# Midterm Project

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## **Exploratory Data Analysis**

```
load("./data/dat1.RData")
load("./data/dat2.RData")
# no missing data
all(is.na(dat1))
## [1] FALSE
all(is.na(dat2))
## [1] FALSE
ifelse(all(names(dat1) == names(dat2)), "train and test data have same structure", "train and test data
## [1] "train and test data have same structure"
str(dat1)
## 'data.frame': 5000 obs. of 14 variables:
## $ id
                : int 12345678910...
## $ age
                : num 50 71 58 63 56 59 67 62 60 64 ...
## $ gender
                : int 0 1 1 0 1 1 0 1 0 1 ...
                : Factor w/ 4 levels "1","2","3","4": 1 1 1 1 3 4 1 4 1 ...
## $ race
## $ smoking
                : Factor w/ 3 levels "0","1","2": 1 1 2 1 1 1 1 1 1 1 ...
## $ height
                : num 176 176 169 167 163 ...
## $ weight
                : num 68.3 69.6 76.9 90 83.9 86.8 91.4 87.7 85.7 76.6 ...
## $ bmi
                        22 22.6 27 32.1 31.7 30.8 29.7 28.1 29 31.5 ...
                 : num
                : int 0000000000...
## $ diabetes
## $ hypertension: num 0 1 0 1 0 1 1 0 0 1 ...
                : num 130 149 127 138 123 132 133 130 129 134 ...
## $ SBP
## $ LDL
                 : num 82 129 101 93 97 108 89 96 120 135 ...
                : num 76 82 168 105 193 143 63 78 61 88 ...
## $ time
## $ log_antibody: num 10.65 9.89 10.9 9.91 9.56 ...
# The 'id' column has no actual meaning, so we remove it.
dat1 <- dat1 %>%
 select(-id)
dat2 <- dat2 %>%
 select(-id)
# Convert categorical variables to labeled factors
convert_factors <- function(df) {</pre>
```

```
df %>%
  mutate(
    gender = factor(gender, levels = c(0, 1), labels = c("Female", "Male")),
    race = factor(race, levels = c(1, 2, 3, 4), labels = c("White", "Asian", "Black", "Hispanic")),
    smoking = factor(smoking, levels = c(0, 1, 2), labels = c("Never", "Former", "Current")),
    diabetes = factor(diabetes, levels = c(0, 1), labels = c("No", "Yes")),
    hypertension = factor(hypertension, levels = c(0, 1), labels = c("No", "Yes"))
}
dat1 <- convert_factors(dat1)
dat2 <- convert_factors(dat2)</pre>
```

### Univariate analysis (continuous & categorical)

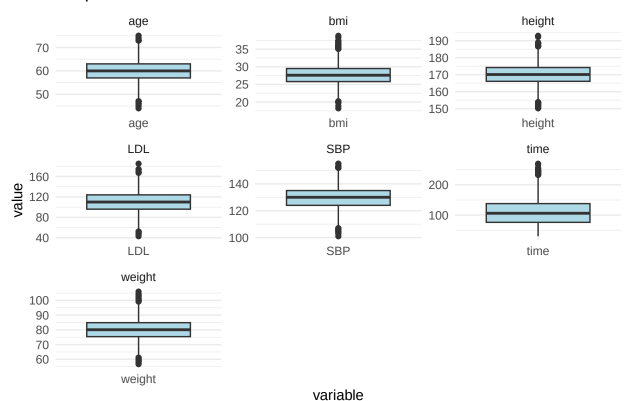
```
continuous_var <- dat1 %>%
   select(age, height, weight, bmi, SBP, LDL, time)

categorical_var <- dat1 %>%
   select(gender, race, smoking, diabetes, hypertension)

# ---- Continuous Variables ----
# summary
summary(continuous_var)
```

```
##
                      height
                                      weight
                                                       bmi
        age
                        :150.2
                                  Min. : 56.70
                                                         :18.20
## Min.
         :44.00
                 Min.
                                                 Min.
## 1st Qu.:57.00
                  1st Qu.:166.1
                                  1st Qu.: 75.40
                                                  1st Qu.:25.80
## Median :60.00
                 Median :170.1
                                  Median : 80.10
                                                 Median :27.60
## Mean :59.97
                  Mean :170.1
                                  Mean : 80.11
                                                  Mean :27.74
                                  3rd Qu.: 84.90
## 3rd Qu.:63.00
                  3rd Qu.:174.2
                                                  3rd Qu.:29.50
          :75.00
                  Max. :192.9
                                        :106.00
                                                  Max. :38.80
## Max.
                                  Max.
##
        SBP
                       LDL
                                       time
## Min.
          :101.0
                  Min.
                         : 43.0
                                  Min. : 30.0
                  1st Qu.: 96.0
## 1st Qu.:124.0
                                  1st Qu.: 76.0
## Median :130.0
                  Median :110.0
                                  Median :106.0
         :129.9
                  Mean :109.9
                                       :108.9
## Mean
                                  Mean
## 3rd Qu.:135.0
                   3rd Qu.:124.0
                                  3rd Qu.:138.0
## Max.
          :155.0
                  Max.
                         :185.0
                                  Max.
                                         :270.0
# boxplots
continuous_var_long <- continuous_var %>%
 tidyr::pivot_longer(cols = everything(), names_to = "variable", values_to = "value")
ggplot(continuous_var_long, aes(x = variable, y = value)) +
 geom boxplot(fill = "lightblue") +
 facet_wrap(~variable, scales = "free", ncol = 3) +
 theme_minimal() +
 labs(title = "Boxplots of Continuous Variables")
```

# **Boxplots of Continuous Variables**



```
# density plots
ggplot(continuous_var_long, aes(x = value)) +
  geom_density(fill = "lightblue", alpha = 0.6) +
  facet_wrap(~variable, scales = "free", ncol = 3) +
  theme_minimal() +
  labs(title = "Density Plots of Continuous Variables", x = "Value", y = "Density")
```

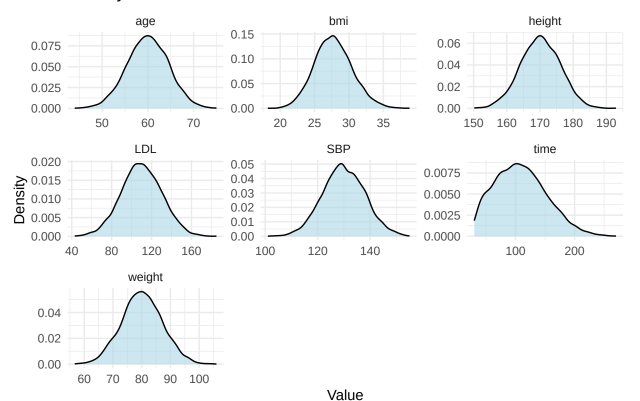
### **Density Plots of Continuous Variables**

