Summary Report of Pairings 1 2 5 12 And 16

Group 6

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In this project, we are implementing 5 algorithms:

- 1. A1+D1 Propensity Matching with Mahalanobis distance measure
- 2. A1+D2+P1 Propensity Matching with propensity score distance measure + logistic regression
- $3.\ A1+D3+P1\ Propensity\ Matching\ with\ linear\ Propensity\ score\ distance\ measure\ +\ logistic\ regression$
- 4. A3+P1 Doubly Robust Estimation + logistic regression
- 5. A5+P1 Stratification + logistic regression

Setting up environment

```
library(MatchIt)
library(dplyr)
library(ggplot2)
library(lmtest)
library(sandwich)
library(optmatch)
library(broom)
```

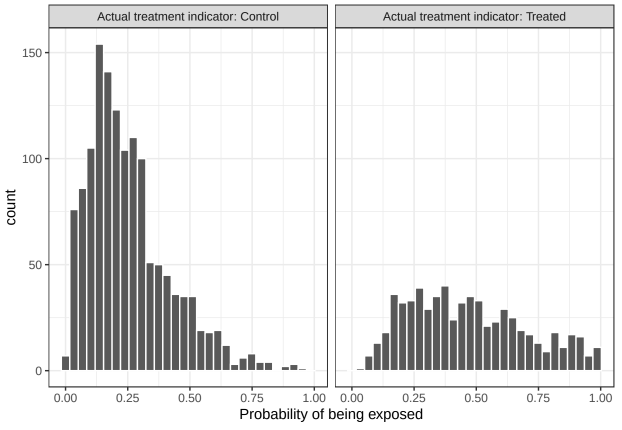
Pre-Analysis

```
# Reading in the datasets as R Dataframes
highDim_data = read.csv("../data/highDim_dataset.csv")
lowDim_data = read.csv("../data/lowDim_dataset.csv")
highDim_data %>%
  group_by(A) %>%
  summarise(n=n(),
            mean outcome=mean(Y),
            std_error = sd(Y) / sqrt(n), .groups = "drop")
## # A tibble: 2 x 4
         Α
               n mean_outcome std_error
## * <int> <int>
                        <dbl>
                                   <dbl>
## 1
         0 1357
                         29.5
                                   0.588
             643
                        -45.7
                                   2.59
with(highDim_data, t.test(Y ~ A))
##
    Welch Two Sample t-test
##
```

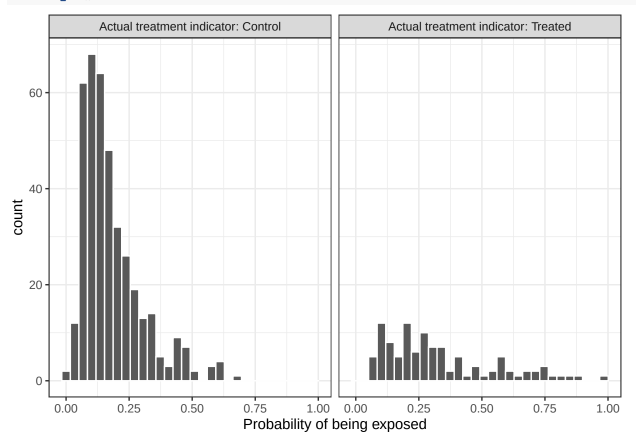
```
## data: Y by A
## t = 28.278, df = 708.98, p-value < 2.2e-16
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## 69.95224 80.39043
## sample estimates:
## mean in group 0 mean in group 1
          29.45850
                         -45.71284
lowDim_data %>%
  group_by(A) %>%
  summarise(n=n(),
            mean_outcome=mean(Y),
            std_error = sd(Y) / sqrt(n), .groups = "drop")
## # A tibble: 2 x 4
        Α
              n mean_outcome std_error
## * <int> <int>
                        <dbl>
                                  <dbl>
                         16.6
                                  0.531
## 1
        0
             394
## 2
         1
             106
                         27.2
                                  1.49
with(lowDim_data, t.test(Y ~ A))
##
   Welch Two Sample t-test
##
## data: Y by A
## t = -6.7071, df = 132.81, p-value = 5.201e-10
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -13.728721 -7.475416
## sample estimates:
## mean in group 0 mean in group 1
          16.56747
                          27.16953
true low ATE<-2.0901
true_high_ATE<- -54.8558
# High dimension dataset
hd_time = system.time({ps_hd_est <- glm(A ~ . -Y,
                 family=binomial(),
                 data=highDim_data)
```

