Propensity Score Analysis in R

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Contents

1	Hov	w to Setup R and RStudio	:
	1.1	Download R, RStudio, and PSA-R	:
	1.2	Run the Code	
	1.3	Troubleshoot Package Errors	:
	1.4	PSA-R Session Info	
2	\mathbf{Gre}	eedy Nearest Neighbor Matching	5
	2.1	Load Packages	Ę
	2.2	Description of Dataset	Ę
	2.3	Load Data and Sort	
	2.4	Check Balance Before Matching	
	2.5	Greedy Nearest Neighbor Matching Without Replacement	
		2.5.1 Check Common Support	
		2.5.2 Check Balance	
	2.6	Greedy Nearest Neighbor Mahalanobis Distance Matching Without Replacement	
		2.6.1 Check Balance	
3	Dro	pensity Score Weighting	14
J	3.1	Load Packages	14
	$\frac{3.1}{3.2}$	Description of Dataset	
	3.2 3.3	Estimate ATE and ATT Weights	
	3.4	Load Data with Propensity Scores and Calculate Weights	
	$\frac{3.4}{3.5}$	Calculate Weights with the WeightIt Package	
	3.6	Outcome Analysis	
		3.6.1 Weighted Regression with ATE Weights	
	0.7	3.6.2 Weighted Regression with ATT Weights	
	3.7	Check Balance	16
4	Pro	pensity Score Estimation Using Generalized Boosted Regression	19
	4.1	Load Package	
	4.2	Load Data and Sort	
	4.3	Fit Generalized Boosted Regression Model	19
	4.4	Estimate Propensity Scores	20
	4.5	Plot Propensity Score Distributions	
	4.6	Summary Statistics of Propensity Scores	
	4.7	GBR Using the WeightIt Package	
		4.7.1 Check Balance	23
5	Mat	tching Estimators	24
		Load Packages	24
		Description of Dataset	24

	5.3	Load Data	24				
	5.4	Breusch-Pagan Test for Heteroskedasticity	24				
	5.5	Matching Estimators	25				
	5.6	Define Outcome (Y), Treatment Index (Tr), and Variables to Match On (X)	26				
	5.7	Get Estimators Individually	26				
	5.8	Get All Estimators	26				
	5.9	Specify Variables in the Bias-Corrected Matching	28				
		Check Balance	29				
6	Pra	ctice Problems	30				
	6.1	Practice 1, Problem 1: Generalized Boosted Regression	30				
		6.1.1 Description of Dataset	30				
		6.1.2 Load and Sort Data	30				
		6.1.3 Estimate Propensity Scores	30				
		6.1.4 Histogram and Density Plots of Propensity Scores	32				
		6.1.5 Boxplot of Propensity Scores	33				
	6.2	Practice 1, Problem 2: Propensity Score Weighting	34				
		6.2.1 Import Stata-Generated Weights for Comparison	34				
		6.2.2 Estimate ATE and ATT Weights	34				
		6.2.3 Outcome Analysis with ATE and ATT Weights	34				
		6.2.4 Check Balance	35				
		6.2.4.1 Hypothesis Tests	35				
		6.2.4.2 Standardized Mean Differences	38				
	6.3	Practice 1: Alternative Solution with the WeightIt Package	39				
		6.3.1 Check Balance	40				
		6.3.2 Outcome Analysis	41				
	6.4	Practice 2: Matching Estimators					
		6.4.1 Load Data	42				
		6.4.2 Breusch-Pagan Test for Heteroskedasticity	42				
		6.4.3 Define Outcome (Y), Treatment Index (Tr), and Variables to Match On (X)	42				
		6.4.4 Define Function for Matching	43				
		6.4.5 Get All Estimators	44				
7	App	pendix A: Function to Replicate Stata's Robust Standard Errors	45				
8	App	pendix B: Rosenbaum's Sensitivity Analysis	46				

1 How to Setup R and RStudio

1.1 Download R, RStudio, and PSA-R

- 1. Download the latest version of R: https://www.r-project.org/
- 2. Download the latest version of RStudio Desktop: https://www.rstudio.com/products/rstudio/download/
- 3. Download the two PSA-R zip files under "R Syntax": https://ssw.unc.edu/psa/

1.2 Run the Code

- 1. To view the code output without running it, extract the "PSA-R_Output.zip" file and open "index.html"
- 2. To run an individual section:
 - Extract "PSA-R_Code_Data.zip"
 - Open "PSA-R.Rproj"
 - Open the desired section code in the file browser (e.g., "01 Section 4.4.1.Rmd")
 - Install packages if necessary
 - Click on "Run All"
- 3. To knit the entire book into HTML output, click on "Build Book"

1.3 Troubleshoot Package Errors

If a line of code using a certain package is not working, try installing an older version of that package. See the output sessionInfo() below for package versions that are known to be compatible with the PSA-R code.

As of March 16, 2022, the latest version of PSweight (1.1.6) will only work if line 142 in "08_Section6.5.2.Rmd" is changed from:

```
data = sur_subclass1,
to
data = as.data.frame(sur_subclass1),
```

An alternative solution is to install an older version of PSweight that works (1.1.2):

```
packageVersion("PSweight")
detach("package:PSweight", unload = T)
remove.packages("PSweight")
library(devtools)
devtools::install_version("PSweight", version = "1.1.2",
    repos = "http://cran.us.r-project.org")
```

1.4 PSA-R Session Info

The following output for sessionInfo() lists package versions that are known to be compatible with the PSA-R code (if the fix for PSweight above is implemented).

```
R version 4.1.3 (2022-03-10)
Platform: x86_64-w64-mingw32/x64 (64-bit)
Running under: Windows 10 x64 (build 19043)

Matrix products: default

locale:
[1] LC_COLLATE=English_United States.1252 LC_CTYPE=English_United States.1252
[3] LC_MONETARY=English_United States.1252 LC_NUMERIC=C
```

```
[5] LC_TIME=English_United States.1252
attached base packages:
[1] stats
              graphics grDevices utils
                                             datasets methods
loaded via a namespace (and not attached):
 [1] tidyr_1.2.0
                             VGAM_1.1-6
                                                    splines_4.1.3
                                                                            foreach_1.5.2
 [5] carData 3.0-5
                             gam 1.20.1
                                                    gtools 3.9.2
                                                                            Formula 1.2-4
[9] assertthat 0.2.1
                             stats4_4.1.3
                                                    coin_1.4-2
                                                                            yaml 2.3.5
[13] numDeriv_2016.8-1.1
                             pillar_1.7.0
                                                    backports_1.4.1
                                                                            lattice_0.20-45
[17] glue_1.6.2
                             digest_0.6.29
                                                    colorspace_2.0-3
                                                                            sandwich_3.0-1
[21] gbm_2.1.8
                             htmltools_0.5.2
                                                    Matrix_1.4-0
                                                                            pkgconfig_2.0.3
[25] broom_0.7.12
                            haven_2.4.3
                                                    gmodels_2.18.1
                                                                            bookdown_0.25
                                                                            gdata_2.18.0
[29] purrr_0.3.4
                            mvtnorm_1.1-3
                                                    scales_1.1.1
[33] tibble_3.1.6
                             generics_0.1.2
                                                    car_3.0-12
                                                                            ggplot2_3.3.5
[37] sjlabelled_1.1.8
                             ellipsis_0.3.2
                                                    cobalt_4.3.2
                                                                            TH.data_1.1-0
[41] nnet_7.3-17
                             maxLik_1.5-2
                                                                            survival_3.2-13
                                                    cli_3.2.0
[45] magrittr_2.0.2
                             crayon_1.5.0
                                                    MatchIt_4.3.4
                                                                            evaluate_0.15
[49] fansi_1.0.2
                             MASS_7.3-55
                                                    SuperLearner_2.0-28
                                                                            forcats_0.5.1
[53] WeightIt_0.12.0
                             rsconnect_0.8.25
                                                    tools_4.1.3
                                                                            hms_1.1.1
[57] mitools_2.4
                             multcomp_1.4-18
                                                    matrixStats_0.61.0
                                                                            lifecycle_1.0.1
                             systemfit_1.1-24
[61] munsell_0.5.0
                                                    compiler_4.1.3
                                                                            rlang_1.0.2
[65] grid_4.1.3
                             Matching_4.9-11
                                                    iterators_1.0.14
                                                                            miscTools_0.6-26
[69] rbounds_2.1
                             rmarkdown_2.13
                                                    gtable_0.3.0
                                                                            codetools_0.2-18
[73] abind 1.4-5
                             DBI 1.1.2
                                                    R6 2.5.1
                                                                            nnls 1.4
                                                    dplyr_1.0.8
                                                                            fastmap_1.1.0
[77] zoo 1.8-9
                            knitr_1.37
[81] utf8 1.2.2
                             libcoin_1.0-9
                                                    insight_0.16.0
                                                                            sampleSelection_1.2-12
[85] modeltools_0.2-23
                             parallel_4.1.3
                                                    Rcpp_1.0.8.2
                                                                            vctrs_0.3.8
[89] tidyselect_1.1.2
                                                                            lmtest_0.9-39
                            xfun_0.30
                                                    PSweight_1.1.6
```

2 Greedy Nearest Neighbor Matching

2.1 Load Packages

The haven and sjlabelled packages are used to load and clean Stata data files (.dta); the MatchIt package contains functions for greedy matching; the cobalt package contains functions for balance checking; and the tidyverse package is loaded for its data manipulation functions.

```
library(haven)
library(sjlabelled)
library(cobalt)
library(MatchIt)
library(tidyverse)
library(kableExtra)
```

2.2 Description of Dataset

This dataset is a sample of 2,758 children from the National Survey of Child and Adolescent Well-Being (NSCAW). The treatment condition is aodserv or caregivers who received (aodserv = 1) or did not receive (aodserv = 0) substance abuse services. Two matching procedures are illustrated here. The full code contains 12 matching schemes and can be found in Section 5.8.1 of the PSA-R code.

