Midterm Exam 2

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Due Friday, November 22, 2024 at 1:00pm

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- This exam consists of 4 problems and 5 pages. The exam is worth 100 points (each question is worth 25 points) with the opportunity to earn 2 additional bonus points.
- Before submitting the exam, read and sign the statement below confirming that you have not cheated on this exam. A digital signature is fine.
- Some parts of a question may require the answer to an earlier part. If you cannot solve the earlier part, you can still receive partial credit for the later parts; make up a reasonable answer for the earlier part to use in subsequent parts of the problem.
- Show your work and explain your reasoning; the final answer is not as important as the process by which you arrived at that answer. We can more easily give partial credit if you have written out your steps clearly.
- This exam is open book and open notes, and you may use the internet. But there can be no discussing the exam with other people, including people who are not in this class.

By signing below, I confirm that I have not cheated while taking this exam: I have not used any unauthorized resources nor copied another student's responses.

Signature: Matthew Krasnow

Problem Scoring		
Problem	Point Value	Points Scored
1	25	
2	25	
3	25	
4	25	
Total	100	

Problem 1: Warm-up [25 points]

Your collaborator is investigating the relationship between educational attainment and cognitive ability. She has collected data on a random sample of adults in Massachusetts in the dataset pl.csv. The variable COG encodes a continuous measure of cognitive ability designed to have a mean around 50 (higher scores are associated with higher cognitive ability). Educational attainment refers to the highest level of education an individual has completed, and is encoded in the EDUC variable as follows:

0: Primary School (grade 5)

1: Junior High School (grade 8)

2: High School (grade 12)

3: College degree or higher

Your collaborator asks you to fit a linear regression to predict cognitive ability from educational attainment.

(a) Write the formal statistical model you intend to fit.

I will use a linear regression model where $EDUC_i$ is treated as a categorical variable (ordinal) with four levels. I will define indicator variables to represent the education levels. \setminus

\$\$

```
D1_i = 1 if EDUC_i = 1, 0 otherwise (middle school) \\
D2_i = 1 if EDUC_i = 2, 0 otherwise (high schoole) \\
D3_i = 1 if EDUC_i = 2, 0 otherwise (college or higher ) \\
$$
```

Thus the formal statistical model is:

$$COG_{i} = \beta_{0} + \beta_{1} * D1 + \beta_{2} * D2 + \beta * D3 + \epsilon_{i}$$

Define

 β_0 = mean cognative ability for people with primary school edu \setminus

 β_1 = mean cognative ability for people with middle school edu\

 β_2 = mean cognative ability for people with high school edu \

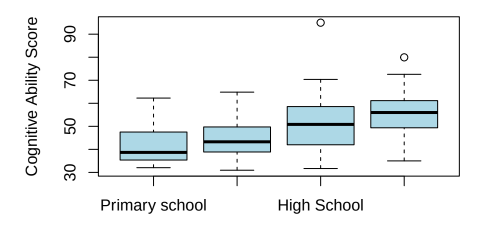
 β_3 = mean cognative ability for people with college or higher education \backslash

 ϵ is the independent error terms. I would like to say that these are $\sim N(0, \sigma^2)$, but we have not properly evaluated our assumptions yet.

For this model to be valid, we need to evaluate our ELI H assumptions, but we have not done that yet

(b) Create an appropriate plot to visualize the association between educational attainment and cognitive ability.

Cognitive Ability by Educational Attainment



Educational Attainment

I chose a boxplot for this question because it demonstrates the spread of the data where the data is separated into categories effectively.

(c) Fit the linear regression that you specified in part (a) and interpret all the mean parameter estimates.

```
model <- lm(COG ~ EDUC, data = data)
summary(model)
##
## Call:</pre>
```

```
## Call.
## lm(formula = COG ~ EDUC, data = data)
##
## Residuals:
## Min 1Q Median 3Q Max
## -21.06 -5.82 -0.78 5.77 43.91
##
## Coefficients:
## Estimate Std. Error t value Pr(>|t|)
```

```
## (Intercept)
                                42.15
                                            2.65
                                                   15.88 < 2e-16 ***
## EDUCJunior High School
                                 2.10
                                            2.91
                                                    0.72
                                                           0.4709
## EDUCHigh School
                                 8.94
                                            2.92
                                                    3.07
                                                           0.0025 **
## EDUCCollege edu or higher
                                                    4.84 2.6e-06 ***
                                13.90
                                            2.87
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 9.19 on 196 degrees of freedom
## Multiple R-squared: 0.244, Adjusted R-squared: 0.232
## F-statistic: 21.1 on 3 and 196 DF, p-value: 7.08e-12
```

Interpretaion: $\beta_0 = 42.148$ on average, individuals with only primary school education have a COG of 42.148

 $\beta_1 = 2.100$ On average, individuals with only junior high school education have a COG that is 2.1 points higher than those with only a primary school education (44.248)

 $\beta_2 = 8.937$ On average, individuals with only high school education have a COG that is 8.937 points higher than those with only a primary school education (42.148+8.937)

 $\beta_3 = 13.902$ On average, individuals with college education or higher have a COG that is 13.902 points higher than those with only a primary school education (42.148+13.902)