Use Logistic regression to get the propensity score

```
## # A tibble: 2 x 2
     treatment mean_prop_score
## *
         <int>
                         <dbl>
## 1
             0
                         0.253
                         0.466
labs <- paste("Actual treatment indicator:", c("Treated", "Control"))</pre>
ps_hd_df %>%
  mutate(treatment = ifelse(treatment == 1, labs[1], labs[2])) %>%
  ggplot(aes(x = prop_score), binwidth = 30) +
  geom_histogram(color = "white", bins=30) +
  facet_wrap(~treatment) +
  xlab("Probability of being exposed") +
  theme_bw()
```



```
summarise(mean_prop_score=mean(prop_score), .groups = "drop")
## # A tibble: 2 x 2
     treatment mean_prop_score
         <int>
                         <dbl>
## *
## 1
                         0.181
             0
## 2
             1
                         0.329
labs <- paste("Actual treatment indicator:", c("Treated", "Control"))</pre>
ps_ld_df %>%
  mutate(treatment = ifelse(treatment == 1, labs[1], labs[2])) %>%
  ggplot(aes(x = prop_score)) +
  geom_histogram(color = "white", bins=30) +
  facet_wrap(~treatment) +
  xlab("Probability of being exposed") +
  theme_bw()
```



Causal Inference Methods

Pairing 1

Algorithm: Propensity Matching
Distance Measure: Mahalanobis
Propensity Score Estimation: NA

Introduction of the algorithm

• Propensity Matching motivations:

key assumption: If a control and treated individual are identical before treatment, the probability of the outcome variable is equal where there is no treatment applied (Y_0) .

Matching on variables X thus ensures independence between treatment and Y_0 Matching on X is often infeasible especially where X is high dimensional. Instead, we use alternate distance measures to match on in order to circumvent this obstacle.

key parameters:

Distance Measure: Measure of similarity between 2 individuals Matching Method: How matching is conducted between individuals

- When estimating the ATE, either subclassification or full matching can be used. Full matching can be more effective because it optimizes a balance criterion, often leading to better balance. With full matching, it's also possible to exact match on some variables and match using the Mahalanobis distance, eliminating the need to estimate propensity scores. However, for large datasets, full matching may not be possible, in which case subclassification is a faster solution.
- The Mahalanobis distance is defined as:

$$D_{i}j = (X_{i} - X_{j})^{T} \Sigma^{-1} (X_{i} - X_{j})$$

Where Σ is the variance covariance matrix of X.

 Mahalanobis Distance does not require propensity score estimation and performs best with continuous variables.

implementation 1 code

```
mahalanobis.ate <- function(data){</pre>
  varnum=dim(data)[2]-2
  xnam <- paste0("V", 1:varnum)</pre>
  start_time <- Sys.time()</pre>
  match_full<-matchit(as.formula(paste("A ~ ", paste(xnam, collapse= "+"))),</pre>
                       data=data, method="full", distance="mahalanobis",
                       link = "logit", estimand = "ATE")
  data.fullMatching <- match.data(match full)</pre>
  x = data.fullMatching %>%
    group by(subclass,A) %>%
    summarise(mean_y = mean(Y), .groups = 'drop')
  group_ate = x %>%
    group_by(subclass) %>%
    summarise(treat_eff = mean_y[A == 1] - mean_y[A == 0], .groups = 'drop')
  group_n = data.fullMatching %>% group_by(subclass) %>% count()
  ate = sum(group_ate$treat_eff*group_n$n/nrow(data))
  end_time <- Sys.time()</pre>
  return(list(ATE=ate,running_time = end_time - start_time))
}
```

