Propensity Score Analysis in R - Basics

Statistical Horizons

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Helpful Resources

- Propensity Score Analysis support site: https://ssw.unc.edu/psa/
- MatchIt package website: https://kosukeimai.github.io/MatchIt/
- Cobalt package website: https://ngreifer.github.io/cobalt/
- List of R software for propensity score analysis: https://www.elizabethstuart.org/psoftware/
- 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
- Greifer, N., & Stuart, E. A. (2023). Choosing the Causal Estimand for Propensity Score Analysis of Observational Studies (arXiv:2106.10577). arXiv. https://doi.org/10.48550/arXiv.2106.10577

Conceptual Review

Basic Concepts

Causal inference is inference about counterfactual outcomes or potential outcomes. An example of a counterfactual is: What would have happened to the treated subjects, had they not received treatment?

(Notation: Let w represent a dichotomous treatment variable, where 1 = treated and 0 = untreated, let Y represent a dichotomous outcome variable. Y_{1i} is the potential outcome for unit i with treatment, and Y_{0i} is the potential outcome for unit i without treatment.)

An individual causal effect is given by:

$$\tau_i = Y_{1i} - Y_{0i}$$

The **Neyman-Rubin counterfactual framework** states that individuals selected into treatment and non-treatment groups have potential outcomes in both states: the one in which they are observed and the one in which they are not observed, which can be expressed as:

$$Y_i = W_i Y_{1i} + (1 - W_i) Y_{0i}$$

This framework assumes the **Stable Unit Treatment Value Assumption (SUTVA)**, which assumes that (1) potential outcomes for an individual must not be affected by treatment for other individuals (e.g., spill-over effect); and (2) there are no different versions of the treatment condition (e.g., individuals receive different or variable levels of treatment).

Because "the fundamental problem of causal inference" is that individual causal effects are unobservable, we generally focus on **average causal effects**. One causal estimand is the **average treatment effect (ATE)**, given by:

$$\tau_{ATE} = \frac{1}{N} \sum_{i=1}^{N} \{Y_{1i} - Y_{0i}\} \text{ or } \tau_{ATE} = \mathbb{E}[Y_{1i} - Y_{0i}]$$

If we let N_1 equal the number of treated units, then the average treatment effect on the treated (ATT) is given by:

$$\tau_{ATT} = \frac{1}{N_1} \sum_{i=1}^{N} W_i \{ Y_{1i} - Y_{0i} \} \text{ or } \tau_{ATT} = \mathbb{E}[Y_{1i} - Y_{0i} | W_i = 1]$$

The **propensity score** is the probability of receiving the treatment given a vector of observed covariates, X_i :

$$\pi(X_i) \equiv \Pr(W_i = 1|X_i)$$

Greedy Matching

Greedy nearest neighbor matching forms matches by pairing a control participant with a treated participant, if the absolute difference of their propensity scores is the smallest among all possible pairs of propensity scores (i.e., they are similar on their covariates). If a caliper is used, then a match is made only if the absolute difference is less than the caliper.

Strengths

- Allows various kinds of post-matching analysis
- Particularly useful if the outcome variable is non-normal, non-continuous (e.g., categorical dependent variable, time-to-event data)
- Mahalanobis metric matching can be used to find well-matched pairs when the number of units is small

Limitations

- Optimization is only done locally
- "Bias due to incomplete matching" when treated subjects are excluded ¹
- Requires a sizeable common support region

Optimal Matching

Optimal matching minimizes a global distance measure to form matches. This circumvents the problem in greedy matching, where the order in which the treated are matched affects the quality of the matches. According to Gu and Rosenbaum (1993), "optimal matching picks about the same controls [as greedy matching] but does a better job of assigning them to treated units." Thus, optimal matching and greedy matching tend to produce similar results, but optimal matching may produce better matched pairs.²

In optimal pair matching, typically a treated participant is matched to a single control. With variable ratio or variable matching, each treated participant matches to a variable number of controls (because some individuals may have more matches than others).