(d) Based on your plot, you might be tempted to use a linear term to model the association between educational attainment and cognitive ability instead, retaining the original coding scheme (that is, treating EDUC as a continuous variable). Statistically test whether this would be a better model. Comment on whether you think this is a better approach.

```
##
## Call:
## lm(formula = COG ~ EDUC_num, data = data)
##
## Residuals:
## Min 1Q Median 3Q Max
## -21.05 -6.44 -1.04 5.70 44.38
```

```
##
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
##
                  39.77
                              1.48
                                      26.83
                                             < 2e-16 ***
## (Intercept)
## EDUC num
                   5.42
                              0.69
                                       7.86
                                            2.4e-13 ***
## ---
## Signif. codes:
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 9.18 on 198 degrees of freedom
## Multiple R-squared: 0.238, Adjusted R-squared:
## F-statistic: 61.8 on 1 and 198 DF, p-value: 2.42e-13
#compare models using an F-test
anova(model2, model) # model is categorical
## Analysis of Variance Table
## Model 1: COG ~ EDUC num
## Model 2: COG ~ EDUC
##
     Res.Df
              RSS Df Sum of Sq
                                  F Pr(>F)
## 1
        198 16700
## 2
        196 16565
                           134 0.79
                                       0.45
```

The F test demonstrates that the data does not imply a statistically significant difference between the categorical model and the continuous model. This indicates that the null hypothesis that the two models are the same should be retained (not accepted) until further data may show that one is better than the other.

Intuitively, I do not think it makes sense to treat education as a continuous scale. The scale does not accurately reflect the actual learning changes that occur between stages—I do not think that partial education is a linear scale – going to half of college or high school is not equally as rewarding as half of total college or high school. This should be treated as categorical.

(e) Suppose instead that you retained the original coding scheme (treating EDUC as a continuous variable) but fit a third degree polynomial (with all lower order terms included as well). Would this improve the fit of the model in part c)? What if you fit an even higher order polynomial?

```
model_poly3 <- lm(COG ~ EDUC_num + I(EDUC_num^2) + I(EDUC_num^3), data = data)
summary(model_poly3)</pre>
```

```
##
## Call:
## lm(formula = COG ~ EDUC_num + I(EDUC_num^2) + I(EDUC_num^3),
       data = data)
##
##
## Residuals:
##
      Min
              1Q Median
                             3Q
                                   Max
## -21.06 -5.82 -0.78
                           5.77
                                 43.91
##
```

```
## Coefficients:
##
                 Estimate Std. Error t value Pr(>|t|)
                   42.148
                                2.654
                                        15.88
                                                <2e-16 ***
## (Intercept)
## EDUC_num
                   -2.470
                                6.306
                                        -0.39
                                                  0.70
## I(EDUC num^2)
                    5.672
                                4.688
                                         1.21
                                                  0.23
## I(EDUC num^3)
                   -1.101
                                0.972
                                        -1.13
                                                  0.26
## ---
## Signif. codes:
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 9.19 on 196 degrees of freedom
## Multiple R-squared: 0.244, Adjusted R-squared: 0.232
## F-statistic: 21.1 on 3 and 196 DF, p-value: 7.08e-12
anova(model, model_poly3)
## Analysis of Variance Table
##
## Model 1: COG ~ EDUC
## Model 2: COG ~ EDUC_num + I(EDUC_num^2) + I(EDUC_num^3)
              RSS Df Sum of Sq F Pr(>F)
     Res.Df
## 1
        196 16565
## 2
        196 16565
                  0 -3.27e-11
```

Based on this, - the polynomial model does not improve uplon the categorical model from part c - the categorical model is better because it 1. makes more sense in the situation 2. there is no significant evidence that the poly is better – we would rather use a simpler model - same RSS and degrees of freedom, they explain the variance in COG equally well - adding new polynomials would not increase the fit because we have already observed polynomial terms which are not significant—adding more will not help

BONUS (2 points): Which of the assumptions of linear regression would you have to test for the model you fit in part c)? Test them and comment on the validity of that model. Is there anything else you should make your collaborator aware of?

We need to test our ELI H assumptions:

Existence

• we standardly assume that the mean exists. This implies that there is finite variance of the residuals. Otherwise, our model would not converge.

Linearity

Single EDUC is categorical, the linearity assumption pertains to peroperly specifying the model to include all categories. We have included all the categories in EDUC, so we are good (assumptino satisfied)

Independence of Errors

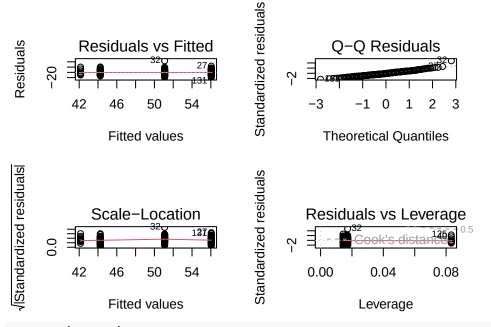
- this depends on the study design, so we don't really know based on this information
- to meet this assumption, we would need to see an SRS and no clustering or repeated measures

• if there is any reason to assume that the values of datapoints affects other datapoints, we cannot meet this assumption

Homoskedasticity

• use the following test:

```
par(mfrow=c(2,2))
plot(model)
```



library(lmtest)

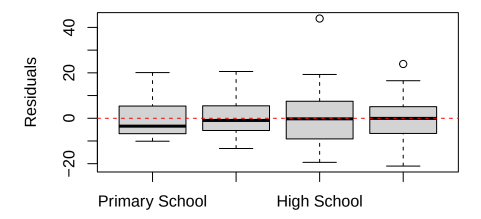
```
## Loading required package: zoo
##
## Attaching package: 'zoo'
## The following objects are masked from 'package:base':
##
## as.Date, as.Date.numeric
bptest(model)
```

```
##
## studentized Breusch-Pagan test
##
## data: model
## BP = 7.1, df = 3, p-value = 0.07
```

- at the = 0.05 value, this is NOT. a statistically significant finding. Thus we do not have fail to reject the null hypothesis
- we do not have data to state that the data is heteroskedastic

Normality

Residuals by Education Level



Education Level

- At the 0.05 level, we would

reject the null hypothesis. This indicates that the residuals are NOT normally distributed.