##

##

gender

Female:2573



```
# ---- Categorical Variables ----
# summary
summary(categorical_var)
```

smoking

Never :3010

race

White

:3221

diabetes

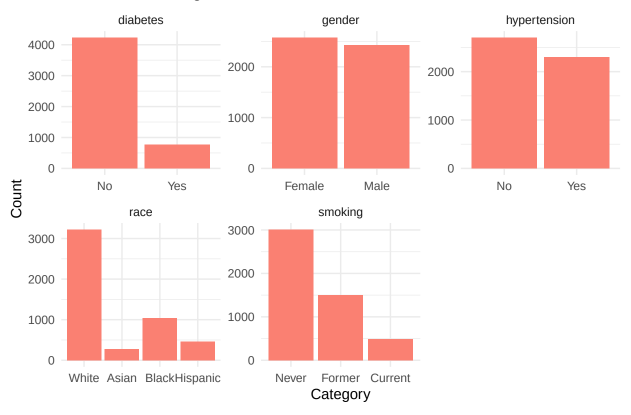
No:4228

hypertension

No :2702

```
##
   Male :2427
                  Asian
                          : 278
                                  Former:1504
                                                 Yes: 772
                                                             Yes:2298
##
                  Black
                          :1036
                                  Current: 486
##
                  Hispanic: 465
# bar plots
categorical_var_long <- categorical_var %>%
  tidyr::pivot_longer(cols = everything(), names_to = "variable", values_to = "value")
ggplot(categorical_var_long, aes(x = value)) +
  geom_bar(fill = "salmon") +
  facet_wrap(~variable, scales = "free", ncol = 3) +
  theme_minimal() +
  labs(title = "Bar Plots of Categorical Variables", x = "Category", y = "Count")
```

## Bar Plots of Categorical Variables



According to the box plot for continuous variables:

Age, BMI, and SBP appear reasonably normally distributed, with expected ranges for an adult population;
 LDL cholesterol and time since vaccination show a wider range, right-skewness and some outliers, which may impact linear models.

According to the bar plot for categorical variables:

- Gender is fairly balanced between Female and Male;
- Race is skewed, with a majority of participants identifying as White (Category 1). Other racial/ethnic groups are underrepresented;
- Smoking status shows that the majority are never smokers (Category 0), with fewer current and former smokers;
- A large proportion of participants do not have diabetes;
- A moderate split exists for hypertension, which may contribute meaningfully to clinical outcome variation
- Demographically, the population is balanced by gender but skewed by race and smoking status.

```
# response variable `log_antibody`

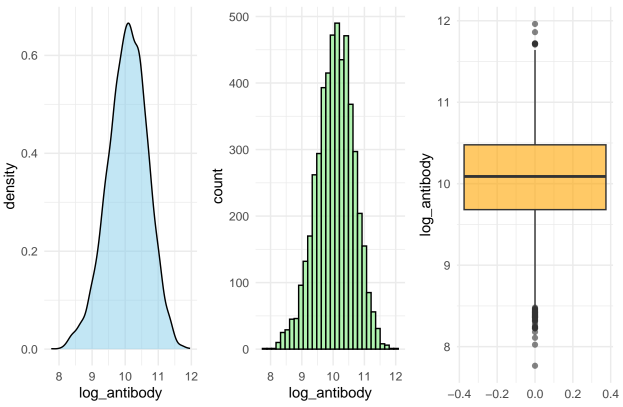
# Density plot
p1 <- ggplot(dat1, aes(x = log_antibody)) +
    geom_density(fill = "skyblue", alpha = 0.5) +
    ggtitle("Density Plot of log_antibody") +
    xlab("log_antibody") +
    theme_minimal()</pre>
```

```
# Histogram
p2 <- ggplot(dat1, aes(x = log_antibody)) +
    geom_histogram(bins = 30, fill = "lightgreen", color = "black", alpha = 0.7) +
    ggtitle("Histogram of log_antibody") +
    xlab("log_antibody") +
    theme_minimal()

# Boxplot
p3 <- ggplot(dat1, aes(y = log_antibody)) +
    geom_boxplot(fill = "orange", alpha = 0.6) +
    ggtitle("Boxplot of log_antibody") +
    ylab("log_antibody") +
    theme_minimal()

grid.arrange(p1, p2, p3, ncol = 3)</pre>
```

# Density Plot of log\_antibodyHistogram of log\_antibodyBoxplot of log\_



log\_antibody (response) appears fairly symmetrical, which supports its use as a continuous response in linear or GAM models.

Overall, we believe the response variable <code>log\_antibody</code> is well-behaved, and further correlation analysis(eg. bivariate) is needed.

Next, we assess correlations and non-linear trends to guide model form.

### **Correlation Analysis**

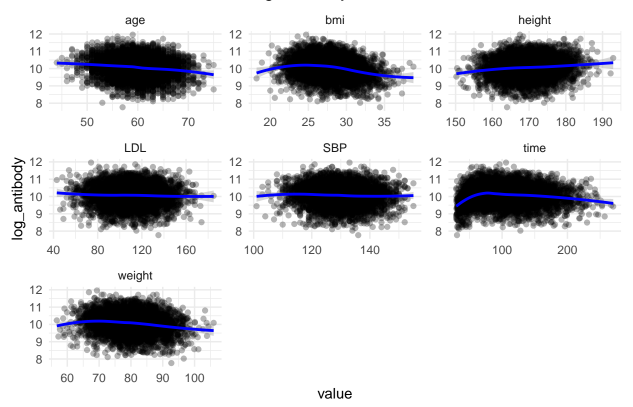
We first analyze the relationship between log\_antibody (response variable) and continuous variables & correlations among continuous variables themselves.

```
continuous_var_long <- dat1 %>%
    select(age, height, weight, bmi, SBP, LDL, time, log_antibody) %>%
    tidyr::pivot_longer(cols = -log_antibody, names_to = "variable", values_to = "value")

# Scatterplots with smoothing lines
ggplot(continuous_var_long, aes(x = value, y = log_antibody)) +
    geom_point(alpha = 0.3) +
    geom_smooth(method = "loess", color = "blue") +
    facet_wrap(~variable, scales = "free", ncol = 3) +
    theme_minimal() +
    labs(title = "Continuous Predictors vs. log_antibody")
```