Strengths

- Finds well-matched pairs
- Full matching uses the entire sample
- Robust against violations of overlap in common support region

Limitations

- Does not necessarily produce better balanced groups than greedy matching
- Variable matching may increase bias due to poor matches

Matching with or without Replacement

According to Stuart (2010), matching with replacement is "particularly helpful in settings where there are few control individuals comparable to the treated individuals (e.g., Dehejia and Wahba, 1999). Additionally, when matching with replacement the order in which the treated individuals are matched does not matter. However, inference becomes more complex when matching with replacement, because the matched controls are no longer independent—some are in the matched sample more than once and this needs to be accounted for in the outcome analysis, for example by using frequency weights. When matching with replacement it is also possible that the treatment effect estimate will be based on just a small number of controls; the number of times each control is matched should be monitored."

A simulation study comparing 12 propensity score algorithms found that "matching with replacement did not result in estimates with less bias compared with the best-performing methods based on caliper matching

¹Austin, P. C. (2014). A comparison of 12 algorithms for matching on the propensity score. *Statistics in Medicine*, 33(6), 1057–1069. https://doi.org/10.1002/sim.6004

²Stuart, 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

³Stuart, 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

without replacement. Furthermore, matching with replacement resulted in estimates that displayed greater variability and that had higher MSE compared with estimates obtained using caliper matching without replacement."⁴

According to Abadie & Imbens (2006), "Matching with replacement produces matches of higher quality than matching without replacement by increasing the set of possible matches. In addition, matching with replacement has the advantage that it allows us to consider estimators that match all units, treated as well as controls, so that the estimand is identical to the population average treatment effect." ⁵

Propensity Score Weighting

Unlike matching, propensity score weighting balances data by using propensity scores to create weights. It makes the estimate of the sample average treatment effect or its inference to the population average treatment effect a weighted average of the difference between observed and potential outcomes. It is similar to the weighted analysis that is conducted when analyzing complex survey designs. To estimate the ATE, the treatment weights are $1/\hat{e}(x)$, where $\hat{e}(x)$ is the propensity score, and the control weights are $1/(1-\hat{e}(x))$ —this is known as the inverse probability of treatment weights (IPTW) estimator. To estimate the ATT, the treatment weight is 1, and the control weights are $\hat{e}(x)/(1-\hat{e}(x))$.

Strengths

- Permits most types of multivariate outcome analyses
- Does not require an outcome variable that is continuous or normally distributed
- Retains most participants in the outcome analysis
- Flexible in the context of complex surveys⁶

Limitations

• Variance can be large if weights are extreme⁷

Matching Estimators

Matching estimators directly impute missing potential outcomes at the unit level using a vector norm. After imputing the missing data, matching estimators can be used to estimate various average treatment effects. Instead of using logistic regression to predict propensity scores, matching estimators use a vector norm to calculate distances on the observed covariates between a treated case and each of its potential control cases. These calculations use either the inverse of the sample variance matrix or the inverse of the sample variance-covariance matrix, the matching estimator calculates Mahalanobis metric distances.

Strengths

- Calculate a variety of average treatment effects
- Estimate effects for both the sample and the population

 $^{^4}$ Austin, P. C. (2014). A comparison of 12 algorithms for matching on the propensity score. Statistics in Medicine, 33(6), 1057–1069. https://doi.org/10.1002/sim.6004

⁵Abadie, A., & Imbens, G. W. (2006). Large Sample Properties of Matching Estimators for Average Treatment Effects. *Econometrica*, 74(1), 235–267. https://doi.org/10.1111/j.1468-0262.2006.00655.x

⁶DuGoff, E. H., Schuler, M., & Stuart, E. A. (2014). Generalizing Observational Study Results: Applying Propensity Score Methods to Complex Surveys. *Health Services Research*, 49(1), 284–303. https://doi.org/10.1111/1475-6773.12090