Since our E.L. assumptions are met, if we assume linearity, we can use our linear model to estimate the mean parameters. however, we cannot do standard inference with this data because the residuals are not distributed normally.

I would want a collaborator to know that we did not formally check for independence and that this should be done before applying this data to a real world scenario.

Problem 2: Open-ended application [25 points]

You have another collaborator who has asked you to help him develop a model to estimate the associations between various predictors and systolic blood pressure (SBP) in a random sample of American adults aged at least 50. For this purpose, he has curated a dataset (p2.csv) which contains the following variables:

```
SBP: Systolic blood pressure, in mmHg. 

AGE: Age in years. 

GENDER: Gender indicator; 0 = \text{Male}, 1 = \text{Female}. 

EDUC: Highest level of education attained. 0 = \text{Primary School}, 1 = \text{Junior High}, 2 = \text{High School}, 3 = \text{College degree or higher}. 

DM: Diabetes mellitus indicator; 0 = \text{No}, 1 = \text{Yes}. 

SMOKE: Smoking status indicator; 0 = \text{No}, 1 = \text{Yes}. 

BMI: Body mass index (\text{kg/m}^2). 

STATIN: Statin use indicator; 0 = \text{No}, 1 = \text{Yes}.
```

(a) Your collaborator does not have a model in mind, but he would like you to do some exploratory analyses, as he believes many of the variables he has collected are important. Used principled model selection procedure(s) to arrive at what you believe to be an appropriate model. For simplicity, do not consider any polynomial terms in this problem. Start by making sure the categorical variables are coded as factors.

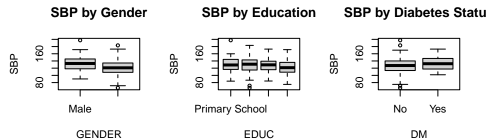
```
library(dplyr)
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:stats':
##
##
       filter, lag
## The following objects are masked from 'package:base':
##
##
       intersect, setdiff, setequal, union
library(ggplot2)
library(corrplot)
## Warning: package 'corrplot' was built under R version 4.3.3
## corrplot 0.95 loaded
data <- read.csv("data/p2.csv")</pre>
# categorical variables => factors with appropriate labels
data$GENDER <- factor(data$GENDER, levels = c(0, 1), labels = c("Male", "Female"))</pre>
```

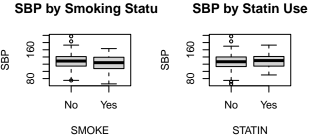
data\$EDUC <- factor(data\$EDUC, levels = c(0, 1, 2, 3),</pre>

```
labels = c("Primary School", "Junior High", "High School", "College or high
data$DM <- factor(data$DM, levels = c(0, 1), labels = c("No", "Yes"))</pre>
data$SMOKE <- factor(data$SMOKE, levels = c(0, 1), labels = c("No", "Yes"))</pre>
data$STATIN <- factor(data$STATIN, levels = c(0, 1), labels = c("No", "Yes"))
str(data)
## 'data.frame':
                    314 obs. of 8 variables:
           : int 55 65 68 70 53 64 70 75 66 64 ...
## $ GENDER: Factor w/ 2 levels "Male", "Female": 2 2 2 1 1 1 1 1 1 2 ...
## $ EDUC : Factor w/ 4 levels "Primary School",..: 3 2 3 3 1 2 4 2 1 2 ...
            : Factor w/ 2 levels "No", "Yes": 2 1 1 1 1 1 1 1 1 1 ...
## $ SMOKE : Factor w/ 2 levels "No", "Yes": 1 1 1 1 1 1 1 1 2 1 ...
## $ BMI
          : num 39.7 29 22.3 32.4 29.2 ...
## $ SBP
           : num 108 124 116 166 159 ...
## $ STATIN: Factor w/ 2 levels "No", "Yes": 1 2 1 1 1 1 1 1 2 1 ...
summary(data)
##
         AGE
                      GENDER
                                               EDUC
                                                          DM
                                                                  SMOKE
## Min.
           :50.0
                   Male :175
                                Primary School
                                                 : 47
                                                        No :284
                                                                  No :242
## 1st Qu.:54.0
                   Female:139
                                Junior High
                                                 :120
                                                        Yes: 30
                                                                  Yes: 72
## Median :61.0
                                High School
                                                 : 66
                                College or higher: 81
## Mean
          :61.8
## 3rd Qu.:69.0
## Max.
           :81.0
##
        BMI
                        SBP
                                   STATIN
          :18.4
                   Min. : 65.3
                                   No :214
## Min.
## 1st Qu.:24.4
                   1st Qu.:113.9
                                  Yes:100
## Median :26.6
                   Median :128.3
## Mean
          :27.4
                   Mean :127.7
                   3rd Qu.:140.3
## 3rd Qu.:29.8
## Max.
           :61.0
                   Max.
                         :197.5
#EDA
# correlation matrix for continuous variables
continuous_vars <- data[c("SBP", "AGE", "BMI")]</pre>
cor_matrix <- cor(continuous_vars)</pre>
corrplot(cor_matrix, method = "number")
```



```
# boxplots for categorical variables
par(mfrow = c(2, 3))
boxplot(SBP ~ GENDER, data = data, main = "SBP by Gender")
boxplot(SBP ~ EDUC, data = data, main = "SBP by Education")
boxplot(SBP ~ DM, data = data, main = "SBP by Diabetes Status")
boxplot(SBP ~ SMOKE, data = data, main = "SBP by Smoking Status")
boxplot(SBP ~ STATIN, data = data, main = "SBP by Statin Use")
par(mfrow = c(1, 1))
```





