# MACS 33002: PSET 2

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```
devtools::install_github("uc-cfss/rcfss")
Skipping install of 'rcfss' from a github remote, the SHA1 (6dd8d8be) has not changed since last :
  Use `force = TRUE` to force installation
library(tidyverse)
#library(ISLR)
library(broom)
library(rsample)
library(rcfss)
Attaching package: 'rcfss'
The following object is masked from 'package:modelr':
   mse
library(yardstick)
For binary classification, the first factor level is assumed to be the event.
Set the global option `yardstick.event_first` to `FALSE` to change this.
Attaching package: 'yardstick'
The following objects are masked from 'package:modelr':
   mae, mape, rmse
The following object is masked from 'package:readr':
    spec
electData <- read_csv("nes2008.csv")</pre>
Parsed with column specification:
cols(
  biden = col_double(),
  female = col_double(),
  age = col_double(),
  educ = col_double(),
  dem = col_double(),
  rep = col_double()
```

1. (10 points) Estimate the MSE of the model using the traditional approach. That is, fit the linear regression model using the entire dataset and calculate the mean squared error for the entire dataset. Present and discuss your results at a simple, high level.

```
reg <- lm(biden ~ female + age + educ + dem + rep, data=electData)
summary(reg)
Call:
lm(formula = biden ~ female + age + educ + dem + rep, data = electData)
Residuals:
   Min
           1Q Median
                          3Q
                                Max
                 1.02
-75.55 -11.29
                       12.78
                              53.98
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept)
             58.8113
                          3.1244
                                   18.82
                                          < 2e-16 ***
              4.1032
                          0.9482
                                    4.33 0.000016 ***
female
              0.0483
                          0.0282
                                    1.71
                                             0.088 .
age
educ
             -0.3453
                          0.1948
                                   -1.77
                                             0.076 .
                          1.0680
dem
             15.4243
                                   14.44
                                          < 2e-16 ***
                                  -12.09
            -15.8495
                          1.3114
                                          < 2e-16 ***
rep
Signif. codes:
                0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 19.9 on 1801 degrees of freedom
Multiple R-squared: 0.282, Adjusted R-squared:
F-statistic: 141 on 5 and 1801 DF, p-value: <2e-16
mse <- MSE(y_pred = reg$fitted.values, y_true = electData$biden)</pre>
mse
[1] 395.3
m <- mean((reg\fitted.values-electData\fiden)^2)
```

#### [1] 395.3

We can see that the mean squared error is 395.3 when we use the whole data for the model. When we look at the coefficients, we can see that they are significant on all the parameters. The coefficient for female shows that females tend to like Biden. The same can be observed for older people. There is a negative correlation between education and liking Biden, so more educated people tend to like Biden less. As we would expect, democrats tends to like Biden more than independents whereas republicans like him less than the independents do. The R-squared is 0.282, which means that the model is not explaining the variation in the data well. Therefore, it is likely that this model is missing features that could predict a person's feelings towards Biden. The mean squared error is 395.3, which shows that the representative residual is approximately 20 points away. This is concerning because 20 points is larger than our coefficients. Therefore, this model might not have good predictive power.

2. (30 points) Calculate the test MSE of the model using the simple holdout validation approach.

(5 points) Split the sample set into a training set (50%) and a holdout set (50%). Be sure to set your seed prior to this part of your code to guarantee reproducibility of results.

(5 points) Fit the linear regression model using only the training observations.

```
reg_train <- lm(biden ~ female + age + educ + dem + rep, data=data_train)
sm <- summary(reg_train)</pre>
Call:
lm(formula = biden ~ female + age + educ + dem + rep, data = data_train)
Residuals:
  Min
          1Q Median
                        3Q
                              Max
-71.60 -11.48
               1.08 13.96 46.03
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 57.8664
                        4.4325 13.06 < 2e-16 ***
             5.4727
                        1.3622 4.02 0.000064 ***
female
             0.0398
                        0.0409
                                  0.97
                                           0.33
age
educ
            -0.3367
                        0.2783 - 1.21
                                           0.23
                        1.5190 9.87 < 2e-16 ***
            14.9923
dem
                        1.8919 -8.97 < 2e-16 ***
rep
           -16.9624
               0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Signif. codes:
Residual standard error: 20.2 on 898 degrees of freedom
Multiple R-squared: 0.284, Adjusted R-squared: 0.28
F-statistic: 71.2 on 5 and 898 DF, p-value: <2e-16
```

(10 points) Calculate the MSE using only the test set observations.