2.3 Load Data and Sort

```
set.seed(1000)
gm_df <- haven::read_dta("data/chpt5_1_original.dta") %>%
  haven::zap_formats() %>%
  sjlabelled::remove_all_labels() %>%
  as_tibble() %>%
  add_column(runif = runif(nrow(.))) %>%
  arrange(runif) %>%
  select(-runif)
```

2.4 Check Balance Before Matching

Use chisq.test to check balance before matching:

```
## # A tibble: 17 x 3
## # Groups:
               variable [17]
##
      variable statistic p.value
##
      <chr>
                    <dbl>
                            <dbl>
                     2.97
##
                            0.085
  1 married
                            0.005
   2 educ
                    10.5
```

```
3 pov
                     11.3
                              0.023
##
    4 employ
                     23.1
                              0
##
##
    5 open
                     58.4
                              0
##
                              0.009
    6 race
                     11.5
##
    7 chdage
                     55.4
                              0
                      3.56
##
                              0.313
    8 cgage
##
    9 CRA47A
                     17.5
                              0
## 10 mental
                     92.5
                              0
## 11 arrest
                    127.
                              0
## 12 PSH17A
                    179.
                              0
## 13 maltx
                     49.7
                              0
                              0
## 14 ra
                    585.
## 15 cidi
                    157.
                              0
## 16 cgneed
                    139.
                              0
                              0
## 17 cwwrep
                   1240.
```

Alternatively, the cobalt package provides several convenient functions for assessing balance.

The standardized mean difference (SMD) is a commonly used balance measure. It is calculated as the difference in means of a covariate across the treatment groups, divided by the standard deviation in the treated group (ATT), the control group (ATC), or the pooled standard deviation (ATE). Stuart et al. (2013) recommend 0.1 or 0.25 as reasonable cut-offs for acceptable standardized biases.¹

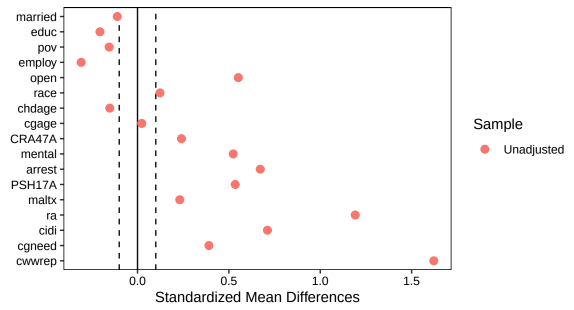
```
# Balance table
bal.tab(select(
   gm_df, married, educ, pov, employ, open, race, chdage, cgage, CRA47A,
   mental, arrest, PSH17A, maltx, ra, cidi, cgneed, cwwrep
),
treat = gm_df$aodserv,
s.d.denom = "treated",
threshold = .1
)
```

```
## Balance Measures
              Type Diff.Un
                               M.Threshold.Un
## married
           Binary -0.0485
                               Balanced, <0.1
## educ
           Contin. -0.2055 Not Balanced, >0.1
           Contin. -0.1550 Not Balanced, >0.1
## pov
            Binary -0.1472 Not Balanced, >0.1
## employ
## open
            Binary 0.2327 Not Balanced, >0.1
           Contin. 0.1231 Not Balanced, >0.1
## race
## chdage
           Contin. -0.1517 Not Balanced, >0.1
## cgage
           Contin. 0.0232
                               Balanced, <0.1
## CRA47A
            Binary 0.1184 Not Balanced, >0.1
## mental
            Binary 0.2619 Not Balanced, >0.1
## arrest
            Binary 0.3254 Not Balanced, >0.1
## PSH17A
                    0.2512 Not Balanced, >0.1
            Binary
## maltx
           Contin.
                    0.2312 Not Balanced, >0.1
## ra
                    0.5598 Not Balanced, >0.1
            Binary
## cidi
            Binary
                    0.3482 Not Balanced, >0.1
                    0.1494 Not Balanced, >0.1
## cgneed
            Binary
## cwwrep
            Binary 0.7036 Not Balanced, >0.1
##
```

¹Stuart, E. A., Lee, B. K., & Leacy, F. P. (2013). Prognostic score—based balance measures for propensity score methods in comparative effectiveness research. *Journal of Clinical Epidemiology*, 66(8 0), S84-S90.e1. https://doi.org/10.1016/j.jclinepi.201 3.01.013

```
## Balance tally for mean differences
##
                      count
## Balanced, <0.1
## Not Balanced, >0.1
## Variable with the greatest mean difference
   Variable Diff.Un
                         M.Threshold.Un
      cwwrep 0.7036 Not Balanced, >0.1
##
##
## Sample sizes
       Control Treated
          2460
                   298
## All
# Love plot
love.plot(select(
  gm_df, married, educ, pov, employ, open, race, chdage, cgage, CRA47A,
  mental, arrest, PSH17A, maltx, ra, cidi, cgneed, cwwrep
),
treat = gm_df$aodserv,
binary = "std",
s.d.denom = "treated",
threshold = .1
  labs(title = "Covariate Balance Before Matching")
```

Covariate Balance Before Matching



2.5 Greedy Nearest Neighbor Matching Without Replacement

By default, the MatchIt::matchit() function performs greedy nearest neighbor matching without replacement, therefore the method = "nearest" and replace = F arguments do not need to be specified.

To avoid dissimilar matches, we can constrain matches so that the absolute distance of propensity scores between two participants is less than a specified tolerance for matching or a caliper. The width of the caliper is by default in standard deviation units and can be specified using the caliper argument. A wide caliper may result in more matches and a larger sample, but inexact matching may occur as indicated by large distances on the propensity score between the treated and nontreated cases. Using varying caliper sizes can test the sensitivity of the findings. Here we use a caliper size of a quarter of a standard deviation, which is suggested by Rosenbaum and Rubin (1985).

The order of the matching can be specified using the m.order argument. Finally, the logit of the predicted probability from a logistic regression model can be supplied to the distance argument. The logit of the predicted probability is used, because the logit is approximately normally distributed.

```
# Logistic regression specification
(gm_f <- cobalt::f.build("aodserv", select(gm_df, PSH17A:other, -aodserv)))</pre>
## aodserv ~ PSH17A + CRA47A + married + high + bahigh + poverty2 +
##
       poverty3 + poverty4 + poverty5 + employ + open + black +
##
       hispanic + natam + cgrage1 + cgrage2 + cgrage3 + chdage1 +
##
       chdage2 + chdage3 + mental + arrest + sexual + provide +
##
       supervis + other
## <environment: 0x0000000242122a0>
# Calculate the logit of the predicted probability as the propensity score
gm_psm <- glm(gm_f, data = gm_df, family = binomial)</pre>
gm_ps <- predict(gm_psm, newdata = gm_df, type = "response")</pre>
gm_ps_logit <- log((1 - gm_ps) / gm_ps)</pre>
# Greedy nearest neighbor matching without replacement
set.seed(1000)
(gm_out <- MatchIt::matchit(</pre>
  gm_f,
 data = gm_df,
  distance = gm ps logit,
 m.order = "random",
  caliper = .25
))
## A matchit object
## - method: 1:1 nearest neighbor matching without replacement
## - distance: User-defined [caliper]
## - caliper: <distance> (0.311)
## - number of obs.: 2758 (original), 574 (matched)
   - target estimand: ATT
   - covariates: PSH17A, CRA47A, married, high, bahigh, poverty2, poverty3, poverty4, poverty5, employ
Notice that a limitation of this matching scheme is that it reduces the sample size from 2758 to 574—287
```

2.5.1 Check Common Support

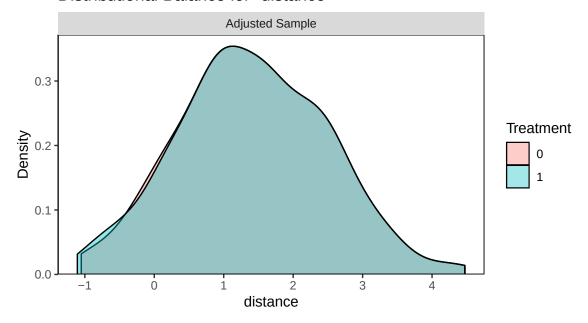
cases in the control group and 287 cases in the treated group.

Greedy matching is criticized because it requires a sizeable common-support region to work. The common support region is defined as the region bounded by the maximum value of estimated propensity scores for

the treated participants and by the minimum value of the estimated propensity scores for the nontreated participants. In this example, a sizeable common-support region exists.

```
cobalt::bal.plot(gm_out, var.name = "distance")
```

Distributional Balance for "distance"



2.5.2 Check Balance

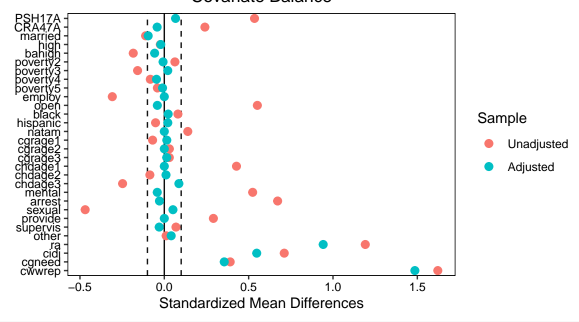
Covariate balance can be assessed using hypothesis tests, such as chisq.test:

```
gm_out_data <- MatchIt::match.data(gm_out)
chisq.test(gm_out_data$ra, gm_out_data$aodserv)</pre>
```

```
##
## Pearson's Chi-squared test with Yates' continuity correction
##
## data: gm_out_data$ra and gm_out_data$aodserv
## X-squared = 111.7, df = 1, p-value < 2.2e-16</pre>
```

The object from matchit() can be directly used in cobalt functions to produce balance tables and plots. To specify additional variables for which to display balance, use the argument addl in conjunction with data.