model selection

```
# fit the full model
full model <- lm(SBP ~ AGE + GENDER + EDUC + DM + SMOKE + BMI + STATIN, data = data)
# stepwise selection aic
step_model_aic <- step(full_model, direction = "both", trace = FALSE)</pre>
# stepwise selection bic
n <- nrow(data)</pre>
step_model_bic <- step(full_model, direction = "both", trace = FALSE,</pre>
                      k = log(n)) # Using BIC penalty
summary(step_model_aic)
##
## Call:
## lm(formula = SBP ~ AGE + GENDER + BMI, data = data)
##
## Residuals:
##
     Min
             1Q Median
                            3Q
                                  Max
## -45.38 -11.95 -1.35 12.79 56.64
##
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
##
                  48,403
                              9.700
                                       4.99 1.0e-06 ***
## (Intercept)
                                       7.46 8.7e-13 ***
## AGE
                   0.907
                              0.122
## GENDERFemale
                  -9.382
                              2.065
                                     -4.54 8.0e-06 ***
## BMI
                   0.996
                              0.225
                                       4.44 1.3e-05 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 17.9 on 310 degrees of freedom
## Multiple R-squared: 0.267, Adjusted R-squared: 0.26
## F-statistic: 37.7 on 3 and 310 DF, p-value: <2e-16
summary(step_model_bic)
##
## Call:
## lm(formula = SBP ~ AGE + GENDER + BMI, data = data)
##
## Residuals:
##
      Min
              1Q Median
                            3Q
                                  Max
## -45.38 -11.95 -1.35 12.79 56.64
##
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
                              9.700
                                       4.99 1.0e-06 ***
## (Intercept)
                  48.403
## AGE
                   0.907
                              0.122
                                       7.46 8.7e-13 ***
```

```
## GENDERFemale
                                    2.065
                     -9.382
                                              -4.54
                                                      8.0e-06 ***
##
   BMI
                      0.996
                                    0.225
                                               4.44
                                                      1.3e-05 ***
##
                      0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
##
## Residual standard error: 17.9 on 310 degrees of freedom
## Multiple R-squared: 0.267, Adjusted R-squared:
## F-statistic: 37.7 on 3 and 310 DF, p-value: <2e-16
# Check model assumptions for the final chosen model
par(mfrow = c(2, 2))
plot(step_model_aic)
                                      Standardized residuals
                                                  Q-Q Residuals
Residuals
           Residuals vs Fitted
           110
                    130
                            150
                                                                 2
                                                                     3
               Fitted values
                                                 Theoretical Quantiles
/Standardized residuals
                                      Standardized residuals
                                              Residuals vs Leverage
            Scale-Location
            110
                    130
                            150
                                               0.00
                                                         0.10
               Fitted values
                                                      Leverage
```

INTERPRETATION: - final model uses: age, gender, BMI to predict systolic BP. - $R^2 = 26\%$ of the variation in SBP is explained by this model - females have lower SBP - higher age => higher SBP - higher BMI => higher SBP - other variabeles are not statistically significant in this dataset ASSUMPTIONS:

Existence: Assumed to be met so that the model can converge Linearity: The residuals vs fitted plot demonstratess a random scatter of points around the 0 line, indicating approximate linearity. Independenc: Since we do not hav eaccess to how to the data was collected, we cannot say for certain if the data points are independent or not. This is something to be cautious of. We would like to see an SRS and no dependencies. Since the problem states that it was a random sample of american adults, we shall say that this assumption was met. Homoskedasticity: We see a relatively constant spread of ressiduals across fitted values in the scale-location plot. This means that our homoskedasticity assumption is met. Normality of residuals: While we are not doing inference in this question, this is not necessary for this question. However, since the QQ plot closely follows the diagonal line, we can say that the residuals are approximately normally distributed.

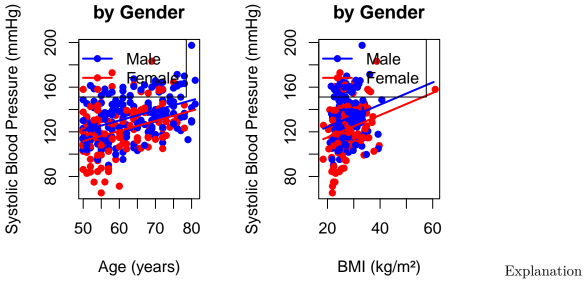
(b) Explain your final model to your collaborator. Remember, your collaborator is not a statistician, and some of the terms in your model might be difficult to explain. It might be helpful to explain