implementation 2 code

```
# Algorithm: Propensity Matching
# Distance Measure: Mahalanobis
# Propensity Score Estimation: NA
# Matching performed on low dimension dataset
```

```
covs = colnames(highDim_data)[-2:-1]
pair1_highdim_time = system.time({
  if (length(na.omit(highDim_data)) != length(highDim_data)) {
   print('There are null values in the dataset')
   break
   } else {
   pair_1 <- matchit(A ~ V1 + V2 + V3 + V4 + V5 + V6 + V7 + V8 + V9 + V10 + V11 + V12 + V13 + V14 + V1
                      data=highDim_data,
                      method="full",
                      distance="mahalanobis",
                      estimand = "ATE")
  }
# Preparing Data Frame based on Pairing 1 Matches High Dim
mpair_1 = match.data(pair_1)
#Estimating Treatment Effects using Pairing 1 Matches High Dim
mpair1_fit <- lm(Y ~ . -Y -weights -subclass,</pre>
                 data = mpair_1,
                 weights = weights)
pair1_highdim_ATE = round(as.numeric(mpair1_fit$coefficients['A']),4)
})
pair1_hd_time <- round(as.numeric(pair1_highdim_time[3]),3)</pre>
# Algorithm: Propensity Matching
# Distance Measure: Mahalanobis
# Propensity Score Estimation: NA
# Matching performed on low dimension dataset
covs = colnames(lowDim_data)[-2:-1]
pair1_lowdim_time = system.time({
  if (length(na.omit(lowDim data)) != length(lowDim data)) {
   print('There are null values in the dataset')
   break
   } else {
   pair_1 <- matchit(A ~ V1 + V2 + V3 + V4 + V5 + V6 + V7 + V8 + V9 + V10 + V11 + V12 + V13 + V14 + V1
                      data=lowDim_data,
                      method="full",
                      distance="mahalanobis",
                      estimand = "ATE")
 }
# Preparing Data Frame based on Pairing 1 Matches Low Dim
mpair_1 = match.data(pair_1)
# Estimating Treatment Effects using Pairing 1 Matches Low Dim
mpair1_fit <- lm(Y \sim . -Y - weights - subclass,
                 data = mpair_1,
                 weights = weights)
pair1_lowdim_ATE = round(as.numeric(mpair1_fit$coefficients['A']),4)
})
pair1_ld_time <- round(as.numeric(pair1_lowdim_time[3]),3)</pre>
High Dim data
```

```
## ATE for implementation 1 is: -63.32722 .   
## ATE error for implementation 1 is: 8.471422 .
```

```
## Processing time for implementation 1 is 1.083517 seconds.
##
## ATE for implementation 2 is: -57.1624 .
## ATE error for implementation 2 is: 2.3066 .
## Processing time for implementation 2 is 63.038 seconds.

Low Dim data
## ATE for implementation 1 is: 2.289205 .
## ATE error for implementation 1 is: -0.199105 .
## Processing time for implementation 1 is 0.34712 seconds.
##
## ATE for implementation 2 is: 2.2017 .
## ATE error for implementation 2 is: -0.1116 .
## Processing time for implementation 2 is 0.35 seconds.
```

Pairing 2

Algorithm: Propensity Matching

Distance Measure: Propensity Score

Propensity Score Estimation: Logistic Regression

Introduction of the algorithm

• The distance measure is defined as:

$$D_{ij} = |e_i - e_j|$$

where e_k is the propensity score for individual k.

• Propensity score definition:

$$p(\mathbf{X}) = P(exposed|\mathbf{X})$$

It is the probability of an individual being exposed (treated) given individual-specific characteristics.

implementation 1 code

```
Propensity.Score.ate <- function(data, methods, link){
   start_time <- Sys.time()
   match_full<-matchit(A ~ .-Y,data=data,method=methods,distance="glm",link = link, estimand = "ATE")
   data.fullMatching <- match.data(match_full)
   x = data.fullMatching %>% group_by(subclass,A) %>% summarise(mean_y = mean(Y), .groups = 'drop')
   group_ate = x %>% group_by(subclass) %>% summarise(treat_eff = mean_y[A == 1] - mean_y[A == 0], .group_ate = sum(group_ate$treat_eff*group_n$n/nrow(data))
   end_time <- Sys.time()
   return(list(ATE=round(ate,4),running_time = round(end_time - start_time, 3)))
}</pre>
```

implementation 2 code

```
# Algorithm: Propensity Matching
# Distance Measure: Propensity Score
# Propensity Score Estimation: Logistic Regression
pair2 <- function(data, method) {</pre>
  pair2_time = system.time({
    if (length(na.omit(data)) != length(data)) {
      print('There are null values in the dataset')
      break
      }
    else {
      pair_2 <- matchit(A ~ .-Y,</pre>
                         data=data,
                         method=method,
                         distance="glm",
                         link="logit",
                         estimand = "ATE")
    }
    # Preparing Data Frame based on Pairing 2 Matches
    mpair_2 = match.data(pair_2)
    # Estimating Treatment Effects using Pairing 2 Matches
    mpair2_fit <- lm(Y ~ . -Y -weights -subclass -distance,</pre>
                      data = mpair_2,
                      weights = weights)
    pair2_ATE = round(as.numeric(mpair2_fit$coefficients['A']),4)
    })
  pair2_time <- round(as.numeric(pair2_time[3]),3)</pre>
  return(list(ATE=pair2_ATE, running_time = pair2_time))
```

