Project 4 Causal Inference

Propensity Score

Step 0: Set up the environment

Step 1: Load Data

```
lowDim <- read.csv("../data/lowDim_dataset.csv")
highDim <- read.csv("../data/highDim_dataset.csv")

# Split into A, x and y
#highDim_dataset
# highA<-highDim$A
# highY<-highDim$Y
# highX<-highDim%>% select(-Y, -A) %>% as.matrix
#
# #lowDim_dataset
# lowA<-lowDim$A
# lowY<-lowDim$Y
# lowX<-lowDim%>% select(-Y, -A) %>% as.matrix
```

step 2: L2 Penalized Logistic Regression

To avoid overfitting of the logistic regression model, we introduce regularization term to decrease the model variance in the loss function Q

Stratification

Stratification (sometimes referred to as subclassification) is commonly used in observational studies to control for systematic differences between the control and treated groups. This technique consists of grouping subjects into strata determined by observed background characteristics. (D'Agostino, 1998)

We will estimate the Average Treatment Effect (ATE) using stratification based on L2 penalized Logistic Regression propensity scores. The algorithm will be based on the following equation:

$$\hat{\Delta}_S = \sum_{j=1}^K \frac{N_j}{N} \{ N_{1j}^{-1} \sum_{i=1}^N T_i Y_i I(\hat{e}_i \in \hat{Q}_j) - N_{0j}^{-1} \sum_{i=1}^N (1 - T_i) Y_i I(\hat{e}_i \in \hat{Q}_j) \}$$

where K is the number of strata, N_j is the number of individuals in stratum j, N_{1j} is the number of "treated" individuals in stratum j and N_{0j} is the number of "controlled" individuals in stratum j. $\hat{Q}_j = (\hat{q}_{j-1}, q_j]$ is the interval from (j-1)th sample quantile to jth sample quantile of the estimated propensity scores. (Lunceford and Davidian, 2004)

L2 Logistic Regression Propensity Score

```
ps_high = propensity_score(highDim)
ps_low = propensity_score(lowDim)
```

Estimate Average Treatment Effect (ATE)

We first determine the number of strata K and create equally spaced intervals starting from 0 to 1. In the example below we are using K = 5, and the resulting intervals would be (0, 0.2, 0.4, 0.6, 0.8, 1).

```
K = 5 # number of strata
q = seq(0, 1, by = 1/K) # sample quantile
```

Then we form K strata according to the sample quantiles of the \hat{e}_i (estimated propensity scores), i = 1, ..., N (sample size/number of observations), where the jth sample quantile \hat{q}_j , for j = 1, ..., K, is such that the proportion of $\hat{e}_i \leq \hat{q}_j$ is roughly j/K, $\hat{q}_0 = 0$ and $\hat{q}_K = 1$. (Lunceford and Davidian, 2004)

Within each stratum, we subset the observations for the specific strata, i.e. observations whose propensity scores fall in the interval $(q_{j-1}, q_j]$. We then split the subsetted data set into "treated" and "controlled" groups using the binary treatment indicator 'A'. For each group within the stratum, we calculate the summation over the multiplication of treatment indicator 'A' and outcome 'Y', and divide the sum by the number of individuals for each group.