some of the relationships visually, and/or in particular subgroups. Every parameter should in some way be included in your interpretation.

```
# For Age effect
age_seq <- seq(min(data$AGE), max(data$AGE), by = 1)
pred_age_male <- data.frame(</pre>
  AGE = age_seq,
  GENDER = factor("Male", levels = c("Male", "Female")),
  BMI = mean(data$BMI)
pred_age_female <- data.frame(</pre>
  AGE = age_seq,
  GENDER = factor("Female", levels = c("Male", "Female")),
  BMI = mean(data$BMI)
)
# For BMI effect
bmi_seq <- seq(min(data$BMI), max(data$BMI), by = 1)</pre>
pred_bmi_male <- data.frame(</pre>
  AGE = mean(data\$AGE),
 GENDER = factor("Male", levels = c("Male", "Female")),
  BMI = bmi seq
)
pred bmi female <- data.frame(</pre>
  AGE = mean(data$AGE),
  GENDER = factor("Female", levels = c("Male", "Female")),
  BMI = bmi_seq
)
# predictions
pred_age_male$SBP <- predict(step_model_aic, pred_age_male)</pre>
pred_age_female$SBP <- predict(step_model_aic, pred_age_female)</pre>
pred_bmi_male$SBP <- predict(step_model_aic, pred_bmi_male)</pre>
pred_bmi_female$SBP <- predict(step_model_aic, pred_bmi_female)</pre>
par(mfrow = c(1, 2))
# Age effect plot
plot(data$AGE, data$SBP, pch = 16, col = ifelse(data$GENDER == "Male", "blue", "red"),
     xlab = "Age (years)", ylab = "Systolic Blood Pressure (mmHg)",
     main = "Effect of Age on Blood Pressure\nby Gender")
lines(pred_age_male$AGE, pred_age_male$SBP, col = "blue", lwd = 2)
lines(pred_age_female$AGE, pred_age_female$SBP, col = "red", lwd = 2)
legend("topleft", c("Male", "Female"), col = c("blue", "red"),
       pch = 16, lty = 1, lwd = 2)
# BMI effect plot
plot(data$BMI, data$SBP, pch = 16, col = ifelse(data$GENDER == "Male", "blue", "red"),
```

```
xlab = "BMI (kg/m²)", ylab = "Systolic Blood Pressure (mmHg)",
    main = "Effect of BMI on Blood Pressure\nby Gender")
lines(pred_bmi_male$BMI, pred_bmi_male$SBP, col = "blue", lwd = 2)
lines(pred_bmi_female$BMI, pred_bmi_female$SBP, col = "red", lwd = 2)
legend("topleft", c("Male", "Female"), col = c("blue", "red"),
    pch = 16, lty = 1, lwd = 2)
```

Effect of Age on Blood PresEffect of BMI on Blood Pres



collaborator:

There are three factors in our dataset that are related to SBP: sex, age, and BMI. We find that: - for every year increase in age, we expect BP to increase by about 1 mmHg if we hold other variables constant - women typically have lower SBP than men (9 mmHg lower) - for every 1 point increase in BMI BP increases by about 1 mmHG with other variables held constant - the other variables like education level, diabetes status, smoking, and statin use didn't show strong enough relationships with BP to be included in our final model

for

We used a linear model here. Linear models are particularly useful. Our model explains about 26% of the variation in the data. We know that our model is not complete – it doesn't have all the factors that may be affecting SBP. But these three variables explain about a quarter of the variation of SBP in the dataset.

Based on our findings, we should more closely monitor "at risk" individuals, specifically people who fall into the higher predicted SBP groups.

Problem 3: The mechanics of prediction [25 points]

A year later, your collaborator returns to you, but he is no longer interested in estimating the associations between predictor variables and SBP. Instead, he is hoping you can help him to build a best predictive model. He has expanded his target population to include all adults, and has collected several more variables. The data (p3.csv) now consists of the following variables:

SBP: Systolic blood pressure, in mmHg.

AGE: Age in years.

GENDER: Gender indicator; 0 = Male, 1 = Female.

EDUC: Highest level of education attained. 0 = Primary School, 1 = Junior High, 2 = High School, 3 = College degree or higher.

DM: Diabetes mellitus indicator; 0 = No, 1 = Yes.

SMOKE: Smoking status indicator; 0 = No, 1 = Yes.

BMI: Boprobdy mass index (kg/m^2) .

STATIN: Statin use indicator; 0 = No, 1 = Yes.

CVD: History of cardiovascular event indicator; 0 = No, 1 = Yes.

HYPERTENSION: Indicator of hypertension; 0 = No, 1 = Yes.

CHOL: Total cholesterol, in mmol/L.

HDL: HDL cholesterol, in mmol/L.

eGFR: Estimated glomerular filtration rate, a measure of kidney function in mL/min (lower values indicate possible kidney damage).

(a) Start by making sure your categorical variables are coded as factors.

```
data <- read.csv("data/p3.csv")

# paranoia
str(data)</pre>
```

```
##
   'data.frame':
                     4065 obs. of
                                   13 variables:
   $ AGE
                          35 35 35 35 35 35 35 35 35 ...
##
   $ GENDER
                    int
                          0 0 0 0 0 0 0 0 0 0 ...
##
   $ CVD
                    int
                          0 0 0 0 0 0 0 0 0 0 ...
   $ DM
                          0 0 0 0 0 0 0 0 0 0 ...
##
                    int
##
   $ HYPERTENSION: int
                          0 0 0 0 0 0 0 0 0 0 ...
                          24.6 21.2 29.2 29.2 29.6 ...
##
   $ BMI
                   : num
   $ SBP
                          116 118 132 130 118 ...
##
                   : num
                          5.5 3.65 6.93 3.95 4.58 5.64 5.35 6.16 4.68 5.09 ...
##
   $ CHOL
                    num
##
   $ HDL
                          0.94 0.87 1.14 0.98 0.92 1.1 0.9 1.19 0.87 1.31 ...
                   : num
                          0 0 0 0 0 0 0 0 0 0 ...
##
    $ STATIN
                   : int
   $ eGFR
                          68.2 104.6 98.5 113.1 90.6 ...
##
                   : num
##
   $ EDUC
                          3 3 2 2 3 1 2 0 2 2 ...
                   : int
```