⁷Austin, P. C., & Stuart, E. A. (2015). Moving towards best practice when using inverse probability of treatment weighting (IPTW) using the propensity score to estimate causal treatment effects in observational studies. *Statistics in Medicine*, 34(28), 3661–3679. https://doi.org/10.1002/sim.6607

Liim tations

- Require a sizeable common support region
 More sensitive to the violation of the strongly ignorable assumption

3.5 Computer Lab: Running Greedy Matching and GBR with R

Greedy Nearest Neighbor Matching

Greedy nearest neighbor matching forms matches by pairing a control participant with a treated participant, if the absolute difference of their propensity scores is the smallest among all possible pairs of propensity scores (i.e., they are similar on their covariates). If a caliper is used, then a match is made only if the absolute difference is less than the caliper.

Load Packages

The haven and sjlabelled packages are used to load and clean Stata data files (.dta); the MatchIt package contains functions for greedy matching⁸; the cobalt package contains functions for balance checking; and the tidyverse package is loaded for its data manipulation functions.

```
if (!require("pacman")) install.packages("pacman")
pacman::p_load(
  haven, sjlabelled, cobalt, MatchIt, tidyverse, kableExtra, survival, psych, here
)
```

Description of Data

This data is a sample of 2,758 children from the National Survey of Child and Adolescent Well-Being (NSCAW), a nationally representative, longitudinal survey of children and families who have been the subjects of investigation by Child Protective Services. Two waves of NSCAW data were used: baseline information between October 1999 and December 2000 and the 18-months follow-up. The sample was limited to children who lived at home (e.g., were not in foster care) and whose primary caregivers were female (because the vast majority of primary caregivers in NSCAW were females). 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. In Section 5.8.1 of the PSA-R code, 12 matching schemes are shown.

Load and Clean Data

```
# Load Data
gm_df0 <- haven::read_dta(here("data", "chpt5_1_original.dta"))
# Inspect Data
str(gm_df0)
# Remove Stata Labels and Formats
gm_df0 <- gm_df0 %>%
    haven::zap_formats() %>%
    sjlabelled::remove_all_labels() %>%
    as_tibble()
# Inspect Data
str(gm_df0)
```

 $^{^{8}}$ https://kosukeimai.github.io/MatchIt/articles/matching-methods.html

⁹https://ssw.unc.edu/psa/

```
psych::describe(gm_df0)
nrow(gm_df0)
table(gm_df0$aodserv)
```

Sort Data

When non-treated cases have the same propensity score values, their matches will depend on the order of the data. Therefore, it's important to order the observations randomly.

```
set.seed(1000)
gm_df <- gm_df0 %>%
  add_column(runif = runif(nrow(.)), .before = "PSH17A") %T>% print() %>%
  arrange(runif) %T>% print() %>%
  select(-runif)
```