##  $geom_smooth()$  using formula = 'y ~ x'

### Continuous Predictors vs. log\_antibody

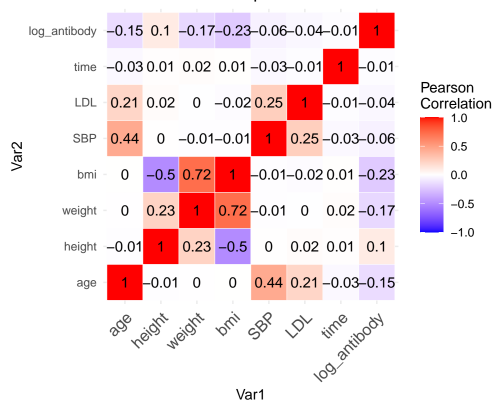


Using LOESS method, we observe linearity between predictors and the response. The plot shows that bmi, time, and weight has clear non linear trend against resopnse log\_antibody, indicating potential need to use GAM or non linear model.

```
continous_name <- c("age", "height", "weight", "bmi", "SBP", "LDL", "time", "log_antibody")
dat_cont <- dat1[ , continous_name]</pre>
# coefficient matrix
cor_matrix <- cor(dat_cont, use = "complete.obs", method = "pearson")</pre>
print(round(cor_matrix, 2))
##
                                              SBP
                  age height weight
                                                    LDL time log_antibody
                                       bmi
## age
                 1.00 -0.01
                                0.00
                                      0.00
                                             0.44
                                                   0.21 -0.03
                                                                      -0.15
## height
                -0.01
                        1.00
                                0.23 - 0.50
                                             0.00
                                                   0.02 0.01
                                                                       0.10
```

```
## weight
                 0.00
                        0.23
                               1.00 0.72 -0.01 0.00 0.02
                                                                    -0.17
                 0.00
                       -0.50
                               0.72 1.00 -0.01 -0.02 0.01
                                                                    -0.23
## bmi
                 0.44
                                          1.00 0.25 -0.03
## SBP
                        0.00
                              -0.01 -0.01
                                                                    -0.06
                                                 1.00 -0.01
## LDL
                 0.21
                        0.02
                               0.00 -0.02 0.25
                                                                    -0.04
## time
                -0.03
                        0.01
                               0.02 0.01 -0.03 -0.01 1.00
                                                                    -0.01
## log antibody -0.15
                        0.10
                              -0.17 -0.23 -0.06 -0.04 -0.01
                                                                     1.00
cor_melt <- melt(cor_matrix)</pre>
ggplot(cor_melt, aes(Var1, Var2, fill = value)) +
  geom_tile(color = "white") +
  scale_fill_gradient2(low = "blue", high = "red", mid = "white",
                       midpoint = 0, limit = c(-1, 1), space = "Lab",
                       name = "Pearson\nCorrelation") +
  geom_text(aes(label = round(value, 2)), color = "black", size = 4) +
  theme minimal() +
  theme(axis.text.x = element_text(angle = 45, vjust = 1,
                                   size = 12, hjust = 1)) +
  coord_fixed() +
  ggtitle("Correlation Heatmap of Continuous Variables")
```

### Correlation Heatmap of Continuous Variables



From the matrix and plot, we can see that the Pearson correlation coefficient between bmi and weight, bmi and height exceed 0.5, indicating that multicollinearity may exist.

```
vif(lm_full)
```

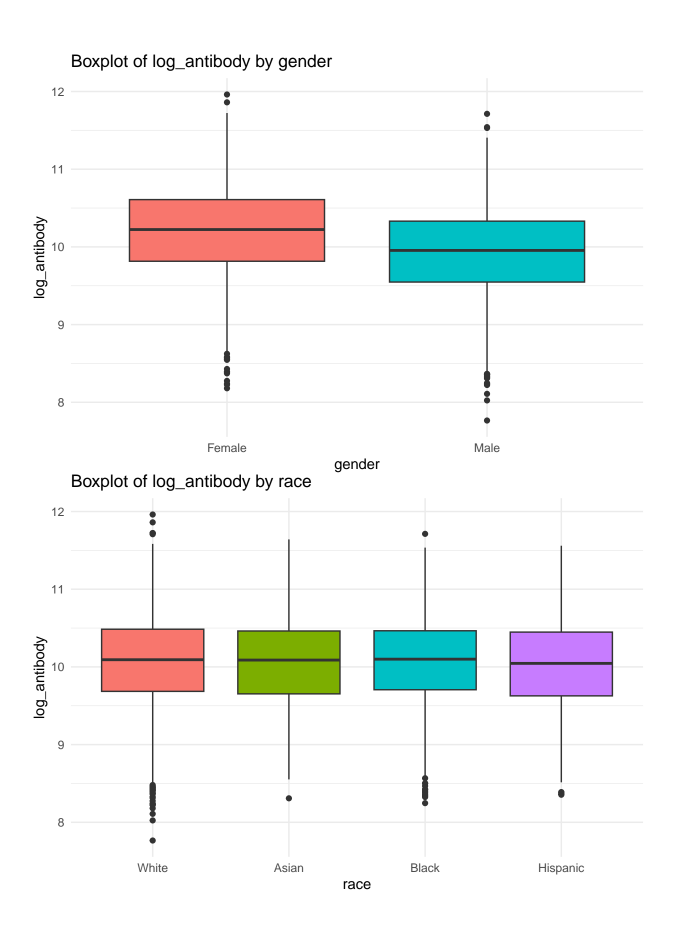
```
GVIF Df GVIF^(1/(2*Df))
## age
                  1.258104 1
                                     1.121652
## gender
                  1.002988 1
                                     1.001493
                  1.002682 2
## smoking
                                     1.000670
## height
                107.111548 1
                                    10.349471
## weight
                169.112707 1
                                    13.004334
                213.764468 1
                                    14.620686
## bmi
## diabetes
                  1.001898 1
                                     1.000949
                                     1.670731
## hypertension
                  2.791341 1
## SBP
                  3.070211 1
                                     1.752202
## LDL
                  1.085268 1
                                     1.041762
## time
                  1.002242 1
                                     1.001120
```