summary(data) ## AGE GENDER CVD DM:35.0 Min. Min. ## Min. :0.000 :0.000 Min. :0.0000 1st Qu.:45.0 1st Qu.:0.000 1st Qu.:0.000 1st Qu.:0.0000 Median:54.0 Median :0.000 Median :0.000 Median :0.0000 ## ## Mean :54.6 Mean :0.476 Mean :0.075 Mean :0.0615 ## 3rd Qu.:63.0 3rd Qu.:1.000 3rd Qu.:0.000 3rd Qu.:0.0000 ## Max. :82.0 Max. :1.000 Max. :1.000 Max. :1.0000 HYPERTENSION BMISBP CHOL ## HDL Min. ## Min. :0.000 Min. :16.8 Min. : 77.5 : 1.58 Min. :0.48 1st Qu.:0.000 1st Qu.:23.7 1st Qu.:112.5 1st Qu.: 4.64 ## 1st Qu.:1.14 ## Median :0.000 Median:26.1 Median :123.0 Median : 5.31 Median:1.37 ## Mean :0.339 Mean :26.8 Mean :125.9 Mean : 5.36 Mean :1.41 ## 3rd Qu.:1.000 3rd Qu.:29.2 3rd Qu.:136.5 3rd Qu.: 6.05 3rd Qu.:1.63 ## Max. :1.000 Max. :61.0 Max. :218.5 Max. :10.02 Max. :3.44 **EDUC** SMOKE ## STATIN eGFR : 20.9 ## Min. :0.000 Min. :0.00 Min. :0.000 Min. 1st Qu.:0.000 1st Qu.: 69.6 1st Qu.:1.00 1st Qu.:0.000 ## ## Median :0.000 Median : 77.9 Median:2.00 Median : 0.000 ## Mean :0.221 Mean : 78.6 Mean :1.85 Mean :0.239 3rd Qu.: 87.7 ## 3rd Qu.:0.000 3rd Qu.:3.00 3rd Qu.:0.000 ## Max. :1.000 Max. :149.5 Max. :3.00 Max. :1.000 # binary variables to factors binary_vars <- c("GENDER", "DM", "SMOKE", "STATIN", "CVD", "HYPERTENSION") data[binary_vars] <- lapply(data[binary_vars], factor)</pre> # Convert EDUC to an ordered factor with appropriate levels data\$EDUC <- factor(data\$EDUC,</pre> levels = 0:3, labels = c("Primary School", "Junior High", "High School", "College degree or higher"), ordered = TRUE) str(data) ## 'data.frame': 4065 obs. of 13 variables: : int 35 35 35 35 35 35 35 35 35 ... ## \$ AGE : Factor w/ 2 levels "0", "1": 1 1 1 1 1 1 1 1 1 ... \$ GENDER ## ## \$ CVD : Factor w/ 2 levels "0", "1": 1 1 1 1 1 1 1 1 1 1 ... : Factor w/ 2 levels "0", "1": 1 1 1 1 1 1 1 1 1 1 ... ## \$ DM ## \$ HYPERTENSION: Factor w/ 2 levels "0","1": 1 1 1 1 1 1 1 1 1 1 ... : num 24.6 21.2 29.2 29.2 29.6 ... ## \$ BMI : num 116 118 132 130 118 ... ## \$ SBP ## \$ CHOL : num 5.5 3.65 6.93 3.95 4.58 5.64 5.35 6.16 4.68 5.09 ...

\$ HDL

: num 0.94 0.87 1.14 0.98 0.92 1.1 0.9 1.19 0.87 1.31 ...

```
## $ STATIN : Factor w/ 2 levels "0","1": 1 1 1 1 1 1 1 1 1 1 1 ...
## $ eGFR : num 68.2 104.6 98.5 113.1 90.6 ...
## $ EDUC : Ord.factor w/ 4 levels "Primary School"<..: 4 4 3 3 4 2 3 1 3 3 ...
## $ SMOKE : Factor w/ 2 levels "0","1": 2 1 2 1 1 2 1 1 2 ...
#lets go</pre>
```

(b) Break the dataset into an 80-20 train-test split.

```
set.seed(139)

train_size <- floor(0.8 * nrow(data))

train_indices <- sample(seq_len(nrow(data)), size = train_size)

train_data <- data[train_indices, ]

test_data <- data[-train_indices, ]

cat("Training size:", nrow(train_data), "observations\n")

## Training size: 3252 observations

cat("Test size:", nrow(test_data), "observations\n")

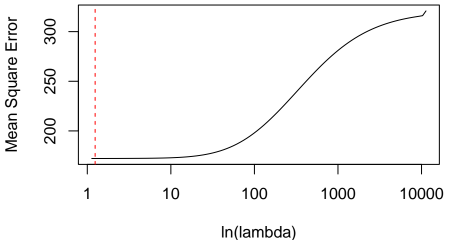
## Test size: 813 observations

cat("Split proportion (train):", nrow(train_data)/nrow(data), "\n")</pre>
```