Covariate Balance



```
## Call
   MatchIt::matchit(formula = gm_f, data = gm_df, distance = gm_ps_logit,
##
       m.order = "random", caliper = 0.25)
##
##
## Balance Measures
                Type Diff.Un Diff.Adj
                                              M.Threshold
## distance Distance -1.3021
                              -0.0040
                                           Balanced, <0.1
## PSH17A
              Binary 0.5347
                               0.0667
                                           Balanced, <0.1
## CRA47A
              Binary 0.2406
                              -0.0425
                                           Balanced, <0.1
## married
              Binary -0.1108
                              -0.0955
                                           Balanced, <0.1
## high
              Binary -0.0236
                              -0.0211
                                           Balanced, <0.1
## bahigh
              Binary -0.1840
                              -0.0574
                                           Balanced, <0.1
              Binary 0.0628
## poverty2
                              -0.0073
                                           Balanced, <0.1
## poverty3
              Binary -0.1584
                               0.0204
                                           Balanced, <0.1
## poverty4
              Binary -0.0829
                              -0.0463
                                           Balanced, <0.1
## poverty5
              Binary -0.0410
                              -0.0103
                                           Balanced, <0.1
## employ
              Binary -0.3082
                               0.0000
                                           Balanced, <0.1
## open
              Binary 0.5516
                              -0.0413
                                           Balanced, <0.1
## black
              Binary 0.0807
                               0.0231
                                           Balanced, <0.1
              Binary -0.0519
                               0.0207
                                           Balanced, <0.1
## hispanic
## natam
              Binary 0.1392
                               0.0000
                                           Balanced, <0.1
                               0.0147
                                           Balanced, <0.1
## cgrage1
              Binary -0.0694
## cgrage2
              Binary 0.0300
                               0.0000
                                           Balanced, <0.1
              Binary 0.0297
                                           Balanced, <0.1
## cgrage3
                               0.0143
                               0.0000
## chdage1
              Binary 0.4279
                                           Balanced, <0.1
## chdage2
                               0.0101
                                           Balanced, <0.1
              Binary -0.0856
## chdage3
              Binary -0.2473
                               0.0864
                                           Balanced, <0.1
## mental
              Binary 0.5238
                              -0.0418
                                           Balanced, <0.1
## arrest
              Binary 0.6718
                              -0.0288
                                           Balanced, <0.1
                               0.0512
## sexual
              Binary -0.4690
                                           Balanced, <0.1
```

```
Binary 0.2909 0.0000
## provide
                                        Balanced, <0.1
## supervis
             Binary 0.0690 -0.0303
                                        Balanced, <0.1
## other
             Binary 0.0110 0.0409
                                        Balanced, <0.1
## ra
             Binary 1.1915
                             0.9419 Not Balanced, >0.1
             Binary 0.7110
                             0.5478 Not Balanced, >0.1
## cidi
## cgneed
             Binary 0.3907
                             0.3554 Not Balanced, >0.1
             Binary 1.6213 1.4853 Not Balanced, >0.1
## cwwrep
##
## Balance tally for mean differences
##
                     count
## Balanced, <0.1
                        27
## Not Balanced, >0.1
## Variable with the greatest mean difference
## Variable Diff.Adj
                           M.Threshold
     cwwrep 1.4853 Not Balanced, >0.1
##
##
## Sample sizes
##
            Control Treated
## All
              2460
                        298
                287
## Matched
                        287
## Unmatched
               2173
                       11
```

2.6 Greedy Nearest Neighbor Mahalanobis Distance Matching Without Replacement

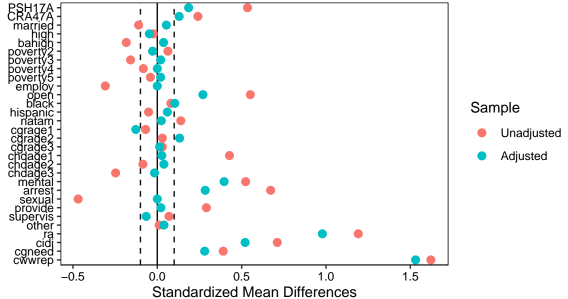
Here we perform Mahalanobis distance matching without replacement and without including an estimated propensity score.

```
set.seed(1000)
(gm_out2 <- MatchIt::matchit(gm_f, data = gm_df,
    method = "nearest", distance = "mahalanobis"))

## A matchit object
## - method: 1:1 nearest neighbor matching without replacement
## - distance: Mahalanobis
## - number of obs.: 2758 (original), 596 (matched)
## - target estimand: ATT
## - covariates: PSH17A, CRA47A, married, high, bahigh, poverty2, poverty3, poverty4, poverty5, employ</pre>
```

2.6.1 Check Balance

Covariate Balance



```
## Call
   MatchIt::matchit(formula = gm_f, data = gm_df, method = "nearest",
##
       distance = "mahalanobis")
##
## Balance Measures
##
              Type Diff.Adj
                                   M.Threshold
## PSH17A
           Binary
                     0.1857 Not Balanced, >0.1
           Binary
                     0.1295 Not Balanced, >0.1
## CRA47A
## married Binary
                     0.0537
                                Balanced, <0.1
```

```
## high
            Binary
                    -0.0474
                                 Balanced, <0.1
## bahigh
                      0.0368
                                 Balanced, <0.1
            Binary
                                 Balanced, <0.1
## poverty2 Binary
                     -0.0281
## poverty3 Binary
                      0.0197
                                 Balanced, <0.1
## poverty4 Binary
                      0.0000
                                 Balanced, <0.1
                                 Balanced, <0.1
## poverty5 Binary
                      0.0199
                                 Balanced, <0.1
## employ
            Binary
                      0.0000
                      0.2705 Not Balanced, >0.1
## open
            Binary
## black
            Binary
                      0.1037 Not Balanced, >0.1
## hispanic Binary
                      0.0597
                                 Balanced, <0.1
## natam
            Binary
                      0.0238
                                 Balanced, <0.1
                     -0.1276 Not Balanced, >0.1
## cgrage1
            Binary
## cgrage2
            Binary
                      0.1320 Not Balanced, >0.1
## cgrage3
            Binary
                      0.0137
                                 Balanced, <0.1
## chdage1
                      0.0269
                                 Balanced, <0.1
            Binary
## chdage2
            Binary
                      0.0390
                                 Balanced, <0.1
## chdage3
            Binary
                     -0.0166
                                 Balanced, <0.1
## mental
                      0.3960 Not Balanced, >0.1
            Binary
## arrest
            Binary
                      0.2841 Not Balanced, >0.1
## sexual
            Binary
                      0.0000
                                 Balanced, <0.1
## provide
            Binary
                      0.0213
                                 Balanced, <0.1
## supervis Binary
                    -0.0656
                                 Balanced, <0.1
                                 Balanced, <0.1
## other
            Binary
                      0.0394
## ra
            Binary
                      0.9786 Not Balanced, >0.1
## cidi
            Binary
                      0.5207 Not Balanced, >0.1
            Binary
## cgneed
                      0.2808 Not Balanced, >0.1
                      1.5310 Not Balanced, >0.1
## cwwrep
            Binary
##
## Balance tally for mean differences
##
                       count
## Balanced, <0.1
                          18
## Not Balanced, >0.1
                          12
##
## Variable with the greatest mean difference
##
    Variable Diff.Adj
                              M.Threshold
                1.531 Not Balanced, >0.1
##
      cwwrep
##
## Sample sizes
##
             Control Treated
## All
                2460
                          298
## Matched
                  298
                          298
## Unmatched
                2162
                            0
```

As seen above, balance has not been achieved in multiple covariates. According to Stuart (2010), "the Mahalanobis distance can work quite well when there are relatively few covariates (fewer than 8), but it does not perform as well when the covariates are not normally distributed or there are many covariates."²

 $^{^2} Stuart,$ E. A. (2010). Matching Methods for Causal Inference: A Review and a Look Forward. Statistical Science, 25(1), 1–21. https://doi.org/10.1214/09-STS313

3 Propensity Score Weighting

3.1 Load Packages

Propensity score weighting can be accomplished with base R. However, we need the lmtest and sandwich packages to estimate clustered covariance matrices in this example. Using these packages, we can obtain estimates and standard errors that are identical to Stata's regress program.

library(lmtest)
library(sandwich)

3.2 Description of Dataset

This dataset is from a study that investigates intergenerational dependence on welfare and its relation to child academic achievement.³

The dependent variable is lwss97 or "letter-word identification" score, and the treatment condition is kuse or children who used Aid to Families With Dependent Children (AFDC). The covariates are:

- male: Child's Gender: Male (Reference: Female)
- black: Child's Race: African American (Reference: Other)
- age97: Child's Age in 1997
- pcged97: Caregiver's Education in 1997 (Years of Schooling)
- mratio96: Ratio of Family Income to Poverty Line in 1996

Additionally, pcg_id is a cluster variable that identifies children nested within families.

3.3 Estimate ATE and ATT Weights

Separate weights need to be calculated for estimating the average treatment effect (ATE) and the average treatment effect for the treated (ATT).

For ATE, the weight estimates are calculated as follows for the treatment group:

$$\omega = \frac{1}{\hat{e}(x)}$$

And for the control group:

$$\omega = \frac{1}{1 - \hat{e}(x)}$$

For ATT, the weight is 1 for a treated case. The weight for a comparison case is:

$$\omega = \frac{\hat{e}(x)}{1 - \hat{e}(x)}$$

³Hofferth, S., Stafford, F. P., Yeung, W. J., Duncan, G. J., Hill, M. S., Lepkowski, J., et al. (2001). *Panel study of income dynamics*, 1968–1999: Supplemental files (computer file), ICPSR version. Ann Arbor: University of Michigan Survey Research Center.