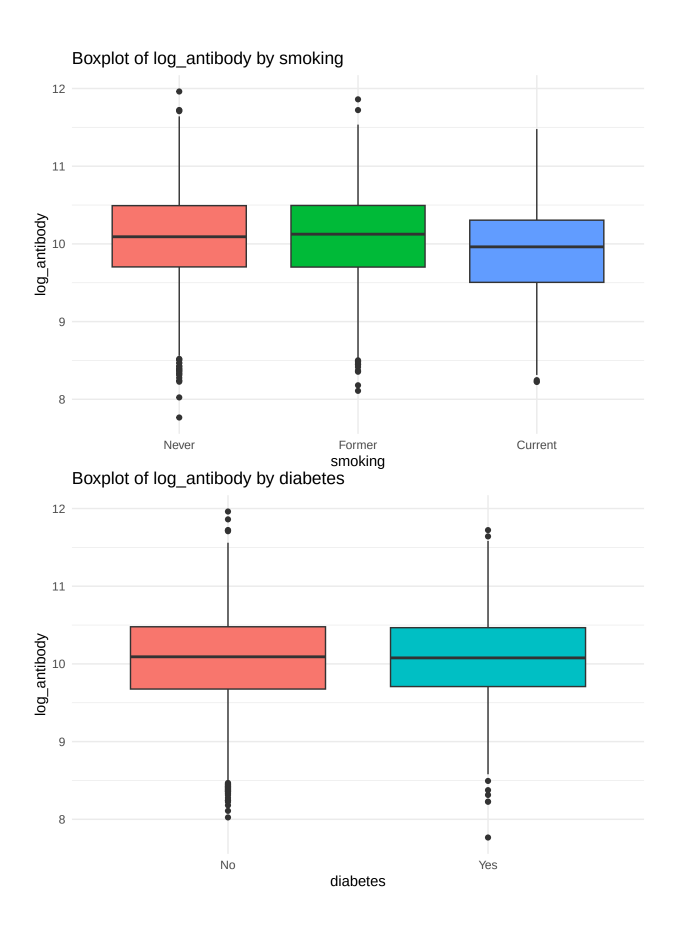
The VIF of bmi, weight and height exceed 10, indicating serious multicollinearity among these variables. Since BMI is a function of weight and height, it is recommended to retain only one of them (e.g., BMI) in the model to avoid redundancy and unstable coefficient estimates.

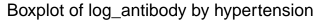
Then for categorical variable, generate boxplots to visualize the distribution of log\_antibody across levels of each categorical variable.

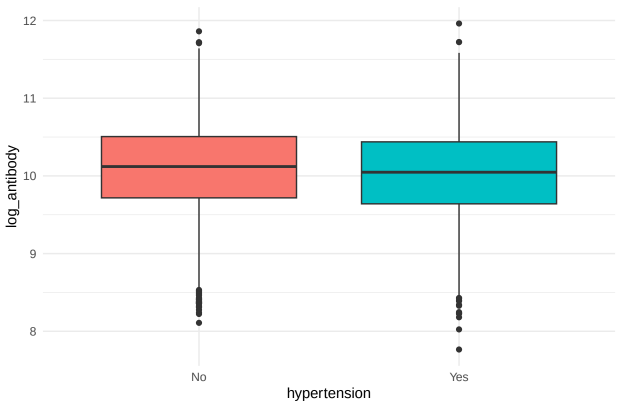
```
categorical_name <- c("gender", "race", "smoking", "diabetes", "hypertension")

for (name in categorical_name) {
   p <- ggplot(dat1, aes_string(x = name, y = "log_antibody", fill = name)) +
        geom_boxplot() +
        ggtitle(paste("Boxplot of log_antibody by", name)) +
        theme_minimal() +
        theme(legend.position = "none")
        print(p)
}</pre>
```









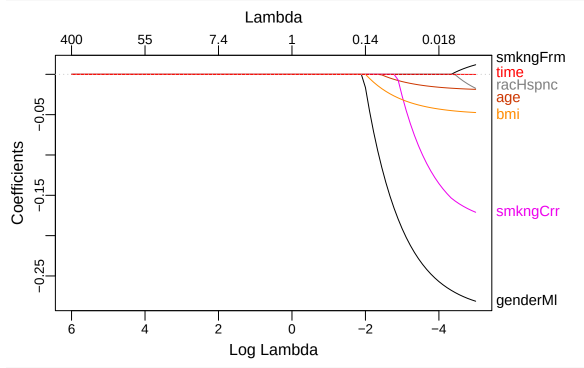
The boxplots suggest that the distribution of log\_antibody does not differ substantially across the categories of each categorical variable, indicating limited evidence of strong group-level effects.

# **Model Training**

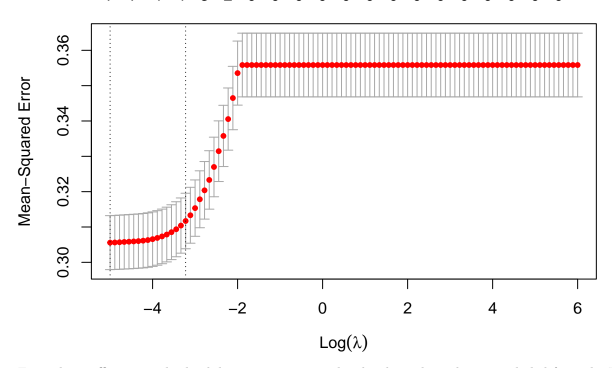
We first fit a Lasso regression model to select important predictors and address multicollinearity. This is particularly useful here, as previous VIF analysis indicated strong multicollinearity among BMI, height, and weight.

```
## genderMale
                   -0.281300893
## raceAsian
## raceBlack
## raceHispanic
                   -0.017347987
## smokingFormer
                    0.011848697
## smokingCurrent
                  -0.171036138
## height
## weight
## bmi
                   -0.047381778
## diabetesYes
## hypertensionYes
## SBP
## LDL
## time
                   -0.000131821
```

## plot\_glmnet(cv.lasso\$glmnet.fit)



plot(cv.lasso)



From the coefficients under lambda.min, we can see that height and weight are excluded from the Lasso model and bmi is retained. The lasso model helps solve multicollinearity.

```
y_pred_train <- predict(cv.lasso, newx = x, s = "lambda.min")

rmse_train <- sqrt(mean((y_pred_train - y)^2))

print(paste("RMSE on training set (lambda.min):", round(rmse_train, 4)))</pre>
```

## [1] "RMSE on training set (lambda.min): 0.5518"

Then we use the test dataset (dat2) to compute the test RMSE of the lasso model and evaluate model generalizability.

```
x_test <- model.matrix(log_antibody ~ ., data = dat2)[, -1]
y_test <- dat2$log_antibody

y_pred_lasso <- predict(cv.lasso, newx = x_test, s = "lambda.min")

rmse_lasso <- sqrt(mean((y_pred_lasso - y_test)^2))

print(paste("RMSE on test set (lambda.min):", round(rmse_lasso, 4)))</pre>
```

## [1] "RMSE on test set (lambda.min): 0.5749"

Since Lasso is a linear model that assumes additive and linear relationships between predictors and the outcome, we next explore two nonlinear modeling approaches (GAM and MARS) to capture potential nonlinearities and interaction effects in the data.