Split proportion (train): 0.8

(c) Using your training data and a k of 5, perform k-fold cross validation with ridge regression to identify the value of λ that produces the lowest mean RMSE for a full model that includes all main effects and 2-way interaction terms. Plot the MSE vs. $\ln(\lambda)$. Report the optimal λ and the RMSE on your test data.

```
")^2"))))
# model matrices for both training and test data
x_train <- model.matrix(formula, data = train_data)[,-1] # Remove intercept</pre>
y_train <- train_data$SBP</pre>
x_test <- model.matrix(formula, data = test_data)[,-1]</pre>
y_test <- test_data$SBP</pre>
# k-fold
cv_ridge <- cv.glmnet(x_train, y_train,</pre>
                       alpha = 0, # ridge!!
                       nfolds = 5,
                       standardize = TRUE)
plot(cv_ridge$lambda, cv_ridge$cvm,
     type = "1",
     xlab = "ln(lambda)",
     ylab = "Mean Square Error",
     log = "x")
abline(v = cv_ridge$lambda.min, col = "red", lty = "dashed")
```



```
cat("Optimal lambda:", optimal_lambda, "\n")
## Optimal lambda: 1.241
cat("Test set RMSE:", test_rmse, "\n")
## Test set RMSE: 13.71
The optimal lambda is 1.241 The test set RMSE is 13.71
(d) Repeat part c) using LASSO.
set.seed(139)
cv_lasso <- cv.glmnet(x_train, y_train,</pre>
                        alpha = 1, # LASSO regression
                        nfolds = 5,
                        standardize = TRUE)
plot(cv_lasso$lambda, cv_lasso$cvm,
     type = "l",
     xlab = "ln(lambda)",
     ylab = "Mean Square Error",
     log = "x")
abline(v = cv_lasso$lambda.min, col = "red", lty = "dashed")
     300
Mean Square Error
     250
     200
                    1e-02
        1e-03
                                1e-01
                                            1e+00
                                                        1e+01
                              In(lambda)
optimal_lambda_lasso <- cv_lasso$lambda.min
final_model_lasso <- glmnet(x_train, y_train,</pre>
                             alpha = 1,
                             lambda = optimal_lambda_lasso,
                             standardize = TRUE)
predictions_lasso <- predict(final_model_lasso, newx = x_test)</pre>
```

test_rmse_lasso <- sqrt(mean((y_test - predictions_lasso)^2))</pre>

```
# Print results
cat("Optimal lambda:", optimal_lambda_lasso, "\n")
## Optimal lambda: 0.2736
cat("Test set RMSE:", test_rmse_lasso, "\n")
```

Test set RMSE: 13.82

Optimal lambda: 0.2736 Test set RMSE: 13.82

Problem 4: A little theory and simulation [25 points]

In this problem, you will explore the concept of shrinkage (or regularization) through an example called "Stein's paradox" (parts a-c) and then draw a connection with ridge regression in part d.

Suppose $Y_i \sim \mathcal{N}(\mu_i, \sigma^2)$ for i = 1, ..., n. You observe $Y_1, ..., Y_n$ and want to estimate $\mu_1, ..., \mu_n$.

(a) The most obvious estimator of μ_i is $\hat{\mu}_i = Y_i$. What is the bias of this estimator? Bias calculation:

$$Bias(\hat{\mu}_i) = E[\hat{\mu}_i] - \mu_i = \mu_i - \mu_i = 0$$

This is an unbiased estimator

- (b) A shrinkage estimator of μ_i is $\hat{\mu}_i(c) = cY_i$ where $c \in [0,1]$ is a user-specified parameter that controls the amount of shrinkage. Write R code to do the following:
 - Set n = 1000. Randomly generate $\mu_i \sim \text{Uniform}(0, 10)$ for i = 1, ..., n.
 - Randomly generate $Y_i \sim \mathcal{N}(\mu_i, \sigma^2)$ for i = 1, ..., n where $\sigma = 4$.
 - For each $c \in \{0.0, 0.01, 0.02, \dots, 0.99, 1.0\}$ compute MSE(c) where:

$$MSE(c) = \frac{1}{n} \sum_{i=1}^{n} (\hat{\mu}_i(c) - \mu_i)^2$$

Plot MSE(c) versus c, along with the variance and the square of the bias. For your dataset, what value of c minimizes MSE(c)? So, does shrinkage help?

```
set.seed(139)

n <- 1000
sigma <- 4
c_values <- seq(0, 1, by=0.01)

mu <- runif(n, min=0, max=10)

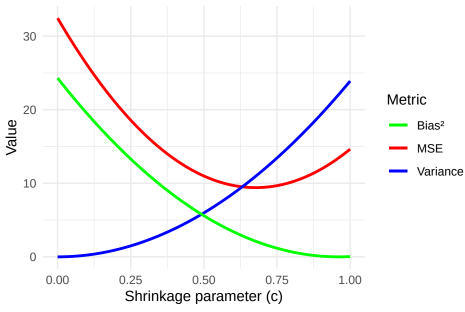
Y <- rnorm(n, mean=mu, sd=sigma)

results <- sapply(c_values, function(c) {
    # Shrinkage estimates
    mu_hat <- c * Y

# Calculate MSE
    mse <- mean((mu_hat - mu)^2)</pre>
```

```
# Bias
    bias <- mean(mu_hat - mu)</pre>
    # Variance
    var <- mean((mu_hat - mean(mu_hat))^2)</pre>
    return(c(mse=mse, bias=bias, var=var))
})
results_df <- data.frame(</pre>
    c = c_values,
    MSE = results[1,],
    Bias = results[2,],
    Variance = results[3,]
)
p <- ggplot(results_df) +</pre>
    geom_line(aes(x=c, y=MSE, color="MSE"), size=1) +
    geom_line(aes(x=c, y=Variance, color="Variance"), size=1) +
    geom_line(aes(x=c, y=Bias^2, color="Bias^2"), size=1) +
    labs(x="Shrinkage parameter (c)",
         y="Value",
         title="MSE, Variance, and Bias2 vs Shrinkage Parameter",
         color="Metric") +
    theme_minimal() +
    scale_color_manual(values=c("MSE"="red", "Variance"="blue", "Bias2"="green"))
## Warning: Using `size` aesthetic for lines was deprecated in ggplot2 3.4.0.
## i Please use `linewidth` instead.
## This warning is displayed once every 8 hours.
## Call `lifecycle::last_lifecycle_warnings()` to see where this warning was
## generated.
print(p)
```