3.4 Load Data with Propensity Scores and Calculate Weights

3.5 Calculate Weights with the WeightIt Package

```
# Load Package
library(WeightIt)

# Estimate ATE and ATT weights and Compare with Previous Results
ate_w2 <- WeightIt::get_w_from_ps(ps = psw_df$ps, treat = psw_df$kuse, estimand = "ATE")
table(ate_w2 == psw_df$ate_w)

##
## TRUE
## 1003
att_w2 <- WeightIt::get_w_from_ps(ps = psw_df$ps, treat = psw_df$kuse, estimand = "ATT")
table(att_w2 == (psw_df$ate_w * psw_df$ps))</pre>
##
## TRUE
## 1003
```

3.6 Outcome Analysis

3.6.1 Weighted Regression with ATE Weights

After creating the weights, use the weights argument in lm() to run a weighted outcome analysis and lmtest::coeftest() to control for clustering effects.

This analysis showed that children who used Aid to Families With Dependent Children (AFDC) had an average letter-word identification score that was 5.16 points lower than children who never used AFDC, p < .01.

```
psw_ate <- lm(lwss97 ~ kuse + male + black + age97 + pcged97 + mratio96,
    data = psw_df, weights = ate_w)
lmtest::coeftest(psw_ate, vcov. = vcovCL(psw_ate, cluster = psw_df$pcg_id))</pre>
```

```
##
## t test of coefficients:
##
##
              Estimate Std. Error t value Pr(>|t|)
                          4.83032 17.4315 < 2.2e-16 ***
## (Intercept) 84.19992
## kuse
              -5.16399
                          1.42438 -3.6254 0.0003031 ***
## male
              -1.62201
                          1.09186 -1.4855 0.1377180
## black
              -2.49898
                          1.34670 -1.8556 0.0638009 .
## age97
               0.73868
                          0.18075 4.0867 4.727e-05 ***
## pcged97
               0.99264
                          0.35596 2.7886 0.0053938 **
                          0.32220 3.5337 0.0004286 ***
## mratio96
              1.13856
## ---
```

```
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
```

3.6.2 Weighted Regression with ATT Weights

When considering only individuals assigned to the treatment condition, children who used AFDC had an average letter-word identification score that was 4.62 points lower than children who never used AFDC, p < .01.

```
psw_att <- lm(lwss97 ~ kuse + male + black + age97 + pcged97 + mratio96,
  data = psw_df, weights = att_w)
lmtest::coeftest(psw_att, vcov. = vcovCL(psw_att, cluster = psw_df$pcg_id))
##
## t test of coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 85.29467
                          5.05331 16.8790 < 2.2e-16 ***
               -4.62058
                          1.41182 -3.2728 0.001102 **
## kuse
## male
               -1.58995
                          1.14705 -1.3861 0.166020
## black
               -2.74605
                          1.41605 -1.9392 0.052756 .
## age97
               0.61577
                          0.20001
                                   3.0787
                                           0.002136 **
## pcged97
               0.92698
                          0.36718 2.5246 0.011738 *
## mratio96
                1.26018
                          0.33556 3.7555 0.000183 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

3.7 Check Balance

To assess balance before and after propensity score weighting, use weighted logistic regression for dummy covariates and weighted simple regression for continuous covariates. Some examples are included below, and the full code can be found in Section 7.3.1 of the PSA-R code.

In model psw_c3 below, the treatment dummy variable is significant, meaning that there is no sufficient balance after the propensity score weighting.

To assess balance before propensity score weighting, remove the weights argument.

```
psw_c1 <- glm(male ~ kuse, family = quasibinomial, data = psw_df, weights = ate_w)
lmtest::coeftest(psw_c1, vcov. = vcovCL(psw_c1, cluster = psw_df$pcg_id))
## z test of coefficients:
##
                Estimate Std. Error z value Pr(>|z|)
##
## (Intercept)
               0.142911
                           0.075538 1.8919
                                              0.0585
## kuse
               -0.060271
                           0.150143 -0.4014
                                              0.6881
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
psw_c2 <- glm(male ~ kuse, family = quasibinomial, data = psw_df, weights = att_w)</pre>
lmtest::coeftest(psw_c2, vcov. = vcovCL(psw_c2, cluster = psw_df$pcg_id))
##
## z test of coefficients:
##
##
                Estimate Std. Error z value Pr(>|z|)
                           0.077410 1.9703
## (Intercept) 0.152522
               -0.079497
                           0.147480 -0.5390
                                              0.5899
## kuse
```

```
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
psw_c3 <- lm(age97 ~ kuse, weights = ate_w, data = psw_df)</pre>
lmtest::coeftest(psw_c3, vcov. = vcovCL(psw_c3, cluster = psw_df$pcg_id))
##
## t test of coefficients:
##
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 6.51389 0.10900 59.7615 < 2.2e-16 ***
## kuse
               0.61064
                          0.21883 2.7905 0.005362 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
psw_c4 <- lm(age97 ~ kuse, weights = att_w, data = psw_df)</pre>
lmtest::coeftest(psw_c4, vcov. = vcovCL(psw_c4, cluster = psw_df$pcg_id))
##
## t test of coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 6.55501 0.11499 57.0057 < 2e-16 ***
               0.56178
                          0.21939 2.5606 0.01059 *
## kuse
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Alternatively, balance can be assessed using standardized mean differences:
# ATE
cobalt::bal.tab(
 x = select(psw_df, male, black, age97, pcged97, mratio96),
 treat = psw_df$kuse,
 weights = psw_df$ate_w,
 binary = "std",
 continuous = "std";
 s.d.denom = "pooled",
 un = T,
 stats = c("mean.diffs"),
 thresholds = c(m = .1)
)
## Balance Measures
              Type Diff.Un Diff.Adj
                                           M.Threshold
## male
            Binary -0.0335 -0.0301
                                        Balanced, <0.1
## black
            Binary 0.9343 0.8006 Not Balanced, >0.1
           Contin. 0.2196 0.2181 Not Balanced, >0.1
## age97
## pcged97 Contin. -0.9067 -0.8244 Not Balanced, >0.1
## mratio96 Contin. -1.0158 -0.8775 Not Balanced, >0.1
##
## Balance tally for mean differences
                     count
## Balanced, <0.1
## Not Balanced, >0.1
## Variable with the greatest mean difference
## Variable Diff.Adj
                        M.Threshold
```

```
## mratio96 -0.8775 Not Balanced, >0.1
##
## Effective sample sizes
           Control Treated
## Unadjusted 729.
                     274.
## Adjusted 725.43 257.71
# ATT
cobalt::bal.tab(
 x = select(psw_df, male, black, age97, pcged97, mratio96),
 treat = psw_df$kuse,
 weights = psw_df$ate_w,
 binary = "std",
 continuous = "std",
 s.d.denom = "treated",
 un = T,
 stats = c("mean.diffs"),
 thresholds = c(m = .1)
## Balance Measures
     Type Diff.Un Diff.Adj
                                        M.Threshold
          Binary -0.0335 -0.0301 Balanced, <0.1
## male
         Binary 1.0129
                           0.8680 Not Balanced, >0.1
## black
## age97 Contin. 0.2181 0.2167 Not Balanced, >0.1
## pcged97 Contin. -1.0082 -0.9167 Not Balanced, >0.1
## mratio96 Contin. -2.1090 -1.8217 Not Balanced, >0.1
## Balance tally for mean differences
                    count
## Balanced, <0.1
## Not Balanced, >0.1
##
## Variable with the greatest mean difference
## Variable Diff.Adj
                      M.Threshold
## mratio96 -1.8217 Not Balanced, >0.1
##
## Effective sample sizes
             Control Treated
## Unadjusted 729. 274.
## Adjusted 725.43 257.71
```

4 Propensity Score Estimation Using Generalized Boosted Regression

4.1 Load Package

Generalized boosted regression (GBR) requires the gbm package.

```
library(gbm)
```

4.2 Load Data and Sort

Generalized boosted regression is an iterative method for creating propensity scores. Therefore, to create reproducible results, we need to use the set.seed() function.

After importing the data, missing data is deleted listwise, and the data is sorted randomly. According to the gbm package vignette, if the data is sorted in a systematic way, then the data should be shuffled before running gbm.

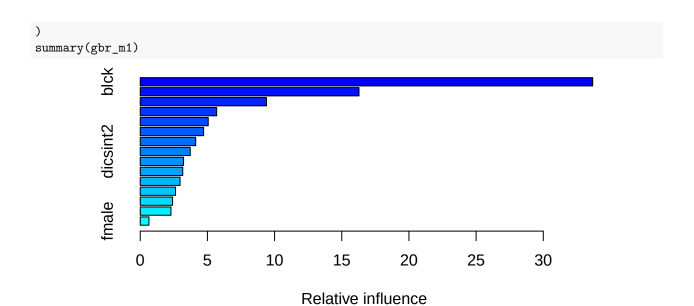
4.3 Fit Generalized Boosted Regression Model

The gbm::gbm() function has many arguments that can be fine-tuned. See ?gbm for a detailed description of each argument.

A summary of the fitted model provides us with *relative influence*, which is the percentage of log likelihood explained by each input variable. The percentages of influence for all predictor variables sum to 100%.

The GBM showed that blck had the strongest influence on the likelihood function (33.7%), followed by ageyc (16.3%) and draggr2 (9.4%).