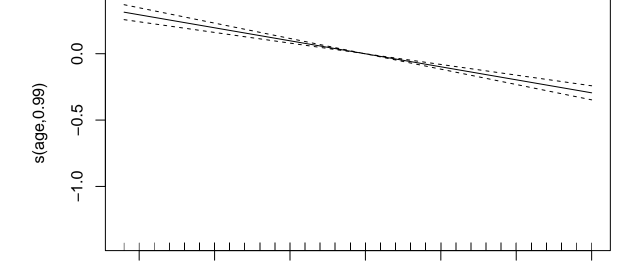
```
# Define trainControl
ctrl1 <- trainControl(method = "cv", number = 10)
train_y <- dat1$log_antibody</pre>
```

```
train_x <- dat1 %>%
  select(-log_antibody)
# GAM model
set.seed(2)
gam.fit <- train(train_x, train_y,</pre>
               method = "gam",
                trControl = ctrl1)
gam.fit$bestTune
    select method
      TRUE GCV.Cp
gam.fit$finalModel
##
## Family: gaussian
## Link function: identity
##
## Formula:
## .outcome ~ gender + diabetes + hypertension + smoking + race +
##
      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
summary(gam.fit$finalModel)
##
## Family: gaussian
## Link function: identity
##
## Formula:
## .outcome ~ gender + diabetes + hypertension + smoking + race +
##
      s(age) + s(SBP) + s(LDL) + s(bmi) + s(time) + s(height) +
##
      s(weight)
##
## Parametric coefficients:
##
                  Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                 ## genderMale
                ## diabetesYes
                 0.014230 0.020640 0.689
                                               0.491
## hypertensionYes -0.007678 0.015995 -0.480
                                               0.631
## smokingFormer 0.022219 0.016660 1.334
                                               0.182
## smokingCurrent -0.193175 0.025834 -7.478 8.9e-14 ***
## raceAsian
                -0.003296 0.033009 -0.100
                                               0.920
                -0.010509
                            0.018837 -0.558
## raceBlack
                                               0.577
## raceHispanic -0.037424
                            0.026176 -1.430
                                               0.153
## 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)
            4.179e+00
                           9 41.897
                                     <2e-16 ***
            7.892e+00
                           9 44.960
                                     <2e-16 ***
## s(time)
## s(height) 1.234e+00
                           9 0.278
                                      0.121
## s(weight) 2.262e-06
                           9 0.000
                                      0.666
## 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
                                      n = 5000
plot(gam.fit$finalModel)
```