```
mse_test <- mean((data_test$biden - predict.lm(reg_train, data_test)) ^ 2)
mse_test</pre>
```

[1] 389.2

(10 points) How does this value compare to the training MSE from question 1? Present numeric comparison and discuss a bit.

The MSE we've found in Question 1 (395.3) is higher than the one we've found in Question 2 (389.2). We can say that this was somehow not expected becausse fitting the model on less data and testing it on data that it hasn't observed before would potentially lead to a worse model performance. However, we should keep in mind that this is a random process so we can expect that some seeds will grab smaller values and other seeds might grab larger values.

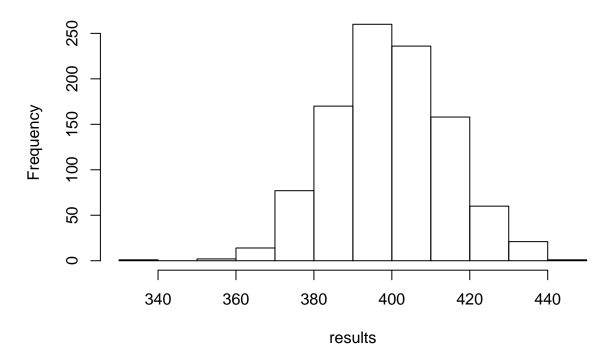
3. (30 points) Repeat the simple validation set approach from the previous question 1000 times, using 1000 different splits of the observations into a training set and a test/validation set. Visualize your results as a sampling distribution (hint: think histogram or density plots). Comment on the results obtained.

```
results <- numeric(1000)

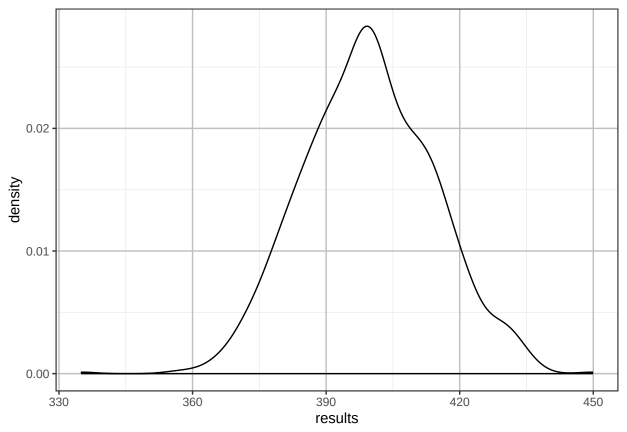
for (i in 1:1000) {
    split_1000 <- initial_split(data = electData, prop = 0.5)
    train_1000 <- training(split_1000)
    test_1000 <- testing(split_1000)
    model_1000 <- lm(biden ~ female + age + educ + dem + rep, data = train_1000)
    mse_test_1000 <- mean((test_1000$biden - predict.lm(model_1000, test_1000)) ^ 2)
    results[i] <- mse_test_1000
}

hist(results)</pre>
```

## Histogram of results



### ggplot(as.data.frame(results), aes(results))+geom\_density()

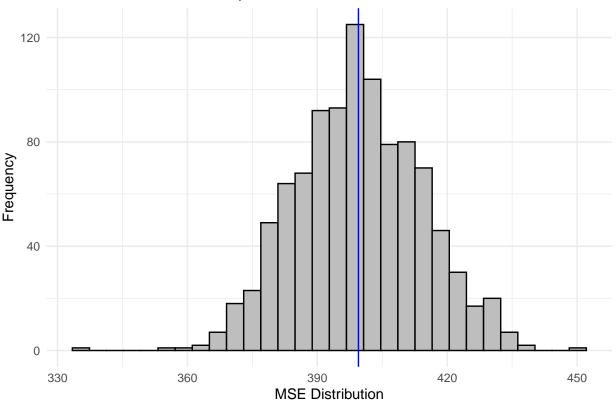