```
(gbr_f <- cobalt::f.build("intbl", select(gbr_df, -intbl)))</pre>
## intbl ~ ageyc + fmale + blck + whit + hisp + pcedu + ipovl +
##
       pcemft + fthr + dicsagg2 + dicsint2 + dccereg2 + dccscom2 +
##
       dccpros2 + draggr2
## <environment: 0x000000027498168>
set.seed(1000)
gbr_m1 <- gbm::gbm(</pre>
 formula = gbr_f,
  data = gbr_df,
 distribution = "bernoulli",
 n.trees = 1000, # number of trees to fit
  train.fraction = 0.8, # a random 80% subsample for estimation
  interaction.depth = 4, # allow all four-way interactions
  shrinkage = 0.0005 # small shrinkage to ensure smooth fit
```



```
##
                 var
                        rel.inf
## blck
                blck 33.6768868
## ageyc
               ageyc 16.2834103
             draggr2 9.3962151
## draggr2
## whit
                whit
                     5.6895119
## ipovl
               ipovl 5.0614691
## pcemft
              pcemft
                      4.7179839
## pcedu
               pcedu
                      4.1280063
## dicsint2 dicsint2
                      3.7223185
## dicsagg2 dicsagg2
                      3.2228567
## dccscom2 dccscom2 3.1648839
                      2.9647178
## dccereg2 dccereg2
## dccpros2 dccpros2
                      2.6269129
## hisp
                hisp
                     2.4079157
## fthr
                fthr
                      2.2874508
## fmale
               fmale 0.6494603
```

4.4 Estimate Propensity Scores

After fitting the model, estimate propensity scores using the predict.gbm() function.

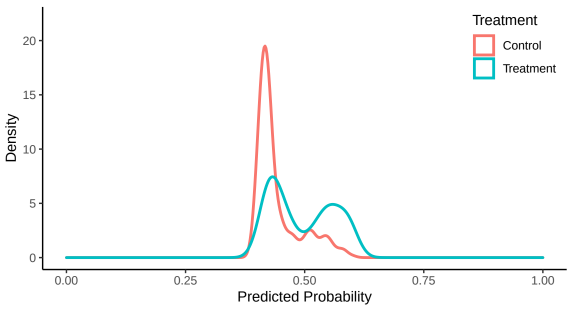
```
psb <- gbm::predict.gbm(gbr_m1, data = gbr_df, type = "response")
head(psb)</pre>
```

[1] 0.4441531 0.4405950 0.5288983 0.5958331 0.3985049 0.5562076

4.5 Plot Propensity Score Distributions

As seen in the figure below, the propensity scores estimated by GBM has sufficient overlap between the control and treatment groups (i.e., "common support"), and the distributions are similar.

Propensity Scores Using Generalized Boosted Regression



4.6 Summary Statistics of Propensity Scores

```
summary(psb)
## Min. 1st Qu. Median Mean 3rd Qu. Max.
## 0.3913 0.4189 0.4383 0.4674 0.5202 0.6082
```

4.7 GBR Using the WeightIt Package

As an alternative to the gbm package, the WeightIt package can also fit GBR models.

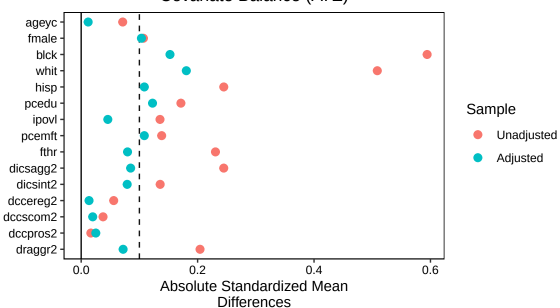
```
set.seed(1000)
(gbr_m2 <- WeightIt::weightit(</pre>
 formula = gbr_f,
 data = gbr_df,
 method = "gbm",
 estimand = "ATE",
 distribution = "bernoulli",
 stop.method = "es.mean",
 n.trees = 10000, # different
 nTrain = 0.8 * nrow(gbr_df), # different
 interaction.depth = 4,
 shrinkage = 0.0005
))
## A weightit object
## - method: "gbm" (propensity score weighting with GBM)
## - number of obs.: 603
## - sampling weights: none
## - treatment: 2-category
## - estimand: ATE
## - covariates: ageyc, fmale, blck, whit, hisp, pcedu, ipovl, pcemft, fthr, dicsagg2, dicsint2, dccer
# ATE Weights
head(gbr_m2$weights)
## [1] 2.644442 1.645871 1.515870 1.287579 1.271966 1.261381
```

4.7.1 Check Balance

Using a standardized mean difference cut-off point of 0.1, it can be seen below that balance has been achieved in most, but not all, of the covariates:

```
cobalt::love.plot(gbr_m2,
   thresholds = c(m = .1),
   binary = "std", abs = T, drop.distance = T
) +
   labs(title = "Covariate Balance (ATE)")
```





5 Matching Estimators

5.1 Load Packages

The Breusch-Pagan test against heteroskedasticity requires the lmtest package, and a variety of matching estimators are implemented in the Matching package.

```
library(lmtest)
library(Matching)
library(broom)
select <- dplyr::select</pre>
```

5.2 Description of Dataset

This example uses the 1997 Child Development Supplement (CDS) to the Panel Study of Income Dynamics (PSID) and the core PSID annual data from 1968 to 1997.

The dependent variable in this dataset is pcss97, a passage comprehension score. Higher scores on this measure indicate higher academic achievement. The treatment variable is kuse or children who ever used Aid to Families With Dependent Children (AFDC). The covariates or matching variables are:

- male: Child's Gender: Male (Reference: Female)
- black: Child's Race: African American (Reference: Other)
- age97: Child's Age in 1997
- pcged97: Caregiver's Education in 1997 (Years of Schooling)
- mratio96: Ratio of Family Income to Poverty Line in 1996
- pcg_adc: Caregiver's History of Using Welfare (Number of Years)

5.3 Load Data

```
me_df <- read_dta("data/cds_pcss97.dta") %>%
  haven::zap_formats() %>%
  sjlabelled::remove_all_labels() %>%
  as_tibble()
head(me_df) %>%
  kbl(booktabs = T, linesep = "", digits = 2) %>%
  kable_styling(position = "center") %>%
  kable_styling(latex_options = c("striped", "hold_position"))
```

kid	pcg_id	age97	pcss97	k_adc	pcged 97	pcg_adc	mratio 96	black	male	kuse
4180	4179	12	93	7.69	9	0	0.41	0	0	1
5032	5170	12	122	0.00	12	0	5.00	0	0	0
7041	7030	6	93	0.00	11	2	1.07	0	1	0
10033	10177	7	101	8.33	12	0	2.40	1	0	1
10034	10006	6	127	0.00	14	0	1.00	1	0	0
14030	14172	11	103	0.00	12	0	3.77	0	1	0

5.4 Breusch-Pagan Test for Heteroskedasticity

The homoskedastic variance estimator assumes that the unit-level treatment effect is constant and that the conditional variance of $Y_i(w)$ given X_i does not vary with either covariates or the treatment. The Breusch-Pagan test is performed for each of the seven independent variables and showed that child's age was statistically significant, p < .01, indicating that the conditional variance of the dependent variable was not

constant across levels of child's age. Therefore, the robust variance estimator should be used in the next stage to allow for heteroskedasticity.

```
# Regress outcome on treatment and matching variables using OLS
me m1 <- lm(pcss97 ~ kuse + male + black + age97 + pcged97 + mratio96 + pcg adc,
  data = me df
# Bresuch-Pagan test
bp <- function(var, df, md) {</pre>
 lmtest::bptest(md, as.formula(paste0("~", var)), data = df, studentize = T) %>%
   broom::tidy() %>%
   mutate(variable = var) %>%
    select(variable, statistic, p.value)
map_dfr(c("kuse", "male", "black", "age97", "pcged97", "mratio96", "pcg_adc"), bp,
 df = me_df, md = me_m1
) %>%
 kbl(
   booktabs = T, linesep = "", digits = 2,
   caption = "Results of Breusch-Pagan Tests for Heteroskedasticity"
  ) %>%
  kable styling(position = "center") %>%
  kable_styling(latex_options = c("striped", "hold_position"))
```

Table 1: Results of Breusch-Pagan Tests for Heteroskedasticity

variable	statistic	p.value
kuse	0.01	0.94
male	1.98	0.16
black	0.92	0.34
age97	16.29	0.00
pcged97	0.12	0.73
mratio 96	0.40	0.53
pcg_adc	0.19	0.66

5.5 Matching Estimators

Of the six matching variables, four are continuous and two are categorical, therefore bias-corrected matching estimator is necessary to correct for bias corresponding to the matching discrepancies between matched units and their matches on the four continuous covariates. Regression adjustment can be used with the BiasAdjust = T argument.

By default, the Matching::Match() function uses the matching variables to make bias adjustments. However, these covariates can be specified using the Z argument (example shown below).

If Var.calc = 0, then homoskedasticity is assumed. Use Var.calc = 4 to request the robust variance estimator using four matches.

The estimand argument is by default "ATT," but can be set to "ATE" or "ATC".

The sample argument is a logical flag for whether the population or sample variance is returned. An example may help illustrate the difference between PATE and SATE: "While the SATE is useful for judging how a job-training program has affected a particular group of participants, the PATE can be used to evaluate

whether another group of participants drawn from the same population is likely to benefit from the program."⁴ Results from the Matching::Match() function are identical to Stata's nnmatch program.

5.6 Define Outcome (Y), Treatment Index (Tr), and Variables to Match On (X)

```
me_Y <- me_df$pcss97
me_Tr <- me_df$kuse
me_X <- select(me_df, male, black, age97, pcged97, mratio96, pcg_adc)</pre>
```

5.7 Get Estimators Individually

Note that by default matching is done with replacement. However, this can be changed with the replace argument.