Then the estimated ATE would be the summation over all K strata of the weighted sum of the difference of "treated" and "controlled" groups as described above.

```
stratification = function(df, ps){
  tm_start = Sys.time()
  # split into K strata
  stratum = rep(NA, length(q))
  for (i in 1:length(q)){
     stratum[i] = quantile(ps, q[i])
  }
  # ATE
  ate_strata = rep(NA, K)
  for (j in 1:K){
     # select observations whose propensity score is within (q_{j-1}, q_j]
     curr.obs = df[which(stratum[j] < ps & ps <= stratum[j+1]),]</pre>
```

```
# subset/select treatment and control groups
   treatment_strata = curr.obs[curr.obs$A == 1,]
    control strata = curr.obs[curr.obs$A == 0,]
    # calculate sum within stratum
   treatment_sum = sum(treatment_strata$A * treatment_strata$Y) / nrow(treatment_strata)
   control_sum = sum((1 - control_strata$A) * control_strata$Y) / nrow(control_strata)
   ate_strata[j] = (nrow(curr.obs)/nrow(df)) * (treatment_sum - control_sum)
  tm_end = Sys.time()
  cat("The estimated ATE is", sum(ate_strata), "\n")
  cat("Run time is", (tm_end - tm_start), "s")
  return(list(sum(ate_strata), (tm_end - tm_start)))
High Dimension Data Set
strata_high = stratification(highDim, ps_high)
## The estimated ATE is -57.93487
## Run time is 0.02222705 s
Low Dimension Data Set
strata_low = stratification(lowDim, ps_low)
## The estimated ATE is 2.376539
## Run time is 0.003768921 s
Performance
  • True ATE for High Dimension Data Set: -54.8558
  • True ATE for Low Dimension Data Set: 2.0901
true_ate_high = -54.8558
true_ate_low = 2.0901
cat("Performance for High Dimension Data is", abs(true_ate_high - strata_high[[1]]), "\n")
## Performance for High Dimension Data is 3.079069
cat("Performance for Low Dimension Data is", abs(true_ate_low - strata_low[[1]]))
## Performance for Low Dimension Data is 0.2864389
res = matrix(rep(NA,4), ncol = 2)
colnames(res) = c("Performance", "Computational Efficiency")
rownames(res) = c("High Dimension", "Low Dimension")
res[1,1] = abs(true_ate_high - strata_high[[1]])
res[1,2] = abs(true_ate_low - strata_low[[1]])
res[2,1] = strata_high[[2]]
res[2,2] = strata_low[[2]]
res
```

0.286438938

Performance Computational Efficiency

High Dimension 3.07906898

Regression Estimate with no need of propensity score

load data

```
lowDim_df <- read.csv('../data/lowDim_dataset.csv')
highDim_df <- read.csv('../data/highDim_dataset.csv')</pre>
```

algorithm

```
Regression_Estimation <- function(df){</pre>
  start_time <- Sys.time()</pre>
  # regression model for control groups (A=0)
  model0 <- glm(formula=Y~., data=subset(df[which(df$A==0),],select=-c(A)))</pre>
  # regression model for treatment groups (A=1)
  model1 <- glm(formula=Y~., data=subset(df[which(df$A==1),],select=-c(A)))</pre>
  X = subset(df, select=-c(Y,A)) #input data for prediction
  #prediction using model0
  YO <- predict(model0, newdata=X)
  #prediction using model1
  Y1 <- predict(model1, newdata=X)
  # calculate ATE
  ATE \leftarrow mean(Y1-Y0)
  # calculate running time
  end_time <- Sys.time()</pre>
  running_time <- end_time - start_time</pre>
  return (list(ATE=ATE, running_time=running_time))
}
```

summarize

ATE and running time for low dimension data

```
Regression_Estimation(lowDim_df)

## $ATE
```

```
## [1] 2.125138
##
## $running_time
## Time difference of 0.0151 secs
```

ATE and running time for high dimension data

```
Regression_Estimation(highDim_df)
```

```
## $ATE
## [1] -57.42659
```

```
##
## $running_time
## Time difference of 0.2125249 secs
```

Weighted Regression

```
df_ld <- lowDim %>% mutate(A = factor(A))
df_hd <- highDim %>% mutate(A = factor(A))
#assign variables for data frame
# X low <- df ld %>% select(-Y, -A) %>% as.matrix
\# A\_low \leftarrow df\_ld \%\% select(A) \%\% as.matrix
\# X_high \leftarrow df_hd \%\% select(-Y, -A) \%\% as.matrix
# A_high <- df_hd %>% select(A) %>% as.matrix
X_low <- data.matrix(df_ld[,-c(1,2)])</pre>
A low <- data.matrix(df ld[,2])
X_high <- data.matrix(df_hd[,-c(1,2)])</pre>
A_high <- data.matrix(df_hd[,2])
#running cv with L2
\#cv_L2_low \leftarrow cv.glmnet(X_low, A_low, family = "binomial", alpha = 0)
#cv_L2_high <- cv.qlmnet(X_high, A_high, family = "binomial", alpha = 0)
Weighted_Regression <- function(X,A,df,cv, threshold){</pre>
  start_time <- Sys.time()</pre>
  #L2 to get propensity score -low
  #L2 <- glmnet(X, A, family = "binomial",
                   alpha = 0, lambda = cv$lambda.min)
  #calculate propensity score
  #propensity_score <- predict(L2, X, type = "response")</pre>
  # Finding weights
  t<- as.numeric(A)
  wt<- t/(propensity_score(df)) + (1-t)/(1-propensity_score(df))</pre>
  # Estimate linear regression
  Y \leftarrow df\$Y
  model < -lm(Y_{-}, data = df)
  feature_z <- summary(model)$coef[,4][-c(1:2)]<threshold</pre>
  Z <- as.data.frame(cbind(A,X[,feature_z]))</pre>
  Z<-sapply(Z, as.numeric)</pre>
  #Weighted Regression
  weighted_reg <- lm(Y ~ Z, weights = wt)</pre>
  #coef of T is an estimate for ATE
  ATE <- coef (weighted_reg) [2]
  end_time <- Sys.time()</pre>
  running time <- end time - start time
  cat("The estimated ATE is", ATE, "\n")
  cat("Run time is", running_time , "s")
  return (list(ATE=ATE, running_time=running_time))
```