```
res_df <- as.data.frame(results)

ggplot(res_df) + geom_histogram(aes(results), boundary = 1, color="black", fill="gray")+ theme_min
geom_vline(xintercept = mean(results), color = "blue") +
    labs(x = "MSE Distribution", y = "Frequency") +
    ggtitle("MSE Distribution with Repetition")</pre>
```

<sup>`</sup>stat\_bin()` using `bins = 30`. Pick better value with `binwidth`.

## MSE Distribution with Repetition



observe that as we take more samples and calculate the mean squared error, we can see that the distribution converges to a notmal with a mean approximately 400, which is very close to the MSE calculated using the full dataset. There are still some variations due to the random sampling that we do. We can calculate the spread of the distribution:

We can

# df <- as.data.frame(results) summary(df)</pre>

#### results

Min.: 335 1st Qu.:389 Median: 399 Mean: 400 3rd Qu.:410 Max.: 450

As observed, the max and min show a difference in fits of the model. The histogram that we've plotted above also shows the importance of iteration to converge to an optimal result in order to have a calibrated model. By just doing sampling once, we could have been misled if we happened to get a value around the tail of this distribution.

4. (30 points) Compare the estimated parameters and standard errors from the original model in question 1 (the model estimated using all of the available data) to parameters and standard errors estimated using the bootstrap (B=1000). Comparison should include, at a minimum, both numeric output as well as discussion on differences, similarities, etc. Talk also about the conceptual use and impact of bootstrapping.

```
library(ISLR)
lm_df <-
  summary(reg)$coefficients %>%
  data.frame() %>%
  mutate(term = names(reg$coefficients)) %>%
  select(term, Lm_Estimate = Estimate, Lm_Sd = Std..Error)
lm_df
                             Lm_Sd
         term Lm_Estimate
1 (Intercept)
                 58.81126 3.12444
2
       female
                  4.10323 0.94823
3
                   0.04826 0.02825
          age
4
         educ
                 -0.34533 0.19478
5
          dem
                 15.42426 1.06803
6
                 -15.84951 1.31136
lm_coefs <- function(splits, ...) {</pre>
  mod <- lm(..., data = analysis(splits))</pre>
  tidy(mod)
}
auto_boot <- electData %>%
  bootstraps(1000) %>%
  mutate(coef = map(splits, lm_coefs, as.formula(biden ~ female + age + educ + dem + rep)))
auto_boot %>%
  unnest(coef) %>%
  group by(term) %>%
  summarize(Boot_Estimate = mean(estimate),
            Boot_Sd= sd(estimate, na.rm = TRUE))
# A tibble: 6 x 3
              Boot_Estimate Boot_Sd
  term
  <chr>>
                       <dbl>
                               <dbl>
1 (Intercept)
                     58.7
                              3.07
                      0.0491 0.0284
2 age
                     15.4
3 dem
                              1.06
4 educ
                     -0.344
                              0.197
5 female
                      4.13
                              0.960
                    -15.8
                              1.41
6 rep
```

Bootstrapping relies on random sampling with replacement to assign measures of accuracy to sample estimates. We know that using a bootstraped estimator is useful because the bootstrap estimator will be more accurate especially when the distributional assumptions are not met. We can observe that the

bootstrap results are similar to the results we got for the original non-bootstrapped model. The standard deviations are also relatively close to each other. When we take the standard errors into account, the coefficients are even the same as the original model that we've found in Q1. This is a very good sign because we would not have many other ways to compute standard errors on the estimates, so we know that we can use bootstrapping to get robust standard errors.