```
# Sample Average Treatment Effect (SATE)
summary(Match(Y = me_Y, Tr = me_Tr, X = me_X, M = 4, BiasAdjust = T, Var.calc = 4,
              estimand = "ATE", sample = T))
# Population Average Treatment Effect (PATE)
summary(Match(Y = me_Y, Tr = me_Tr, X = me_X, M = 4, BiasAdjust = T, Var.calc = 4,
              estimand = "ATE", sample = F))
# Sample average treatment effect for the treated (SATT)
summary(Match(Y = me_Y, Tr = me_Tr, X = me_X, M = 4, BiasAdjust = T, Var.calc = 4,
              estimand = "ATT", sample = T))
# Population average treatment effect for the treated (PATT)
summary(Match(Y = me_Y, Tr = me_Tr, X = me_X, M = 4, BiasAdjust = T, Var.calc = 4,
              estimand = "ATT", sample = F))
# Sample average treatment effect for the controls (SATC)
summary(Match(Y = me_Y, Tr = me_Tr, X = me_X, M = 4, BiasAdjust = T, Var.calc = 4,
              estimand = "ATC", sample = T))
# Population average treatment effect for the controls (PATC)
summary(Match(Y = me_Y, Tr = me_Tr, X = me_X, M = 4, BiasAdjust = T, Var.calc = 4,
             estimand = "ATC", sample = F))
```

5.8 Get All Estimators

```
# Function for extracting estimate, SE, t-stat, and p-value from Match()
get_match <- function(estimand, sample, Y = me_Y, Tr = me_Tr, X = me_X) {
    m <- Matching::Match(
        Y = Y, Tr = Tr, X = X, M = 4, BiasAdjust = T, Var.calc = 4,
        estimand = estimand, sample = sample
)
    return(list(
        est = m$est[, 1],
        se = m$se,
        t.stat = m$est[, 1] / m$se,
        p = (1 - pnorm(abs(m$est[, 1] / m$se))) * 2</pre>
```

 $^{^4}$ https://journals.sagepub.com/doi/pdf/10.1177/1536867X0400400307

```
))
}
# Estimate different matching estimators
tribble(
  ~estimator, ~estimand, ~sample,
  "SATE", "ATE", T,
  "PATE", "ATE", F,
  "SATT", "ATT", T,
  "PATT", "ATT", F,
  "SATC", "ATC", T,
  "PATC", "ATC", F
) %>%
  rowwise() %>%
  mutate(match = list(get_match(estimand, sample))) %>%
  unnest_wider(match) %>%
  select(-estimand, -sample) %>%
  kbl(
    booktabs = T, linesep = "",
    caption = "Bias-Corrected Matching Estimators with Robust Standard Errors"
  ) %>%
  kable_styling(position = "center") %>%
  kable_styling(latex_options = c("striped", "hold_position"))
```

Table 2: Bias-Corrected Matching Estimators with Robust Standard Errors

estimator	est	se	t.stat	p
SATE	-4.703773	1.769696	-2.657956	0.0078616
PATE	-4.703773	1.765187	-2.664746	0.0077047
SATT	-5.229651	1.781217	-2.935999	0.0033248
PATT	-5.229651	1.720590	-3.039451	0.0023701
SATC	-4.467254	2.133536	-2.093827	0.0362754
PATC	-4.467254	2.135647	-2.091757	0.0364602

When looking at the ATE, the results suggest that childhood poverty strongly affected children's academic achievement. Children who used AFDC had a passage comprehension score 4.7 units lower than that of children who never used AFDC. When considering only the treated population, this treatment effect was much larger.

Notice also that "inference on the PATE is made with respect to another sample drawn from the same population; inference on the SATE is made conditional on the sample at hand. (Although this does not affect the choice of estimator, for this reason, the standard error of the estimated PATE is generally larger than the standard error of the estimated SATE.)"

 $^{^5} https://journals.sagepub.com/doi/pdf/10.1177/1536867X0400400307$

5.9 Specify Variables in the Bias-Corrected Matching

5.10 Check Balance

Matched (Unweighted)

418.

188.

The Matching::Match() function works well in conjunction with the cobalt::bal.tab() function for checking covariate balance.

By default, the denominator for standardized mean differences uses a pooled estimate (square root of the average of the group variances) for ATE and the standard deviation of the treated group for ATT, and both standard deviations are computed using the sample before matching. This option can also be manually set with the s.d.denom option.

```
# Covariate Balance for SATE
me_SATE <- Match(</pre>
 Y = me_Y, Tr = me_Tr, X = me_X, M = 4, BiasAdjust = T, Var.calc = 4,
  estimand = "ATE", sample = T
)
bal.tab(
  me_SATE, kuse ~ male + black + age97 + pcged97 + mratio96 + pcg_adc,
  data = me_df,
  abs = T,
  un = T,
  thresholds = c(m = .1),
  s.d.denom = "pooled"
)
## Balance Measures
##
               Type Diff.Un Diff.Adj
                                             M. Threshold
## male
             Binary 0.0239
                               0.0050
                                          Balanced, <0.1
                                          Balanced, <0.1
## black
             Binary 0.4318
                               0.0245
## age97
            Contin.
                     0.0713
                               0.0010
                                          Balanced, <0.1
            Contin. 0.8084
                               0.3478 Not Balanced, >0.1
## pcged97
## mratio96 Contin.
                     1.0066
                               0.5831 Not Balanced, >0.1
## pcg_adc Contin.
                     0.6508
                               0.0450
                                          Balanced, <0.1
##
## Balance tally for mean differences
##
                       count
## Balanced, <0.1
## Not Balanced, >0.1
                           2
##
## Variable with the greatest mean difference
   Variable Diff.Adj
                              M.Threshold
##
               0.5831 Not Balanced, >0.1
##
    mratio96
##
## Sample sizes
##
                         Control Treated
## All
                          418.
                                  188.
## Matched (ESS)
                          318.46
                                   85.38
```

6 Practice Problems

6.1 Practice 1, Problem 1: Generalized Boosted Regression

6.1.1 Description of Dataset

This dataset is a subset of the experimental dataset used by LaLonde (1986).⁶ The LaLonde study is very famous among observational researchers. LaLonde is one of few pioneering researchers who used experimental data to cross-validate estimates of treatment effects generated by nonexperimental approaches. In this study, LaLonde's original dataset was created by a randomized experiment—a study examining trainee earnings of an employment program where participants were randomly assigned to treatment and control conditions. LaLonde's study compares the "true" treatment effect from this randomized experiment to a set of estimates generated by nonexperimental approaches. His study shows that "many of the nonexperimental procedures do not replicate the experimentally determined results, and suggests that researchers should be aware of the potential for specification errors in other nonexperimental evaluations" (LaLonde, 1986, p.604).

In this example, we are interested in the effect of participation in a job-training program on individuals' earnings in 1978. Thus, the dependent variable is re78 or earnings in 1978 (in thousands of 1978 dollars). The binary treatment variable is t or participation in a job-training program (1 = treated; 0 = control). The observable covariates are:

```
age: Age (in years)
educ: Years of education
black: African-American
hisp: Hispanic
married: Married
u74: Unemployed in 1974
u75: Unemployed in 1975
re74: Earnings in 1974 (in thousands of 1978 dollars)
re75: Earnings in 1975 (in thousands of 1978 dollars)
```

6.1.2 Load and Sort Data

```
set.seed(1000)
p11_df <- read_dta("data/ldw_exper.dta") %>%
  haven::zap_formats() %>%
  sjlabelled::remove_all_labels() %>%
  as_tibble() %>%
  add_column(runif = runif(nrow(.))) %>%
  arrange(runif) %>%
  select(-runif)
```

6.1.3 Estimate Propensity Scores

```
set.seed(1000)
p11_m1 <- gbm::gbm(
  formula = t ~ age + educ + black + hisp + married + re74 + re75 + u74 + u75,
  data = p11_df,
  distribution = "bernoulli",
  n.trees = 1000,
  train.fraction = 0.8,
  interaction.depth = 4,
  shrinkage = 0.0005</pre>
```

⁶LaLonde, R. J. (1986). Evaluating the Econometric Evaluations of Training Programs with Experimental Data. *The American Economic Review*, 76(4), 604–620. JSTOR.

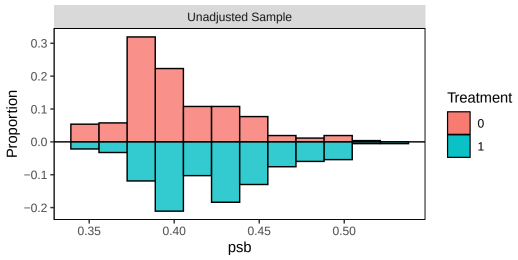
```
# Estimate Propensity Scores and Obtain Summary Statistics
p11_df$psb <- gbm::predict.gbm(p11_m1, data = p11_df, type = "response")
summary(p11_df$psb)

## Min. 1st Qu. Median Mean 3rd Qu. Max.
## 0.3394 0.3855 0.4006 0.4108 0.4346 0.5214</pre>
```

6.1.4 Histogram and Density Plots of Propensity Scores

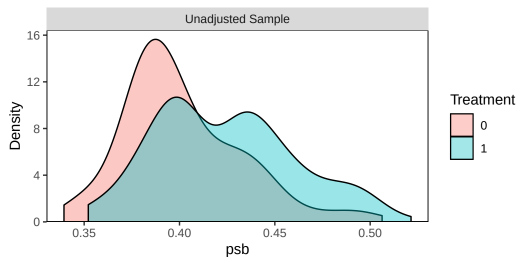
```
cobalt::bal.plot(
    t ~ psb,
    data = p11_df,
    var.name = "psb",
    type = "histogram",
    mirror = T
)
```

Distributional Balance for "psb"



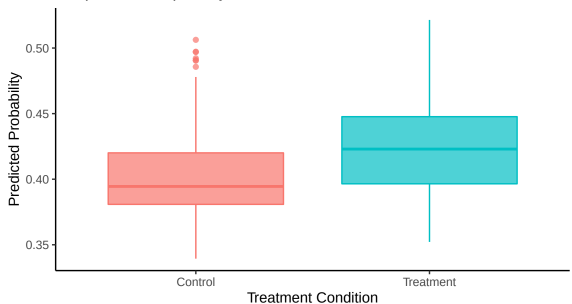
```
cobalt::bal.plot(
  t ~ psb,
   data = p11_df,
   var.name = "psb"
)
```

Distributional Balance for "psb"



6.1.5 Boxplot of Propensity Scores

Boxplots of Propensity Scores



6.2 Practice 1, Problem 2: Propensity Score Weighting

6.2.1 Import Stata-Generated Weights for Comparison

6.2.2 Estimate ATE and ATT Weights

6.2.3 Outcome Analysis with ATE and ATT Weights

