctrl1)
ggplot(mars.fit)
```

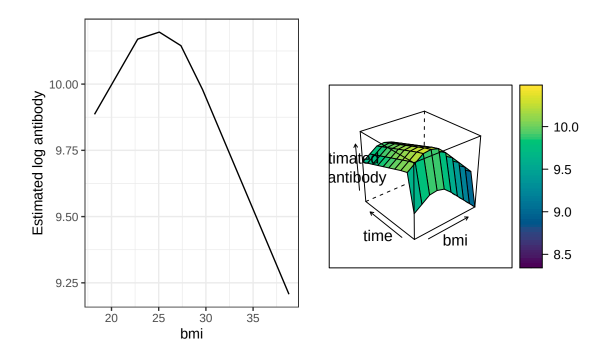


3.2.2 Multivariate Adaptive Regression Splines (MARS)

```
mars.fit$bestTune
##
     nprune degree
## 8
coef(mars.fit$finalModel)
##
      (Intercept)
                     h(27.8-bmi)
                                      h(time-57)
                                                     h(57-time)
                                                                     genderMale
     10.847446930
                    -0.061997354
                                    -0.002254182
                                                                   -0.296290451
##
                                                    -0.033529326
##
        h(age-59)
                       h(59-age) smokingCurrent
                                                    h(bmi-23.7)
     -0.022957648
                     0.016138468
                                    -0.205126851
                                                    -0.084380175
##
summary(mars.fit$finalModel)
## Call: earth(x=matrix[5000,15], y=c(10.65,9.889,1...), keepxy=TRUE, degree=1,
               nprune=9)
##
##
##
                  coefficients
## (Intercept)
                    10.8474469
## genderMale
                    -0.2962905
## smokingCurrent
                    -0.2051269
## h(59-age)
                     0.0161385
## h(age-59)
                    -0.0229576
## h(bmi-23.7)
                    -0.0843802
## h(27.8-bmi)
                    -0.0619974
## h(57-time)
                    -0.0335293
## h(time-57)
                    -0.0022542
##
## Selected 9 of 10 terms, and 5 of 15 predictors (nprune=9)
## Termination condition: RSq changed by less than 0.001 at 10 terms
```

```
## Importance: bmi, genderMale, time, age, smokingCurrent, raceAsian-unused, ...
## Number of terms at each degree of interaction: 1 8 (additive model)
## GCV 0.2787787 RSS 1384.431 GRSq 0.2166152 RSq 0.2216218
```

We fitted a MARS model with 10-fold cross-validation. The optimal model selected 9 basis functions with degree 1 (additive model). Among 15 candidate predictors, 5 variables were retained (bmi, gender, time, age, and smoking). The fitted model explained approximately 22% of the variation in log-transformed antibody levels. The model identified several meaningful breakpoints such as bmi = 23.7 and bmi = 27.8, suggesting non-linear relationships. For example, BMI showed a negative effect on antibody levels when exceeding 23.7, and males had lower antibody levels compared to females.



```
set.seed(123)
gam.fit = train(x, y, method = "gam", trControl = ctrl1)
summary(gam.fit)
```

3.2.3 Generalized Additive Model (GAM)

```
## Family: gaussian
## Link function: identity
## Formula:
## .outcome ~ genderMale + raceAsian + raceBlack + raceHispanic +
      smokingFormer + smokingCurrent + diabetesYes + hypertensionYes +
##
##
      s(age) + s(sbp) + s(ldl) + s(bmi) + s(time) + s(height) +
##
      s(weight)
## Parametric coefficients:
                  Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                  10.228177
                             0.015328 667.269 < 2e-16 ***
## genderMale
                -0.297837
                             0.014933 -19.945 < 2e-16 ***
## raceAsian
                -0.003296
                            0.033009 -0.100
                                                0.920
## raceBlack
                 -0.010509 0.018837 -0.558
                                                 0.577
## raceHispanic
                 -0.037424
                             0.026176 -1.430
                                                 0.153
                                       1.334
## smokingFormer
                  0.022219
                            0.016660
                                                 0.182
## smokingCurrent -0.193175
                            0.025834 -7.478 8.9e-14 ***
## diabetesYes
                  0.014230
                             0.020640 0.689
                                                 0.491
## hypertensionYes -0.007678
                            0.015995 -0.480
                                                 0.631
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Approximate significance of smooth terms:
                  edf Ref.df
##
                                 F p-value
## s(age)
            9.908e-01
                      9 13.733 <2e-16 ***
## s(sbp)
                         9 0.000
                                    0.765
          6.175e-07
## s(ldl)
            6.648e-07
                          9 0.000
                                    0.639
## s(bmi)
                          9 41.897 <2e-16 ***
            4.179e+00
## s(time)
          7.892e+00
                          9 44.960 <2e-16 ***
## s(height) 1.234e+00
                          9 0.278 0.121
                          9 0.000 0.666
## s(weight) 2.262e-06
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## R-sq.(adj) =
                 0.22
                       Deviance explained = 22.4%
## GCV = 0.27867 Scale est. = 0.27738
gam.fit$bestTune
    select method
##
      TRUE GCV.Cp
## 2
gam.fit$finalModel
##
## Family: gaussian
## Link function: identity
##
## Formula:
```