optimal full matching

```
pair2fhigh = Propensity.Score.ate(highDim_data, methods="full", link="logit")
pair2fhigh_alt = pair2(highDim_data, method="full")

High Dim data

## ATE for implementation 1 is: -59.5639 .

## ATE error for implementation 1 is: 4.7081 .

## Processing time for implementation 1 is 3.845 seconds.

##

## ATE for implementation 2 is: -60.4208 .

## ATE error for implementation 2 is: 5.565 .

## Processing time for implementation 2 is 4.094 seconds.

## Processing time for calculating the propensity score is 0.394 seconds.

pair2flow = Propensity.Score.ate(lowDim_data, methods="full", link="logit")
pair2flow_alt = pair2(lowDim_data, method="full")
```

```
Low Dim data
## ATE for implementation 1 is: 1.8199 .
## ATE error for implementation 1 is: 0.2702 .
## Processing time for implementation 1 is 0.216 seconds.
## ATE for implementation 2 is: 2.4034 .
## ATE error for implementation 2 is: -0.3133 .
## Processing time for implementation 2 is 0.198 seconds.
## Processing time for calculating the propensity score is 0.008 seconds.
subclassification
pair2shigh = Propensity.Score.ate(highDim_data, methods="subclass", link="logit")
pair2shigh_alt = pair2(highDim_data, method="subclass")
High Dim data
## ATE for implementation 1 is: -60.2547 .
## ATE error for implementation 1 is: 5.3989 .
## Processing time for implementation 1 is 0.482 seconds.
##
## ATE for implementation 2 is: -58.5693 .
## ATE error for implementation 2 is: 3.7135 .
## Processing time for implementation 2 is 0.48 seconds.
##
## Processing time for calculating the propensity score is 0.394 seconds.
pair2slow = Propensity.Score.ate(lowDim_data, methods="subclass", link="logit")
pair2slow_alt = pair2(lowDim_data, method="subclass")
Low Dim data
## ATE for implementation 1 is: 2.4099 .
## ATE error for implementation 1 is: -0.3198 .
## Processing time for implementation 1 is 0.023 seconds.
## ATE for implementation 2 is: 2.233 .
## ATE error for implementation 2 is: -0.1429 .
## Processing time for implementation 2 is 0.018 seconds.
```

Processing time for calculating the propensity score is 0.008 seconds.

##

Pairing 5

Algorithm: Propensity Matching

Distance Measure: Linear Propensity Score

Propensity Score Estimation: Logistic Regression

Introduction of the algorithm

The distance of Propensity Score is defined as:

$$D_{ij} = |e_i - e_j|$$

Obtained by applying the logit function on the Propensity Scores. Matching on the linear propensity score can be particularly effective in terms of reducing bias.

Linear propensity score matching is same with propensity score to entail forming matched sets of treated and untreated subjects who share a similar value of the propensity score. Once a matched sample has been formed, the treatment effect can be estimated by directly comparing outcomes between treated and untreated subjects in the matched sample. Once the effect of treatment has been estimated in the propensity score matched sample, the variance of the estimated treatment effect and its statistical significance can be estimated.

After the matched sets are obtained, Linear Propensity Score asks to calculate a "subclass effects" for each matched set/subclass, and then estimate overall ATE by an weighted average of the subclass effects where weights would be the number of individuals in each subclass.

Linear Propensity score performed well for low dimensional dataset as explained above for standard propensity score.

Linear Propensity Score didn't perform as well for high dimension as low dimensional dataset for the same reason discussed above for standard Propensity Score.

optimal full matching

```
pair5fhigh = Propensity.Score.ate(highDim_data, methods="full", link="linear.logit")
pair5flow = Propensity.Score.ate(lowDim_data, methods="full", link="linear.logit")
```

High Dim data

```
## ATE for high dimensional data is: -60.1412 .
```

- $\mbox{\tt \#\#}$ ATE error for high dimensional data is: 5.2854 .
- ## Processing time for calculating the propensity score is 0.394 seconds.
- ## Processing time for pairing 5 with full matching is 3.865 seconds.