```
# Define outcome formula
p12_f <- as.formula(re78 ~ t + age + educ + black + hisp +
 married + re74 + re75 + u74 + u75)
# Weighted OLS with R-Generated Propensity Scores
p12_m1 <- lm(p12_f, data = p12_df, weights = ate_w)
tidy(lmtest::coeftest(p12_m1, vcov. = vcovHC(p12_m1, "HC1"))) %>% filter(term == "t")
## # A tibble: 1 x 5
##
   term estimate std.error statistic p.value
             <dbl>
                       <dbl>
                               <dbl> <dbl>
                                  2.52 0.0122
              1.65
                       0.655
## 1 t.
p12_m2 <- lm(p12_f, data = p12_df, weights = att_w)
tidy(lmtest::coeftest(p12_m2, vcov. = vcovHC(p12_m2, "HC1"))) %>% filter(term == "t")
## # A tibble: 1 x 5
    term estimate std.error statistic p.value
##
     <chr>
             <dbl>
                       <dbl>
                                 <dbl> <dbl>
## 1 t
                       0.663
                                  2.60 0.00965
               1.72
# Weighted OLS with Stata-Generated Propensity Scores
p12_m1_stata <- lm(p12_f, data = p12_df, weights = stata_ate_w)
tidy(lmtest::coeftest(p12_m1_stata, vcov. = vcovHC(p12_m1_stata, "HC1"))) %>%
filter(term == "t")
## # A tibble: 1 x 5
   term estimate std.error statistic p.value
## <chr> <dbl>
                       <dbl>
                              <dbl> <dbl>
## 1 t
              1.63
                       0.656
                                  2.48 0.0135
p12_m2_stata <- lm(p12_f, data = p12_df, weights = stata_att_w)
tidy(lmtest::coeftest(p12_m2_stata, vcov. = vcovHC(p12_m2_stata, "HC1"))) %>%
filter(term == "t")
```

6.2.4 Check Balance

6.2.4.1 Hypothesis Tests We can use logistic regression and OLS regression to check the balance of categorical and continuous covariates, respectively:⁷

```
# Categorical Covariates
i1 <- glm(black ~ t, family = quasibinomial, data = p12_df, weights = stata_ate_w)
i2 <- glm(hisp ~ t, family = quasibinomial, data = p12_df, weights = stata_ate_w)
i3 <- glm(married ~ t, family = quasibinomial, data = p12_df, weights = stata_ate_w)
i4 <- glm(u74 ~ t, family = quasibinomial, data = p12_df, weights = stata_ate_w)
i5 <- glm(u75 ~ t, family = quasibinomial, data = p12_df, weights = stata_ate_w)
robustse(i1, coef = "odd.ratio")
robustse(i2, coef = "odd.ratio")
robustse(i3, coef = "odd.ratio")
robustse(i4, coef = "odd.ratio")
robustse(i5, coef = "odd.ratio")
# Continuous Covariates
i6 <- lm(age ~ t, data = p12_df, weights = stata_ate_w)</pre>
i7 <- lm(educ ~ t, data = p12_df, weights = stata_ate_w)
i8 <- lm(re74 ~ t, data = p12_df, weights = stata_ate_w)
i9 <- lm(re75 ~ t, data = p12_df, weights = stata_ate_w)
lmtest::coeftest(i6, vcov. = vcovHC(i6, "HC1"))
lmtest::coeftest(i7, vcov. = vcovHC(i7, "HC1"))
lmtest::coeftest(i8, vcov. = vcovHC(i8, "HC1"))
lmtest::coeftest(i9, vcov. = vcovHC(i9, "HC1"))
```

⁷See Appendix A for the custom function robustse() that is used to replicate the robust standard errors in Stata.

```
# Hypothesis testing for categorical covariates
p12_df %>%
  select(black, hisp, married, u74, u75, ate_w:stata_att_w, t) %>%
  pivot_longer(black:u75, names_to = "cat_covs", values_to = "cat_val") %%
  pivot_longer(ate_w:stata_att_w, names_to = "estimand", values_to = "weight") %>%
  group_by(estimand, cat_covs) %>%
 nest() %>%
  mutate(p.value = map(data, ~robustse(
   glm(.$cat_val ~ .$t, family = quasibinomial, weights = .$weight),
    coef = "odd.ratio")[2,4])) %>%
  unnest(p.value) %>%
  ungroup() %>%
  select(-data)
## # A tibble: 20 x 3
##
      cat_covs estimand
                           p.value
##
      <chr>
              <chr>
                             <dbl>
## 1 black ate_w
                             0.896
            att_w
## 2 black
                             0.977
## 3 black
            stata_ate_w 0.716
## 4 black stata_att_w 0.772
## 5 hisp
                             0.222
            ate_w
## 6 hisp
              att w
                             0.215
## 7 hisp
             stata_ate_w 0.127
## 8 hisp
               stata_att_w 0.128
## 9 married ate_w
                            0.452
## 10 married att_w
                             0.458
## 11 married stata_ate_w 0.402
## 12 married stata_att_w 0.383
## 13 u74
              \mathtt{ate}_{\mathtt{w}}
                             0.534
## 14 u74
              \mathtt{att}_{\mathtt{w}}
                             0.492
## 15 u74
              stata_ate_w 0.593
## 16 u74
               stata_att_w 0.492
## 17 u75
                             0.260
               ate_w
## 18 u75
               \mathtt{att}_{\mathtt{w}}
                             0.236
## 19 u75
               stata_ate_w
                             0.241
## 20 u75
              stata_att_w
                             0.181
# Hypothesis testing for continuous covariates
p12_df %>%
  select(age, educ, re74, re75, ate_w:stata_att_w, t) %>%
  pivot_longer(age:re75, names_to = "cont_covs", values_to = "cont_val") %>%
  pivot_longer(ate_w:stata_att_w, names_to = "estimand", values_to = "weight") %%
  group_by(estimand, cont_covs) %>%
 nest() %>%
  mutate(lm = map(data, ~lm(.$cont_val ~ .$t, weights = .$weight)),
         p.value = map(lm, ~lmtest::coeftest(., vcov. = vcovHC(., "HC1"))[2,4])) %>%
  unnest(p.value) %>%
  ungroup() %>%
  select(-data, -lm)
## # A tibble: 16 x 3
##
      cont covs estimand
                            p.value
##
      <chr>
            <chr>
                              <dbl>
```

0.520

1 age

ate_w

```
att_w
                                   0.512
## 2 age
                stata_ate_w 0.455
## 3 age
                stata_att_w 0.426
## 4 age
## 5 educ
                ate_w
                                 0.299
## 6 educ
                \mathtt{att}\_\mathtt{w}
                                   0.315
## 7 educ
                stata_ate_w 0.356
## 8 educ stata_att_w 0.302

## 9 re74 ate_w 0.863

## 10 re74 att_w 0.908

## 11 re74 stata_ate_w 0.722
## 12 re74
                stata_att_w 0.850
## 13 re75
                                   0.612
                \mathtt{ate}_{\mathtt{w}}
## 14 re75
                                   0.585
                \mathtt{att}_{\mathtt{w}}
## 15 re75
                stata_ate_w 0.666
             stata_att_w 0.553
## 16 re75
```

The survey package can also be used to incorporate weights:

```
library(survey)
i.svy <- survey::svydesign(~1, weights = p12_df$stata_ate_w, data = p12_df)
survey::svychisq(~black + t, design = i.svy)
survey::svychisq(~hisp + t, design = i.svy)
survey::svychisq(~married + t, design = i.svy)
survey::svychisq(~u74 + t, design = i.svy)
survey::svychisq(~u75 + t, design = i.svy)
survey::svyttest(age ~ t, design = i.svy)
survey::svyttest(educ ~ t, design = i.svy)
survey::svyttest(educ ~ t, design = i.svy)
survey::svyttest(re74 ~ t, design = i.svy)
survey::svyttest(re75 ~ t, design = i.svy)</pre>
```

6.2.4.2 Standardized Mean Differences Finally, the cobalt package can be used to check balance using standardized mean differences.⁸

```
cobalt::bal.tab(
  p12_df %% select(black, hisp, married, u74, u75, age, educ, re74, re75),
  treat = p12_df$t,
  weights = p12_df$ate_w,
 abs = T,
  threshold = .1,
  binary = "std",
  s.d.denom = "pooled",
 un = T
)
## Balance Measures
##
              Type Diff.Un Diff.Adj
                                           M.Threshold
                                        Balanced, <0.1
## black
            Binary 0.0440
                             0.0126
## hisp
            Binary 0.1749
                             0.1245 Not Balanced, >0.1
## married Binary 0.0939
                             0.0721
                                        Balanced, <0.1
## u74
            Binary 0.0944
                             0.0595
                                        Balanced, <0.1
```

0.1084 Not Balanced, >0.1

0.1009 Not Balanced, >0.1

Balanced, <0.1

Balanced, <0.1

Balanced, <0.1

```
## Balance tally for mean differences
## count
```

Contin. 0.0022

Contin. 0.0839

Binary 0.1772

0.1073

0.1412

0.0615

0.0162

0.0490

Contin.

Contin.