50

45



60

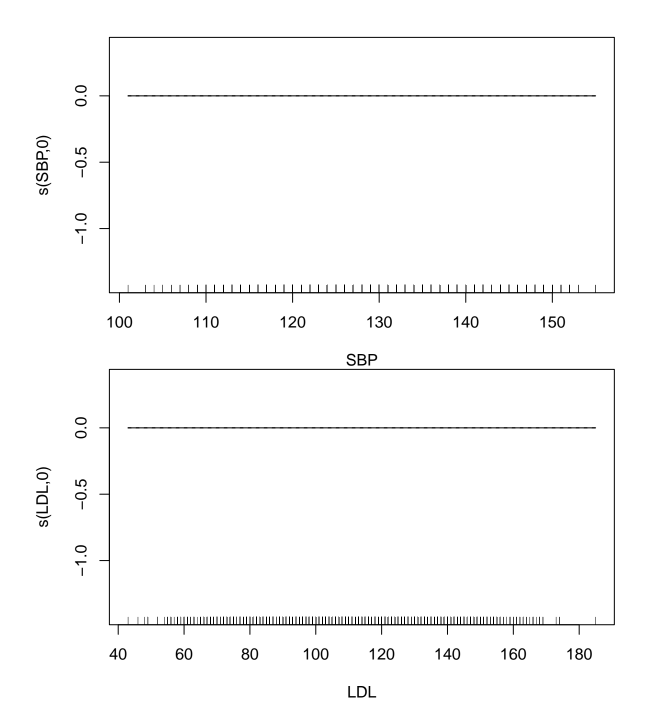
age

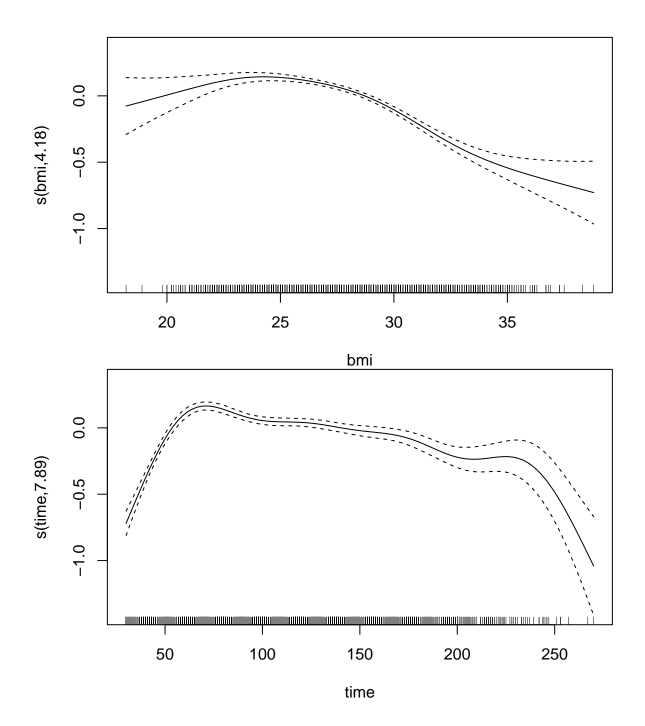
65

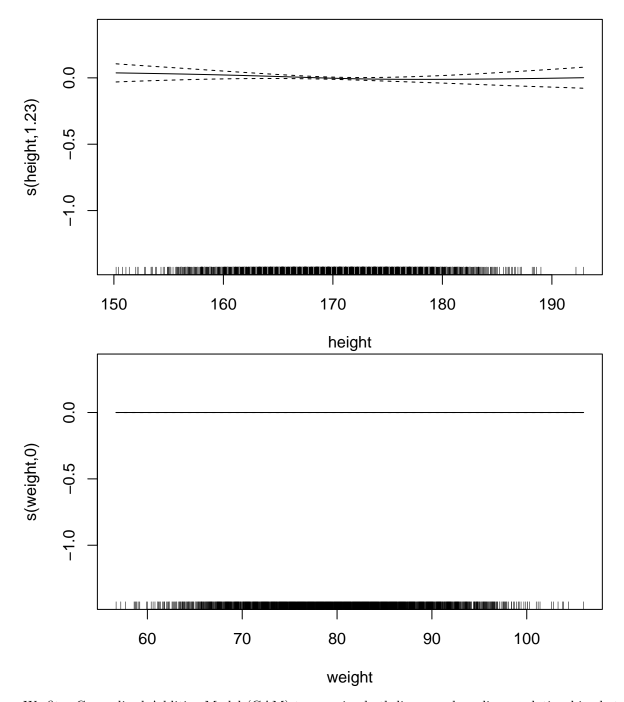
70

75

55





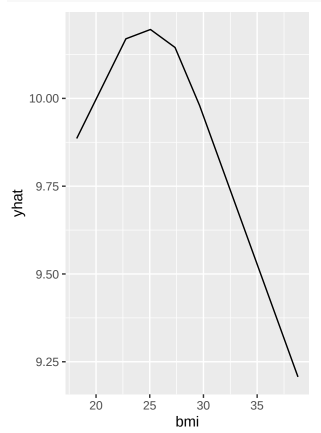


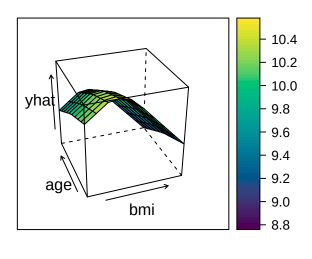
We fit a Generalized Additive Model (GAM) to examine both linear and nonlinear relationships between antibody levels and a set of demographic, clinical, and time-related predictors. The model achieved an adjusted R<sup>2</sup> of 0.22 and explained 22.4% of the deviance, indicating a moderate fit that captures key patterns in the data. Among the categorical predictors, male gender and current smoking were significantly associated with lower antibody responses, while race, diabetes, and hypertension did not show statistically significant effects. Several continuous variables exhibited nonlinear effects: notably, BMI and time since vaccination demonstrated strong and statistically significant nonlinear associations with antibody levels. Antibody responses peaked at a moderate BMI and declined at higher levels, while time since vaccination showed a characteristic waning immunity curve, with predicted antibody levels decreasing nonlinearly over time. These findings support the relevance of GAM for capturing complex decay behavior in vaccine-induced immune responses against both demographic and clinical factors.

```
# MARS model
mars_grid <- expand.grid(degree = 1:3,</pre>
                           nprune = 2:15)
set.seed(2)
mars.fit <- train(train_x, train_y,</pre>
                   method = "earth",
                   tuneGrid = mars_grid,
                   trControl = ctrl1)
ggplot(mars.fit)
   0.58 -
   0.57 -
RMSE (Cross-Validation)
                                                                              Product Degree
   0.56 -
   0.55 -
  0.54 -
  0.53 -
                                                        12
                                      8
                   4
                                     #Terms
mars.fit$bestTune
     nprune degree
## 8
           9
coef(mars.fit$finalModel)
##
       (Intercept)
                      h(27.8-bmi)
                                        h(time-57)
                                                        h(57-time)
                                                                         genderMale
     10.847446930
                                      -0.002254182
                                                     -0.033529326
                                                                       -0.296290451
##
                     -0.061997354
##
        h(age-59)
                         h(59-age) smokingCurrent
                                                       h(bmi-23.7)
##
     -0.022957648
                       0.016138468
                                      -0.205126851
                                                      -0.084380175
p1 <- pdp::partial(mars.fit, pred.var = c("bmi"), grid.resolution = 10) |> autoplot()
p2 <- pdp::partial(mars.fit, pred.var = c("bmi", "age"),</pre>
                     grid.resolution = 10) |>
      pdp::plotPartial(levelplot = FALSE, zlab = "yhat", drape = TRUE,
```

screen = list(z = 20, x = -60))

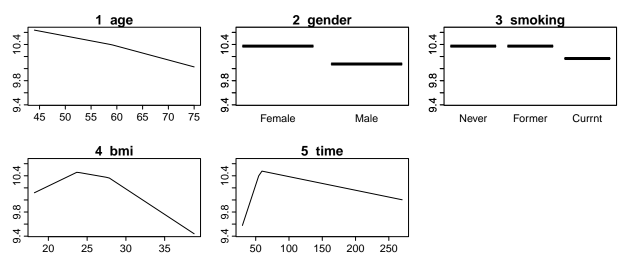
#### gridExtra::grid.arrange(p1, p2, ncol = 2)





```
plotmo(mars.fit,
nresponse = 1,
degree2 = FALSE,
varnames = "age")
```

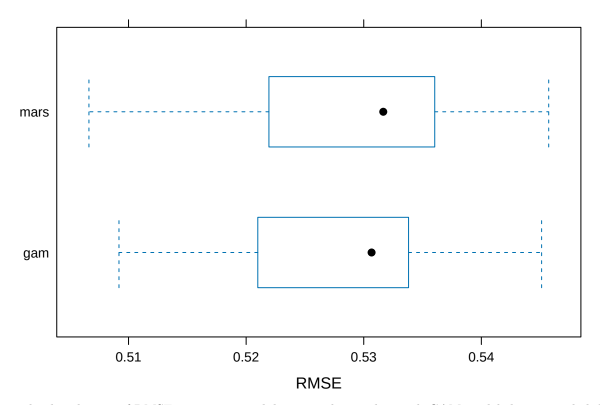
type=raw train.default(x=train\_x, y=train\_y, method="earth", trControl=ctr...



### Model selection

To select the best model, we can compare the cross-validated metrics of the MARS and GAM models using resampling, this helps evaluate and visualize the relative performance of the two models.

```
resamp <- resamples(list(mars = mars.fit, gam = gam.fit))</pre>
summary(resamp)
##
## Call:
  summary.resamples(object = resamp)
##
## Models: mars, gam
## Number of resamples: 10
##
## MAE
                                            Mean
##
             Min.
                    1st Qu.
                                Median
                                                    3rd Qu.
                                                                 Max. NA's
## mars 0.4120189 0.4180233 0.4203065 0.4224208 0.4285348 0.4360995
                                                                         0
       0.4127242 0.4190074 0.4202804 0.4224455 0.4273258 0.4352565
                                                                          0
##
##
## RMSE
                    1st Qu.
##
                                Median
                                                                 Max. NA's
             Min.
                                            Mean
                                                    3rd Qu.
## mars 0.5066327 0.5230870 0.5316602 0.5282995 0.5354905 0.5457286
                                                                         0
##
   gam 0.5091877 0.5223781 0.5306669 0.5279212 0.5336806 0.5451253
                                                                          0
##
## Rsquared
             Min.
                    1st Qu.
                                Median
                                            Mean
                                                    3rd Qu.
                                                                 Max. NA's
## mars 0.1766328 0.1941155 0.2028183 0.2159220 0.2369173 0.2730827
                                                                          0
  gam 0.1795042 0.1955026 0.2071224 0.2170568 0.2376473 0.2735385
bwplot(resamp, metric = "RMSE")
```



The distribution of RMSE across cross-validation is also similar, with GAM model showing a slightly lower median RMSE compared to MARS model.