Low Dim data

```
## ATE for low dimensional data is: 1.7251 .
```

- ## ATE error for low dimensional data is: 0.365 .
- ## Processing time for calculating the propensity score is 0.008 seconds.

Processing time for pairing 5 with full matching is 0.205 seconds.

```
pair5shigh = Propensity.Score.ate(highDim_data, methods="subclass", link="linear.logit")
pair5slow = Propensity.Score.ate(lowDim_data, methods="subclass", link="linear.logit")
```

subclassification

High Dim data

```
## ATE for high dimensional data is: -60.2547 .
```

ATE error for high dimensional data is: 5.3989 .

Processing time for calculating the propensity score is 0.394 seconds.

Processing time for runing pairing 5 is 0.495 seconds.

Low Dim data

```
## ATE for low dimensional data is: 2.4099 .
```

ATE error for low dimensional data is: -0.3198 .

Processing time for calculating the propensity score is 0.008 seconds.

Processing time for runing pairing 5 is 0.022 seconds.

Pairing 12

Algorithm: Doubly Robust Estimation
Distance Measure: Propensity Score

Propensity Score Estimation: Logistic Regression

Introduction of the algorithm

- The Doubly Robust Estimation has the smallest asymptotic variance. It remains consistent if the outcome models are wrong but the propensity model is right, or if the propensity model is wrong but the outcome models are right.
- Doubly Robust Estimator formula:

$$\hat{\Delta_{DR}} = N^{-1} \sum_{i=1}^{N} \frac{T_i Y_i - (T_i - \hat{e_i}) \hat{m_1}(X_i)}{\hat{e_i}} - N^{-1} \sum_{i=1}^{N} \frac{(1 - T_i) Y_i - (T_i - \hat{e_i}) \hat{m_0}(X_i)}{1 - \hat{e_i}}$$

where \hat{e}_i is the estimated propensity score for individual i, $\hat{m}_t(X)$ is a consistent estimate for E(Y|T=t,X) and is usually obtained by regressing the observed response Y on X in group t.

```
DoublyRobustEst <- function(data) {

X = data %>% select(-Y, -A)
n <- dim(data)[1]</pre>
```

```
start_time1 <- Sys.time()</pre>
  #get propensity scores by using logistic regression
  logit_model <- glm(A ~., data=data[,-1], family="binomial")</pre>
  propensity <- predict(logit_model, X, type="response")</pre>
  data$ps <- propensity</pre>
  end_time1 <- Sys.time()</pre>
  start time2 <- Sys.time()</pre>
  #split treatment and control group, and do regression for each group
  control <- data[data$A == 0, -2]</pre>
  treatment <- data[data$A == 1, -2]
  control_model <- lm(Y ~., data=control)</pre>
  treatment_model <- lm(Y ~., data=treatment)</pre>
  data$m0 <- predict(control_model, data[,-c(1,2)])</pre>
  data$m1 <- predict(treatment_model, data[,-c(1,2)])</pre>
  #calculate ATE
  ATE <- sum((data$A*data$Y-(data$A-data$ps)*data$m1)/data$ps)/n -
          sum(((1-data\$A)*data\$Y+(data\$A-data\$ps)*data\$m0)/(1-data\$ps))/n
  end_time2 <- Sys.time()</pre>
  time1 <- round(end_time1 - start_time1,3)</pre>
  time2 <- round(end_time2 - start_time1,3)</pre>
  df <- data.frame(cbind(round(ATE,4), time1, time2))</pre>
  return(df)
DRE.high <- DoublyRobustEst(highDim data)</pre>
```

```
DRE.high <- DoublyRobustEst(highDim_data)
diff.high <- true_high_ATE-DRE.high[,1]</pre>
```

highDim data

```
## ATE for high dimensional data is: -56.422 .
```

ATE error for high dimensional data is: 1.5662 .

Processing time for calculating the propensity score is 0.395 seconds.

Processing time for running the algorithm is 0.548 seconds.

```
DRE.low <- DoublyRobustEst(lowDim_data)
diff.low <- true_low_ATE-DRE.low[,1]</pre>
```

lowDim data

```
## ATE for low dimensional data is: 2.0678 .
```

ATE error for low dimensional data is: 0.0223 .

```
## Processing time for calculating the propensity score is 0.008 seconds.
```

Processing time for running the algorithm is 0.019 seconds.