Check Balance Before Matching

Because all of the variables that will be used for predicting the propensity scores of service receipt are categorical, we can use chisq.test to check their balance before matching. Before we do that, we convert these categorical variables from numeric variables to factor variables using as.factor(). This will help functions such as bal.tab() to automatically calculate appropriate balance statistics.

```
# Change categorical independent variables to factor variables, except the id and
# and aodserv treatment variables
gm_df <- gm_df %>%
  mutate(across(c(-id, -aodserv), as.factor))
# Marital Status (large sample size, therefore continuity correction not needed)
chisq.test(gm_df$aodserv, gm_df$married, correct = F)
##
## Pearson's Chi-squared test
##
## data: gm_df$aodserv and gm_df$married
## X-squared = 2.9705, df = 1, p-value = 0.0848
# All Variables
gm df %>%
  select(
   married, educ, pov, employ, open, race, chdage, cgage, CRA47A, mental,
   arrest, PSH17A, maltx, ra, cidi, cgneed, cwwrep, aodserv
  pivot_longer(-aodserv, names_to = "variable") %>%
  group_by(variable) %>%
  nest() %>%
  mutate(bivariate.test = map(data, ~chisq.test(.$aodserv, .$value, correct = F))) %>%
  mutate(statistic = map(bivariate.test, ~ round(.$statistic, 3))) %>%
  mutate(p.value = map(bivariate.test, ~ round(.$p.value, 3))) %>%
  unnest(cols = c(statistic, p.value)) %>%
  select(variable, statistic, p.value)
```

```
## # A tibble: 17 x 3
## # Groups:
               variable [17]
      variable statistic p.value
##
##
                    <dbl>
      <chr>>
                            <dbl>
##
    1 married
                    2.97
                            0.085
                    10.5
                            0.005
##
    2 educ
                            0.023
##
    3 pov
                    11.3
##
    4 employ
                    23.1
                            0
##
   5 open
                    58.4
                            0
                            0.009
##
    6 race
                    11.5
   7 chdage
                    55.4
                    3.56
##
    8 cgage
                            0.313
##
  9 CRA47A
                    17.5
                            0
## 10 mental
                   92.5
                            0
## 11 arrest
                   127.
                            0
## 12 PSH17A
                   179.
                            0
## 13 maltx
                   49.7
                            0
## 14 ra
                  585.
                            0
## 15 cidi
                   157.
                            0
## 16 cgneed
                   139.
                            0
## 17 cwwrep
                  1240.
                            0
# Fisher's Exact Test for cgage
(c1 <- chisq.test(gm_df$aodserv, gm_df$cgage, correct = F))</pre>
##
##
   Pearson's Chi-squared test
##
## data: gm_df$aodserv and gm_df$cgage
## X-squared = 3.5619, df = 3, p-value = 0.3128
c1$expected
                gm_df$cgage
##
                                             2
  gm_df$aodserv
                         0
                                   1
                                                        3
##
               0 38.35388 1698.2741 582.4438 140.92821
##
               1 4.64612 205.7259 70.5562
                                                17.07179
fisher.test(gm_df$aodserv, gm_df$cgage)
##
##
    Fisher's Exact Test for Count Data
##
## data: gm_df$aodserv and gm_df$cgage
## p-value = 0.288
## alternative hypothesis: two.sided
```

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

The standardized mean difference (SMD) (also referred to as the "normalized difference") is a commonly used balance measure.¹⁰ It is calculated as the difference in means of a covariate across the treatment groups,

¹⁰The disadvantage of hypothesis tests is "they are influenced by sample size, which fluctuates during adjustment, and the theory behind them is inappropriate because balance is a quality solely of the sample in question, not in relation to a population" (https://cran.r-project.org/web/packages/cobalt/vignettes/cobalt.html).

divided by the standard deviation in the treated group (ATT), the control group (ATC), or the pooled standard deviation (ATE). See ?cobalt::col_w_smd for additional options. Stuart et al. (2013) recommend 0.1 or 0.25 as reasonable cut-offs for acceptable standardized biases.¹¹

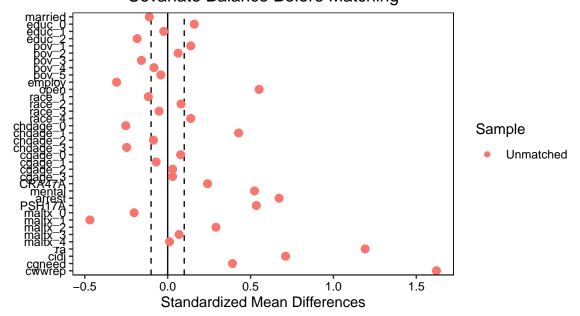
```
# Balance table
cobalt::bal.tab(
    select(
        gm_df, married, educ, pov, employ, open, race, chdage, cgage, CRA47A,
        mental, arrest, PSH17A, maltx, ra, cidi, cgneed, cwwrep
),
    treat = gm_df$aodserv,
    s.d.denom = "treated", # ATT
    threshold = .1,
    stats = "mean.diff",
 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