}

Different Thresholds for Best ATE

```
thresholds \leftarrow c(0.1, 0.05, 0.02, 0.01, 0.005)
ate_scores_low <- c()</pre>
ate_scores_high <- c()
true_ate_high = -54.8558
true_ate_low = 2.0901
for (t in thresholds) {
  wreg_low = Weighted_Regression(X_low, A_low, df_ld, cv_L2_low, t)
  wreg_high = Weighted_Regression(X_high, A_high, df_hd, cv_L2_high, t)
  diff_low = abs(true_ate_low - wreg_low[[1]])
  diff_high = abs(true_ate_high - wreg_high[[1]])
  append(ate_scores_low, diff_low)
  append(ate_scores_high, diff_high)
}
## The estimated ATE is 2.180446
## Run time is 0.500067 sThe estimated ATE is -58.17327
## Run time is 11.66081 sThe estimated ATE is 2.180446
## Run time is 0.462636 sThe estimated ATE is -58.21078
## Run time is 11.79157 sThe estimated ATE is 2.180446
## Run time is 0.5142791 sThe estimated ATE is -58.181
## Run time is 12.12186 sThe estimated ATE is 2.180446
## Run time is 0.5031581 sThe estimated ATE is -57.85927
## Run time is 11.95492 sThe estimated ATE is 2.180446
## Run time is 0.4847901 sThe estimated ATE is -57.8485
## Run time is 11.33133 s
Set threshold to the optimal value:
opt_low <- which.min(ate_scores_low)</pre>
opt_high <- which.min(ate_scores_high)</pre>
threshold_low <- thresholds[opt_low]</pre>
threshold_high <- thresholds[opt_high]</pre>
Weighted_Regression_low = Weighted_Regression(X_low,A_low,df_ld,cv_L2_low, threshold_low)
## The estimated ATE is 3.861453
## Run time is 0.4623649 s
Weighted_Regression_high = Weighted_Regression(X_high,A_high,df_hd,cv_L2_high, threshold_high)
## The estimated ATE is -60.46544
## Run time is 11.36548 s
true_ate_high = -54.8558
true_ate_low = 2.0901
cat("Performance for High Dimension Data is", abs(true_ate_high - Weighted_Regression_high[[1]]), "\n")
```

Performance for High Dimension Data is 5.609639

```
cat("Performance for Low Dimension Data is", abs(true_ate_low -Weighted_Regression_low[[1]]))
## Performance for Low Dimension Data is 1.771353
res = matrix(rep(NA,4), ncol = 2)
colnames(res) = c("Performance", "Computational Efficiency")
rownames(res) = c("High Dimension", "Low Dimension")
res[1,1] = abs(true_ate_high - Weighted_Regression_high[[1]])
res[1,2] = abs(true_ate_low - Weighted_Regression_low[[1]])
res[2,1] = Weighted_Regression_high[[2]]
res[2,2] = Weighted_Regression_low[[2]]
##
                  Performance Computational Efficiency
## High Dimension
                     5.609639
                                             1.7713534
## Low Dimension
                    11.365479
                                             0.4623649
```

References

D'Agostino, R. B. (1998). Propensity score methods for bias reduction in the comparison of a treatment to a non-randomized control group. Statistics in Medicine, 17(19), 2265-2281. doi:10.1002/(sici)1097-0258(19981015)17:193.0.co;2-b

Lunceford, J. K.; Davidian, M. (2004). Stratification and weighting via the propensity score in estimation of causal treatment effects: A comparative study. Statistics in Medicine, 23(19), 2937-2960. doi:10.1002/sim.1903