Pairing 16

Algorithm: Stratification

Distance Measure: Propensity Score

Propensity Score Estimation: Logistic Regression

Introduction of the algorithm

• Stratification method definition:

$$\hat{\Delta_S} = \sum_{i=1}^K \frac{N_j}{N} \{ N_{1j}^{-1} \sum_{i=1}^K N T_i Y_i I(\hat{e_i} \in \hat{Q_j}) - N_{0j}^{-1} \sum_{i=1}^N (1 - T_i) Y_i I(\hat{e_i} \in \hat{Q_j}) \}$$

Where K is the number of strata, N_j is the number of individuals in stratum j, N_{ij} is the number of treated individuals in stratum j, and N_{0j} is the number of controlled individuals in stratum j.

Here we use K=7 as advised by the first article (Chan, Ge, Gershony, Hesterberg & Lambert).

```
n_strat <- 7
#High Dimensional Data
hd_prep <- system.time({
  #Stratify the Data by Propensity score
  hd_strat_divider <- n_strat/nrow(ps_hd_df);</pre>
  strat_hd_df <- ps_hd_df %>% arrange(prop_score) %>%
   mutate(stratum = as.integer((1:n() -.5)*hd_strat_divider ) + 1);
  #Differences in means
  diff_mean_hd <- strat_hd_df %>%
    group_by(treatment, stratum) %>%
    summarise(mean_treat = mean(Y), .groups = "keep") %>%
   mutate(mean_treat= if_else(treatment == 0, -mean_treat, mean_treat)) %>%
   ungroup(treatment) %>%
   mutate(diff_mean = sum(mean_treat)) %>%
   filter(treatment == 0) %>%
    select(stratum, diff_mean);
  #Weights for each stratum
  diff_weight_hd <- strat_hd_df %>%
    group_by(stratum) %>%
   tally() %>%
   mutate(weight = n/nrow(strat_hd_df)) %>%
   left_join(diff_mean_hd, by = "stratum") %>%
   select(weight, diff_mean);
  #Calculate the ATE
```

```
ATE_hd <- sum(diff_weight_hd$diff_mean * diff_weight_hd$weight)
})
time_hd <-sum(hd_prep)
#Low Dimensional Data
ld_prep <- system.time({</pre>
  #Stratify the Data by Propensity score
  ld strat divider <- n strat/nrow(ps ld df);</pre>
  strat_ld_df <- ps_ld_df %>% arrange(prop_score) %>%
    mutate(stratum = as.integer((1:n() -.5)*ld_strat_divider ) + 1);
  #Differences in means
  diff_mean_ld <- strat_ld_df %>%
    group_by(treatment, stratum) %>%
    summarise(mean_treat = mean(Y), .groups = "keep") %>%
    mutate(mean_treat= if_else(treatment == 0, -mean_treat, mean_treat)) %>%
    ungroup(treatment) %>%
    mutate(diff_mean = sum(mean_treat)) %>%
    filter(treatment == 0) %>%
    select(stratum, diff_mean);
  #Weights for each stratum
  diff_weight_ld <- strat_ld_df %>%
    group_by(stratum) %>%
    tally() %>%
    mutate(weight = n/nrow(strat_ld_df)) %>%
    left join(diff mean ld, by = "stratum") %>%
    select(weight, diff mean);
  #Calculate the ATE
  ATE_ld <- sum(diff_weight_ld$diff_mean * diff_weight_ld$weight)
})
time_ld <- sum(ld_prep)</pre>
```

Perform stratification and find difference of means and weights.

Alternate Method that performs regression on each stratum by Y with A and Vs. Then averages the coefficients

• alternate Method definition:

$$\hat{\Delta^{(j)}} = n_j^{-1} \sum_{i=1}^n I(\hat{e_i} \in \hat{Q_j}) \{ m^{(j)} (1, X_i, \alpha^{(j)}) - m^{(j)} (0, X_i, \alpha^{(j)}) \}$$