Moreover, both models achieve lower average RMSE on the training set (dat1) compared to the Lasso model (Lasso model train RMSE: 0.5518, test RMSE: 0.5749), suggesting better performance. This may be attributed to the presence of nonlinear relationships between the response variable (log\_antibody) and some predictors (such as bmi and time), which cannot be effectively captured by the linear structure of Lasso regression.

We should also evaluate the generalizability of the trained MARS and GAM models by computing RMSE on the test set (dat2).

```
# test RMSE of MARS model
mars.pred <- predict(mars.fit, newdata = dat2)
mars_test_rmse = sqrt(mean((mars.pred - dat2[, "log_antibody"])^2))
print(paste("RMSE on test set (MARS model):", round(mars_test_rmse, 4)))
## [1] "RMSE on test set (MARS model): 0.5328"
# test RMSE of GAM model
gam.pred <- predict(gam.fit, newdata = dat2)
gam_test_rmse = sqrt(mean((gam.pred - dat2[, "log_antibody"])^2))
print(paste("RMSE on test set (GAM model):", round(gam_test_rmse, 4)))</pre>
```

```
## [1] "RMSE on test set (GAM model): 0.5701"
```

Both MARS and GAM models show very similar performance in cross-validation, with nearly identical mean RMSE values (0.5283 vs 0.5279).

However, on the test set, the MARS model achieves a lower RMSE (0.5328) compared to the GAM model (0.5701), suggesting better generalization.

Given this gap in test performance while maintaining comparable training performance, the MARS model appears to be the better choice in this case.

### Influence of Demographic and Clinical Factors on Antibody Responses

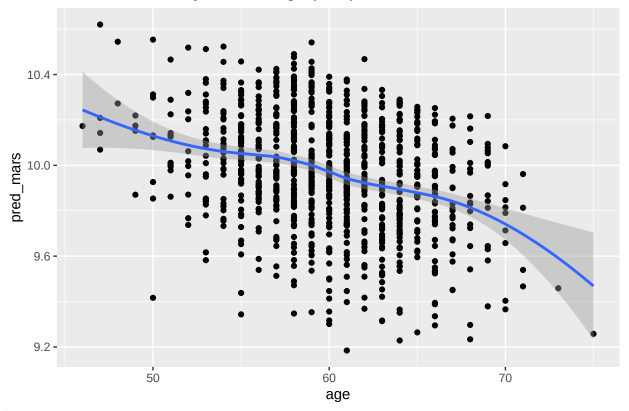
```
dat2$pred mars <- predict(mars.fit, newdata = dat2)</pre>
dat2 %>%
  group_by(gender) %>%
  summarise(mean_pred = mean(pred_mars), sd_pred = sd(pred_mars))
## # A tibble: 2 x 3
##
     gender mean_pred sd_pred
##
     <fct>
                <dbl>
                         <dbl>
## 1 Female
                10.1
                         0.215
## 2 Male
                 9.83
                         0.204
dat2 %>%
  group_by(smoking) %>%
  summarise(mean_pred = mean(pred_mars), sd_pred = sd(pred_mars))
## # A tibble: 3 x 3
##
     smoking mean_pred sd_pred
##
     <fct>
                 <dbl>
                          <dbl>
## 1 Never
                  9.99
                          0.238
                          0.260
## 2 Former
                  9.98
                          0.222
## 3 Current
                  9.80
```

For categorical predictors, the predicted antibody levels from the MARS model on dat2 are consistent with the trends observed during model training on dat1. Specifically, females show higher predicted antibody levels than males (10.10 vs. 9.83), reflecting potential gender-related differences in immune response. Similarly, current smokers exhibit the lowest predicted antibody levels (mean = 9.80) compared to former and never smokers (means = 9.98 and 9.99, respectively), aligning with immunological evidence that smoking impairs vaccine response. These results reinforce the reliability of the model and suggest meaningful differences in antibody responses across demographic and behavioral subgroups.

```
# 1. PDP-like plots in dat2 using predicted values
ggplot(dat2, aes(x = age, y = pred_mars)) +
  geom_point() +
  geom_smooth(method = "loess") +
  labs(title = "Predicted Antibody Level vs Age (dat2)")
```

```
## `geom_smooth()` using formula = 'y ~ x'
```

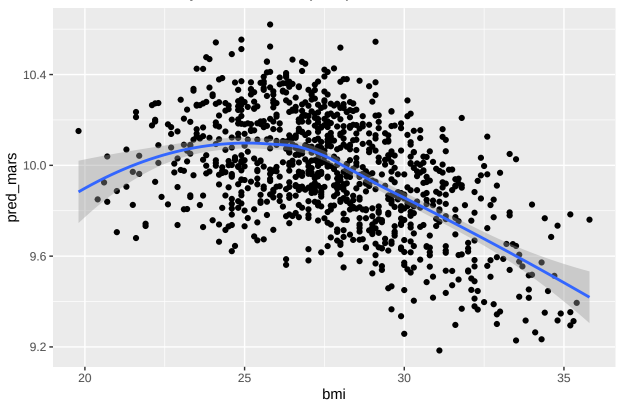
# Predicted Antibody Level vs Age (dat2)



```
ggplot(dat2, aes(x = bmi, y = pred_mars)) +
  geom_point() +
  geom_smooth(method = "loess") +
  labs(title = "Predicted Antibody Level vs BMI (dat2)")
```

## `geom\_smooth()` using formula = 'y ~ x'

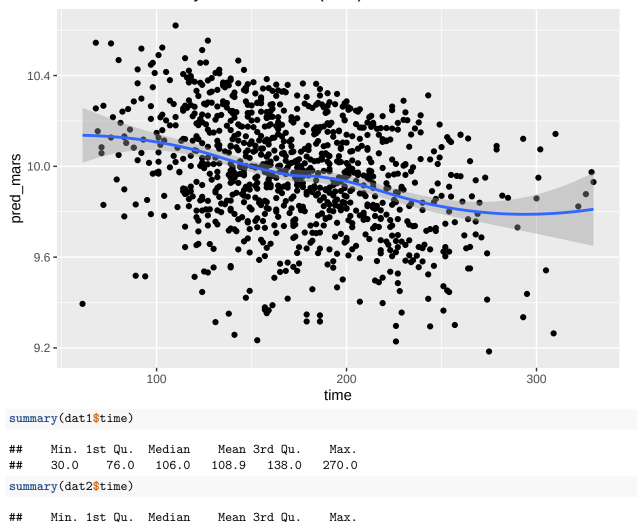
# Predicted Antibody Level vs BMI (dat2)



```
ggplot(dat2, aes(x = time, y = pred_mars)) +
  geom_point() +
  geom_smooth(method = "loess") +
  labs(title = "Predicted Antibody Level vs Time (dat2)")
```