```
## .outcome ~ genderMale + raceAsian + raceBlack + raceHispanic +
## smokingFormer + smokingCurrent + diabetesYes + hypertensionYes +
## s(age) + s(sbp) + s(ldl) + s(bmi) + s(time) + s(height) +
## s(weight)
##
## Estimated degrees of freedom:
## 0.991 0.000 0.000 4.179 7.892 1.234 0.000
## total = 23.3
##
## GCV score: 0.2786734
```

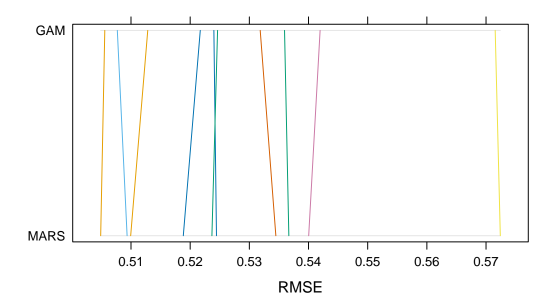
A Generalized Additive Model (GAM) with Gaussian distribution and identity link was fitted. The model included both parametric terms and smooth terms. Among parametric terms, male gender and current smoking status were significantly associated with lower log-transformed antibody levels. Smooth terms indicated non-linear effects of age (edf 1), BMI (edf 4.18), and time since vaccination (edf 7.89) on antibody levels, all statistically significant (p < 0.001). Variables such as SBP, LDL, and weight showed non-significant effects. The model explained approximately 22% of the variability in log-transformed antibody levels.

```
resamp_non_linear <- resamples(list(
   MARS = mars.fit,
   GAM = gam.fit
))
summary(resamp_non_linear)</pre>
```

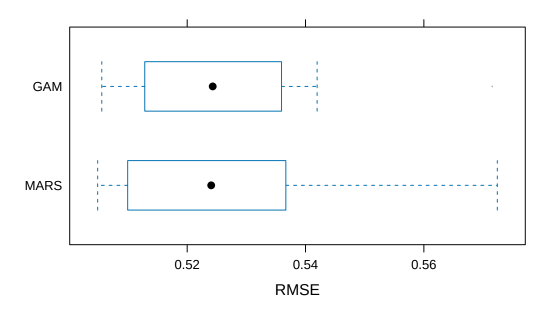
3.2.4 Model Comparison (resampling)

```
##
## Call:
## summary.resamples(object = resamp_non_linear)
##
## Models: MARS, GAM
## Number of resamples: 10
##
## MAE
##
             Min.
                     1st Qu.
                                Median
                                            Mean 3rd Qu.
                                                                Max. NA's
## MARS 0.4078837 0.4098168 0.4163554 0.4220725 0.428591 0.4641090
                                                                        0
       0.4078470 0.4112544 0.4168810 0.4225034 0.428161 0.4635896
                                                                        0
##
## RMSE
             Min.
                    1st Qu.
                                Median
                                            Mean
                                                    3rd Qu.
                                                                 Max. NA's
## MARS 0.5048757 0.5121685 0.5240607 0.5274715 0.5361255 0.5724126
                                                                         0
  GAM 0.5055708 0.5150506 0.5243183 0.5277724 0.5349189 0.5715483
                                                                         0
##
## Rsquared
##
             Min.
                    1st Qu.
                                Median
                                            Mean
                                                    3rd Qu.
                                                                 Max. NA's
## MARS 0.1566346 0.1954579 0.2274358 0.2187657 0.2465799 0.2627660
                                                                         0
## GAM 0.1586929 0.1935757 0.2259290 0.2177415 0.2449893 0.2595686
                                                                         0
```

```
parallelplot(resamp_non_linear, metric = "RMSE")
```



bwplot(resamp_non_linear, metric = "RMSE")



3.1.6 Test Set Performance

''' r
Make predictions using each mode
pred_ss = predict(fit.ss, x = dat2\$bmi)
pred_mars = predict(mars.fit, newdata = x_test)

4. Model Comparison between Linear and non-linear Models

Model	MSE
MARS	0.2838458
Linear Regression	0.3229854
Smoothing Spline	0.3247256
GAM	0.3249953
Ridge Regression	0.3258077
Lasso Regression	0.3288173
Elastic Net Regression	0.3584620

The best model is the MARS model with the smallest test error.