- This is an alternative method for calculating the ATE once the rows are stratified by performing regression on each stratum for (Y ~ A + Covariates) and using the mean of the coefficients among strata as the ATE.
- In larger numbers of strata, there is a risk of getting NAs for coefficients for the last few Covariates, if the number of observations in a stratum is less than the number of covariates + 1

for A in each stratum to get the ATE

```
n_strat <- 7
#High Dimensional Data
```

```
hd_prep_alt <- system.time({
  hd_strat_divider <- n_strat/nrow(ps_hd_df);</pre>
  #Append propensity score to the initial dataset
  hd_pscore <- matrix(ps_hd_df$prop_score, ncol = 1);</pre>
  hd_df_pscore <- cbind(highDim_data, hd_pscore);</pre>
  strat_hd_df_alt <- hd_df_pscore %>% arrange(hd_pscore) %>%
    mutate(stratum = as.integer((1:n() -.5)*hd_strat_divider ) + 1) %>%
    select(-hd_pscore);
  strat_hd_reg <- strat_hd_df_alt %>% group_by(stratum) %>%
    do(strat_reg = lm(Y ~ A + .-stratum, data = .)) %>%
    mutate(coefs = list(strat_reg[["coefficients"]])) %>%
    select(-strat_reg) %>%
    summarize(ajz = coefs[["A"]], .groups = "drop");
  #Calculate the ATE
  ATE_hd_alt <- mean(strat_hd_reg$ajz)</pre>
})
time_hd_alt <- sum(hd_prep_alt)</pre>
#Low Dimensional Data
ld_prep_alt <- system.time({</pre>
  ld_strat_divider <- n_strat/nrow(ps_ld_df);</pre>
  #Append propensity score to the initial dataset
  ld_pscore <- matrix(ps_ld_df$prop_score, ncol = 1);</pre>
  ld_df_pscore <- cbind(lowDim_data, ld_pscore);</pre>
  strat_ld_df_alt <- ld_df_pscore %>% arrange(ld_pscore) %>%
    mutate(stratum = as.integer((1:n() -.5)*ld_strat_divider ) + 1) %>%
    select(-ld_pscore);
  strat_ld_reg <- strat_ld_df_alt %>% group_by(stratum) %>%
    do(strat_reg = lm(Y ~ A + .-stratum, data = .)) %>%
    mutate(coefs = list(strat_reg[["coefficients"]])) %>%
    select(-strat_reg) %>%
    summarize(ajz = coefs[["A"]], .groups = "drop")
  #Calculate the ATE
  ATE_ld_alt <- mean(strat_ld_reg$ajz)</pre>
})
time_ld_alt <- sum(ld_prep_alt)</pre>
```

Calculates the ATE and process time

```
High Dim data
```

```
## ATE for stratification is: -59.85656 .
## ATE error for stratification is: 5.000763 .
```

```
## Processing time for stratification is 0.086 seconds.
##
## ATE for stratification alternative is: -53.97352 .
## ATE error for stratification alternative is: -0.88228 .
## Processing time for stratification is 0.534 seconds.
##
## Processing time for calculating the propensity score is 0.394 seconds.

Low Dim data
## ATE for stratification is: 2.012627 .
## ATE error for stratification is: 0.07747324 .
## Processing time for stratification is 0.078 seconds.
##
## ATE for stratification alternative is: 2.243051 .
## ATE error for stratification alternative is: -0.1529513 .
## Processing time for stratification is 0.104 seconds.
##
```

Processing time for calculating the propensity score is 0.008 seconds.

Summary

The table shows the ATEs, ATE errors, and times for all of our algorithms for both low dimensional and high dimensional datasets.

Summary

Algorithm / Results	Low Dimension Dataset			High Dimension Dataset		
	ATE	ATE Error	Time taken/s	ATE	ATE Error	Time taken/s
Mahalanobis Distance	2.2017	0.1116	0.44	-57.1624	2.3066	73.082
Propensity Score - Logistic Regression	2.2330	0.1429	0.009	-59.5639	4.812	4.7081
Linear Propensity Score - Logistic Regression	2.4050	-0.3198	0.0271	-60.1412	5.2854	3.1152
Stratification	2.0126	0.0775	0.028	-59.8565	5.0008	0.03
Alternate Stratification Method	2.2431	0.1530	0.045	-53.9735	0.8823	0.257
Doubly Robust Estimation	2.0678	0.0223	0.0113	-56.4220	1.5662	0.1573

Based on our calculation on ATE and processing time for the five algorithms that we implemented, we conclude that Doubly Robust Estimation works best on low dimension dataset and the stratification with alternative method is the most efficient on high dimension dataset.

Sources

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