MSE, Variance, and Bias² vs Shrinkage Parameter



```
opt_c <- c_values[which.min(results_df$MSE)]
cat("Optimal c value:", opt_c, "\n")</pre>
```

Optimal c value: 0.68

cat("Minimum MSE:", min(results_df\$MSE), "\n")

Minimum MSE: 9.413

Optimal c value: 0.68 Minimum MSE: 9.413

YES shrinkagedoes help because the c value is not 1. If the optimal shrinkage was 1 or close to 1, it would not help because it wouldn't change hte Y_i much, but it is .68

(c) Explain your findings from (b) in terms of the bias-variance tradeoff. What happens to the bias and the variance when $c \to 0$?

bias vairance tradeoff - variance increases quadratically as C increases - bias 2 decreases as C increase

SO when C goes to 0, - variance approaches 0 - bias approaches the maximum of around 24 - this happens because as we are shrinking hteestimates, this means that all of the estimates get closer together (lower variance) and farther away from the true means (bias increase)

This is the whole point of shrinkage – we want to optimize bias variance tradeoff

(d) Suppose we defined our ridge regression estimates as:

$$\hat{\boldsymbol{\beta}}_{\lambda}^{R} = \left(\mathbf{X}'\mathbf{X} + \lambda \mathbf{I}\right)^{-1} \mathbf{X}' \vec{\mathbf{y}}$$

where $\vec{\mathbf{y}} \in \mathbb{R}^{n \times 1}$. Note this is a slightly different definition than was used in class. But suppose you want to frame the shrinkage problem in parts a) - c) as a ridge regression problem. How should

you define X and I? Make sure to include dimensions.

% Model definition $\beta_i \colon Y_i = \beta_i + \varepsilon_i$

In matrix notation this is: $\vec{\mathbf{y}} = \mathbf{X}\boldsymbol{\beta} + \boldsymbol{\varepsilon}$

To match the shrinkage estimator, define: **X**: An $n \times n$ identity matrix. This means each Y_i is directly associated with its own β_i .

$$\mathbf{X} = \begin{bmatrix} 1 & 0 & \cdots & 0 \\ 0 & 1 & \cdots & 0 \\ \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & \cdots & 1 \end{bmatrix}$$

 ${f X}$ is the identity matrix. Since ${f X}$ is the identity matrix:

$$\mathbf{X}^{\mathsf{T}}\mathbf{X} = \mathbf{I}$$

Thus our estimator simplifies to be:

$$\hat{\boldsymbol{\beta}}_{\lambda}^{R} = (\mathbf{I} + \lambda \mathbf{I})^{-1} \, \mathbf{\vec{y}} = \frac{1}{1 + \lambda} \mathbf{\vec{y}}$$

Thus when we set $c = \frac{1}{1+\lambda}$, this matches our shrinkage estimator $\hat{\mu}(c) = cY_i$

Dimensions:

- X is an $n \times n$ identity matrix
- I is also the $n \times n$ identity matrix

When we make X an identity matrix, the shrinkage estimator becomes a ridge regression estimator. This shows that the shrinkage in this context is the same as applying RIDGE with a λ .

(e) Using the design matrix you defined in part d), show that $\hat{\boldsymbol{\beta}}_{\lambda}^{R} = c\vec{\mathbf{y}}$ for some $c \in \mathbb{R}$. Give the actual mathematical expression for c as a function of λ .

The ridge regression estimator is:

$$\hat{\boldsymbol{\beta}}_{\lambda}^{R} = \left(\mathbf{X}^{\top}\mathbf{X} + \lambda\mathbf{I}\right)^{-1}\mathbf{X}^{\top}\vec{\mathbf{y}}$$

Since \mathbf{X} is the identity matrix, we have:

$$\mathbf{X}^{\top}\mathbf{X} = \mathbf{I}^{\top}\mathbf{I} = \mathbf{II} = \mathbf{I}$$

Therefore, the term $\mathbf{X}^{\top}\mathbf{X} + \lambda \mathbf{I}$ simplifies to:

$$\mathbf{X}^{\top}\mathbf{X} + \lambda \mathbf{I} = \mathbf{I} + \lambda \mathbf{I} = (1 + \lambda)\mathbf{I}$$

The inverse of this term is:

$$\left(\mathbf{X}^{\top}\mathbf{X} + \lambda \mathbf{I}\right)^{-1} = \left((1 + \lambda)\mathbf{I}\right)^{-1} = \frac{1}{1 + \lambda}\mathbf{I}$$

Next, since $\mathbf{X}^{\top} = \mathbf{I}^{\top} = \mathbf{I}$, we have:

$$\mathbf{X}^{ op} \ddot{\mathbf{y}} = \mathbf{I} \ddot{\mathbf{y}} = \ddot{\mathbf{y}}$$

Substituting back into the ridge regression estimator:

$$\begin{split} \hat{\boldsymbol{\beta}}_{\lambda}^{R} &= \left(\mathbf{X}^{\top}\mathbf{X} + \lambda\mathbf{I}\right)^{-1}\mathbf{X}^{\top}\vec{\mathbf{y}} \\ &= \left(\frac{1}{1+\lambda}\mathbf{I}\right)\vec{\mathbf{y}} \\ &= \frac{1}{1+\lambda}\vec{\mathbf{y}} \end{split}$$

Thus, we have shown that:

$$\hat{\boldsymbol{\beta}}_{\lambda}^{R} = c\vec{\mathbf{y}}$$

Using the identity matrix for both ${\bf X}$ and ${\bf I}$, the ridge regression estimator simplifies to scaling the vector ${\bf y}$ by a constant factor c which depends on λ .