```
## Balanced, <0.1
                          6
## Not Balanced, >0.1
                          3
##
## Variable with the greatest mean difference
   Variable Diff.Adj
##
                             M.Threshold
               0.1245 Not Balanced, >0.1
##
        hisp
##
## Effective sample sizes
              Control Treated
## Unadjusted
               260.
                       185.
## Adjusted
               259.23 183.68
```

u75

age

educ ## re74

re75

##

 $^{^{86}}$ The "effective sample size" (ESS) is a measure of the sample size a non-weighted sample would have to have to achieve the same level of precision as the weighted sample (Ridgeway 2006)" (https://cran.r-project.org/web/packages/cobalt/vignettes/cobalt.html#effective-sample-size-for-weighting)

6.3 Practice 1: Alternative Solution with the WeightIt Package

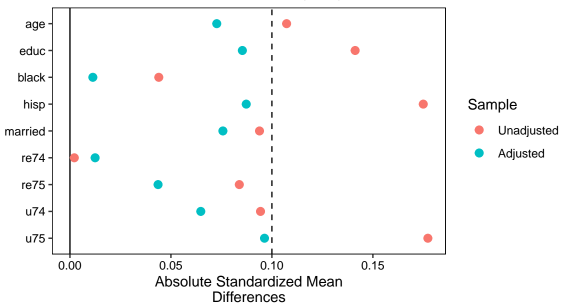
Use the GBM implementation in WeightIt::weightit() to estimate ATE and ATT:

```
set.seed(1000)
p1_ate <- WeightIt::weightit(</pre>
 formula = t ~ age + educ + black + hisp + married + re74 +
    re75 + u74 + u75,
  data = p11_df,
 method = "gbm",
  distribution = "bernoulli",
  stop.method = "es.mean",
 n.trees = 1000,
 nTrain = 0.8 * nrow(p11_df),
 interaction.depth = 4,
 shrinkage = 0.0005,
 estimand = "ATE"
set.seed(1000)
p1_att <- WeightIt::weightit(</pre>
 formula = t ~ age + educ + black + hisp + married + re74 +
   re75 + u74 + u75,
 data = p11_df,
 method = "gbm",
  distribution = "bernoulli",
  stop.method = "es.mean",
 n.trees = 1000,
 nTrain = 0.8 * nrow(p11_df),
 interaction.depth = 4,
 shrinkage = 0.0005,
  estimand = "ATT"
)
```

6.3.1 Check Balance

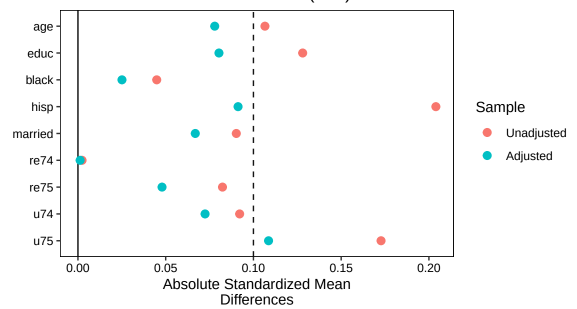
```
cobalt::love.plot(p1_ate, thresholds = .1, binary = "std", abs = T, drop.distance = T) +
labs(title = "Covariate Balance (ATE)")
```





cobalt::love.plot(p1_att, thresholds = .1, binary = "std", abs = T, drop.distance = T) +
labs(title = "Covariate Balance (ATT)")

Covariate Balance (ATT)



6.3.2 Outcome Analysis

For the outcome analysis, the ATE and ATT weights can be obtained with p1_ate\$weights (ATE) and p1_att\$weights (ATT):

```
p1_m1 <- lm(p12_f, data = p11_df, weights = p1_ate$weights)</pre>
tidy(lmtest::coeftest(p1_m1, vcov. = vcovHC(p1_m1, "HC1"))) %>% filter(term == "t")
## # A tibble: 1 x 5
##
    term estimate std.error statistic p.value
##
     <chr>
              <dbl>
                        <dbl>
                                  <dbl>
                                         <dbl>
                        0.646
                                   2.43 0.0155
## 1 t
               1.57
# ATT
p1_m2 <- lm(p12_f, data = p11_df, weights = p1_att$weightit)
tidy(lmtest::coeftest(p1_m2, vcov. = vcovHC(p1_m2, "HC1"))) %>% filter(term == "t")
## # A tibble: 1 x 5
##
   term estimate std.error statistic p.value
              <dbl>
                        <dbl>
                                  <dbl> <dbl>
##
     <chr>
                        0.662
                                   2.53 0.0119
## 1 t
               1.67
```

6.4 Practice 2: Matching Estimators

6.4.1 Load Data

```
p2_df <- haven::read_dta("data/prac2.dta") %>%
haven::zap_formats() %>%
sjlabelled::remove_all_labels() %>%
as_tibble()
```

6.4.2 Breusch-Pagan Test for Heteroskedasticity

The Breusch-Pagan test fits a linear regression model to the residuals of a linear regression model. A statistically significant result indicates that too much of the variance is explained by the additional explanatory variables.

The homoscedasticity assumption is not valid (e.g., p-value of the test for age97 is < .05), indicating that the conditional variance of the outcome variable was not constant across levels of child's age, therefore a robust estimation of variance is warranted.

```
# Fit a linear regression model, regressing the outcome on the covariates
p2_m1 <- lm(lwss97 ~ kuse + male + black + age97 + pcged97 + mratio96 + pcg_adc,
  data = p2_df
)
# Breusch-Pagan test
bp <- function(var) {</pre>
 lmtest::bptest(p2_m1, as.formula(paste0("~", var)), data = p2_df, studentize = T) %>%
   broom::tidy() %>%
   mutate(variable = var) %>%
    dplyr::select(variable, statistic, p.value)
map_dfr(c("kuse", "male", "black", "age97", "pcged97", "mratio96", "pcg_adc"), bp) %>%
 kbl(
    booktabs = T, linesep = "", digits = 2,
    caption = "Results of Breusch-Pagan Tests for Heteroskedasticity"
  ) %>%
  kable_styling(position = "center") %>%
  kable_styling(latex_options = c("striped", "hold_position"))
```

Table 3: Results of Breusch-Pagan Tests for Heteroskedasticity

variable	statistic	p.value
kuse	1.46	0.23
male	0.70	0.40
black	0.94	0.33
age 97	6.99	0.01
pcged97	3.62	0.06
mratio 96	5.60	0.02
pcg_adc	0.49	0.48

6.4.3 Define Outcome (Y), Treatment Index (Tr), and Variables to Match On (X)

```
p2_Y <- p2_df$lwss97
p2_Tr <- p2_df$kuse
```

```
p2_X <- select(p2_df, male, black, age97, pcged97, mratio96, pcg_adc)</pre>
```

6.4.4 Define Function for Matching

```
get_match <- function(estimand, sample, Y = p2_Y, Tr = p2_Tr, X = p2_X) {
    m <- Matching::Match(
        Y = Y, Tr = Tr, X = X, M = 4, BiasAdjust = T, Var.calc = 4,
        estimand = estimand, sample = sample
)
    return(list(
        est = m$est[, 1],
        se = m$se,
        t.stat = m$est[, 1] / m$se,
        p = (1 - pnorm(abs(m$est[, 1] / m$se))) * 2
))
}</pre>
```

6.4.5 Get All Estimators

```
tribble(
   ~estimator, ~estimand, ~sample,
   "SATE", "ATE", T,
   "PATE", "ATE", F,
   "SATT", "ATT", T,
   "PATT", "ATC", T,
   "PATC", "ATC", F
) %>%
   rowwise() %>%
   mutate(match = list(get_match(estimand, sample))) %>%
   tidyr::unnest_wider(match) %>%
   select(-estimand, -sample) %>%
   kbl(booktabs = T, linesep = "") %>%
   kable_styling(position = "center") %>%
   kable_styling(latex_options = c("striped", "hold_position"))
```

estimator	est	se	t.stat	p
SATE	-5.448863	1.646936	-3.3084850	0.0009380
PATE	-5.448863	1.652232	-3.2978811	0.0009742
SATT	-1.277287	1.683284	-0.7588067	0.4479682
PATT	-1.277287	1.695820	-0.7531973	0.4513314
SATC	-7.016781	1.965677	-3.5696503	0.0003575
PATC	-7.016781	1.969424	-3.5628594	0.0003668

7 Appendix A: Function to Replicate Stata's Robust Standard Errors

Function by Jorge Cimentada to replicate robust standard errors in Stata:⁹

```
robustse <- function(x, coef = c("logit", "odd.ratio", "probs")) {</pre>
  suppressMessages(suppressWarnings(library(lmtest)))
  suppressMessages(suppressWarnings(library(sandwich)))
  sandwich1 <- function(object, ...) {</pre>
    sandwich(object) *
      nobs(object) / (nobs(object) - 1)
  mod1 <- coeftest(x, vcov = sandwich1)</pre>
  if (coef == "logit") {
    return(mod1)
  } else if (coef == "odd.ratio") {
    mod1[, 1] <- exp(mod1[, 1])
    mod1[, 2] <- mod1[, 1] * mod1[, 2]
    return(mod1)
  } else {
    mod1[, 1] <- (mod1[, 1] / 4)
    mod1[, 2] <- mod1[, 2] / 4
    return(mod1)
  }
}
```

 $^{^9} https://cimentadaj.github.io/blog/2016-09-19-obtaining-robust-standard-errors-and-odds-ratios/obtaining-robust-standard-errors-and-odds-ratios/obtaining-robust-standard-errors-and-odds-ratios-for-logistic-regression-in-r/$

8 Appendix B: Rosenbaum's Sensitivity Analysis

The R package rbounds can be used to carry out Rosenbaum's sensitivity analysis. Please refer to Section 11.5.1 and 11.5.2 of the PSA-R code. Below is an example from the rbounds package documentation:¹⁰

```
library(rbounds)
# Data: Matched Data of Lead Blood Levels in Children
trt <- c(38, 23, 41, 18, 37, 36, 23, 62, 31, 34, 24, 14, 21, 17, 16, 20, 15,
         10, 45, 39, 22, 35, 49, 48, 44, 35, 43, 39, 34, 13, 73, 25, 27)
ctrl <- c(16, 18, 18, 24, 19, 11, 10, 15, 16, 18, 18, 13, 19, 10, 16, 16, 24, 13,
          9, 14, 21, 19, 7, 18, 19, 12, 11, 22, 25, 16, 13, 11, 13)
hlsens(trt, ctrl)
##
    Rosenbaum Sensitivity Test for Hodges-Lehmann Point Estimate
##
##
##
  Unconfounded estimate .... 15.5
##
##
    Gamma Lower bound Upper bound
##
        1
                 15.5
                              15.5
        2
                              19.6
##
                 10.5
        3
                  8.0
                              23.1
##
##
                  6.5
                              25.1
##
        5
                  5.0
                              26.6
##
        6
                  4.0
                              28.1
##
   Note: Gamma is Odds of Differential Assignment To
##
##
    Treatment Due to Unobserved Factors
##
```

 $^{^{10} \}rm https://cran.r-project.org/web/packages/rbounds/rbounds.pdf$