##  $geom_smooth()$  using formula = 'y ~ x'

### Predicted Antibody Level vs Time (dat2)



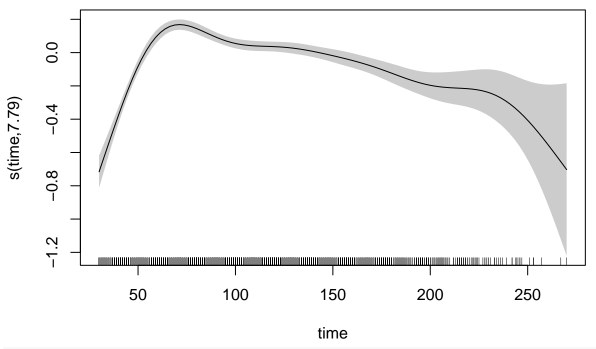
```
## 61.0 140.0 171.0 173.8 205.0 330.0

For continuous varibles, we observed consistent trends in the marginal effects of BMI and age on predicted antibody levels between dat1 and dat2, indicating stable model behavior across datasets. However, the relationship between time since vaccination and predicted antibody levels differs noticeably: while dat1 shows an initial sharp rise followed by a gradual decline (consistent with typical post-vaccination antibody dynamics), the prediction on dat2 displays a more uniformly decreasing trend. Upon examining the distribution of
```

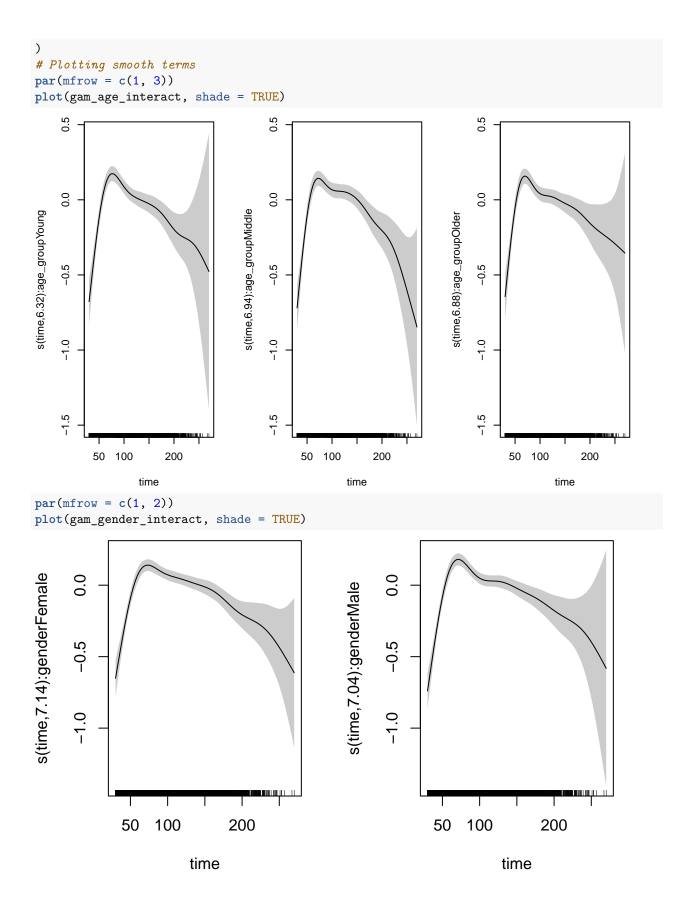
an initial sharp rise followed by a gradual decline (consistent with typical post-vaccination antibody dynamics), the prediction on dat2 displays a more uniformly decreasing trend. Upon examining the distribution of the time variable, we found that this shift in behavior is due to the absence of early time points in dat2 — specifically, no observations exist for time < 61. As a result, the model cannot capture the early rise phase in dat2, and instead predicts a smoother, gradual decline.

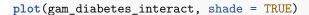
#### How Antibody Levels Change Over Time

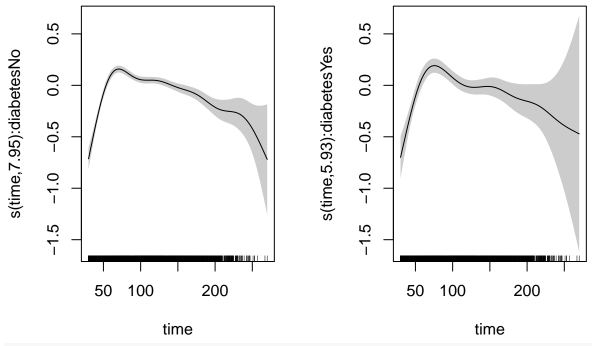
```
dat1_ageGroup <- dat1 %>%
  mutate(age_group = ntile(age, 3)) %>%
  mutate(age_group = factor(age_group, labels = c("Young", "Middle", "Older"))) %>%
  mutate(
    gender = factor(gender, labels = c("Female", "Male")),
    diabetes = factor(diabetes, labels = c("No", "Yes")),
    hypertension = factor(hypertension, labels = c("No", "Yes"))
```



```
library(mgcv)
gam_age_interact <- gam(</pre>
  log_antibody ~ s(time, by = age_group) + age_group + gender + bmi + SBP + LDL +
    race + smoking + diabetes + hypertension,
  data = dat1_ageGroup,
  method = "GCV.Cp"
gam_gender_interact <- gam(</pre>
  log_antibody ~ s(time, by = gender) + gender + age + bmi + SBP + LDL +
    race + smoking + diabetes + hypertension,
  data = dat1_ageGroup,
  method = "GCV.Cp"
)
gam_diabetes_interact <- gam(</pre>
  log_antibody ~ s(time, by = diabetes) + diabetes + age + bmi + SBP + LDL +
    race + gender + smoking + hypertension,
  data = dat1_ageGroup,
  method = "GCV.Cp"
gam_hypertension_interact <- gam(</pre>
  log_antibody ~ s(time, by = hypertension) + diabetes + age + bmi + SBP + LDL +
    race + gender + smoking + hypertension,
  data = dat1_ageGroup,
  method = "GCV.Cp"
```







plot(gam\_hypertension\_interact, shade = TRUE)