```
## Balance Measures
##
              Type Diff.Un
                               M.Threshold.Un
           Binary -0.1108 Not Balanced, >0.1
## married
## educ_0
            Binary 0.1605 Not Balanced, >0.1
            Binary -0.0236
## educ 1
                               Balanced, <0.1
            Binary -0.1840 Not Balanced, >0.1
## educ_2
## pov_1
           Binary 0.1395 Not Balanced, >0.1
## pov 2
           Binary 0.0628
                               Balanced, <0.1
## pov_3
           Binary -0.1584 Not Balanced, >0.1
           Binary -0.0829
                               Balanced, <0.1
## pov_4
## pov_5
           Binary -0.0410
                               Balanced, <0.1
## employ
           Binary -0.3082 Not Balanced, >0.1
## open
           Binary 0.5516 Not Balanced, >0.1
## race 1
           Binary -0.1167 Not Balanced, >0.1
## race 2
           Binary 0.0807
                               Balanced, <0.1
## race 3
           Binary -0.0519
                               Balanced, <0.1
           Binary 0.1392 Not Balanced, >0.1
## race_4
## chdage_0 Binary -0.2527 Not Balanced, >0.1
## chdage_1 Binary 0.4279 Not Balanced, >0.1
## chdage 2 Binary -0.0856
                               Balanced, <0.1
## chdage_3 Binary -0.2473 Not Balanced, >0.1
## cgage_0 Binary 0.0781
                               Balanced, <0.1
## cgage_1 Binary -0.0694
                               Balanced, <0.1
## cgage_2 Binary 0.0300
                               Balanced, <0.1
## cgage 3 Binary 0.0297
                               Balanced, <0.1
## CRA47A
           Binary 0.2406 Not Balanced, >0.1
## mental
           Binary 0.5238 Not Balanced, >0.1
## arrest
           Binary 0.6718 Not Balanced, >0.1
## PSH17A
           Binary 0.5347 Not Balanced, >0.1
## maltx_0 Binary -0.2025 Not Balanced, >0.1
## maltx 1 Binary -0.4690 Not Balanced, >0.1
## maltx_2 Binary 0.2909 Not Balanced, >0.1
## maltx_3 Binary 0.0690
                               Balanced, <0.1
```

 $^{^{11}\}mathrm{Stuart},$ 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. 2013.01.013

```
## maltx_4 Binary 0.0110
                               Balanced, <0.1
## ra
            Binary 1.1915 Not Balanced, >0.1
## cidi
            Binary 0.7110 Not Balanced, >0.1
## cgneed
            Binary 0.3907 Not Balanced, >0.1
## cwwrep
            Binary 1.6213 Not Balanced, >0.1
##
## Balance tally for mean differences
##
                      count
## Balanced, <0.1
                         13
                         23
## Not Balanced, >0.1
## Variable with the greatest mean difference
##
   Variable Diff.Un
                        M.Threshold.Un
##
      cwwrep 1.6213 Not Balanced, >0.1
##
## Sample sizes
##
       Control Treated
## All
          2460
                   298
# Love plot
cobalt::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,
  sample.names = c("Unmatched")
) +
  labs(title = "Covariate Balance Before Matching")
```

Covariate Balance Before Matching



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. Non-replacement means that once a treated case is matched to a non-treated case, both cases are removed from the pool. Matching with replacement allows each control unit to be matched with any number of treated units.

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). Austin (2011) recommends a caliper size of 0.2 of the standard deviation.¹²

The order of the matching can be specified using the m.order argument. If this argument is set to largest, then matching begins with the treated subject with the highest propensity score; if set to smallest, then matching takes places in ascending order of the distance measures; and if random, matching takes place in a random order. "The default is to go in descending order from the highest propensity score; doing so allows the units that would have the hardest time finding close matches to be matched first." Note that when non-treated cases have the same propensity score values, their matches will depend on the order of the data. Thus, it is still important to randomly shuffle your data prior to matching.

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. Matching on the logit also improves balance, compared to matching on the raw propensity score. To calculate this automatically, set distance="glm" and link="logit".

The choice of explanatory variables (i.e., conditioning variables) in the model predicting propensity scores of service receipt serves a paramount role in the propensity score analysis. We chose these variables based on a review of substance abuse literature to determine what characteristics were associated with treatment receipt:

```
# Logistic regression specification
(gm_f <- cobalt::f.build("aodserv", select(gm_df, PSH17A:other, -aodserv)))

## 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: 0x0000020386bb1578>
```

We then calculate the logit of the predicted probability as the propensity score:

```
gm_psm <- glm(gm_f, data = gm_df, family = binomial)
gm_ps <- predict(gm_psm, newdata = gm_df, type = "response")
gm_ps_logit <- log((1 - gm_ps) / gm_ps)</pre>
```

And then perform greedy nearest neighbor matching without replacement

¹²https://pubmed.ncbi.nlm.nih.gov/20925139/

 $^{^{13} \}rm https://kosukeimai.github.io/MatchIt/articles/matching-methods.html$

```
(gm_out <- MatchIt::matchit(
  gm_f,
  data = gm_df,
  distance = gm_ps_logit,
  m.order = "largest", # descending order
  caliper = .25
))

## A 'matchit' object
## - method: 1:1 nearest neighbor matching without replacement
## - distance: User-defined [caliper]
## - caliper: <distance> (0.311)
```

- covariates: PSH17A, CRA47A, married, high, bahigh, poverty2, poverty3, poverty4, poverty5, employ

Notice that a limitation of using calipers is that only 283 out of 298 of the original treated cases were matched. This matching scheme reduces the sample size from 2758 to 566—283 cases in the control group and 283 cases in the treated group.

- number of obs.: 2758 (original), 566 (matched)

```
set.seed(1000)
(gm_out_20 <- MatchIt::matchit(
   gm_f,
   data = gm_df,
   distance = gm_ps_logit,
   m.order = "largest", # descending order
   caliper = .20
))</pre>
```

```
## A 'matchit' object
## - method: 1:1 nearest neighbor matching without replacement
## - distance: User-defined [caliper]
## - caliper: <distance> (0.249)
## - number of obs.: 2758 (original), 566 (matched)
## - target estimand: ATT
## - covariates: PSH17A, CRA47A, married, high, bahigh, poverty2, poverty3, poverty4, poverty5, employ
```

The matchit object will return a match.matrix, which contains the treated units as the rownames and the values in each row the names or indices of the control units matched to the treated units:

```
head(gm_out$match.matrix)
```

```
## [,1]
## 2 "420"
## 3 "1852"
## 10 "2130"
## 14 NA
## 41 "782"
## 49 "1162"
```

set.seed(1000)

- target estimand: ATT

Remember to use weights when estimating the treatment effect (but this is not necessary when 1:1 matching without replacement was performed):

```
head(gm_out$weights)
```

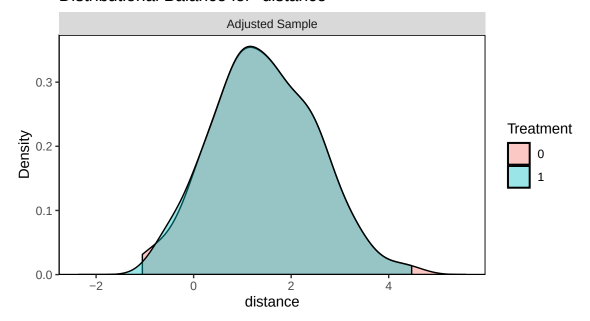
```
## 1 2 3 4 5 6
## 0 1 1 0 0 0
```

Check Common Support 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. The discard argument in matchit() can be used to discard units outside a region of common support.

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

```
## Ignoring unknown labels:
## * colour : "Treatment"
```

Distributional Balance for "distance"



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

```
# Extract matched data
gm_out_data <- MatchIt::match.data(gm_out)

# Assess balance on "ra"
chisq.test(gm_out_data$ra, gm_out_data$aodserv)</pre>
```

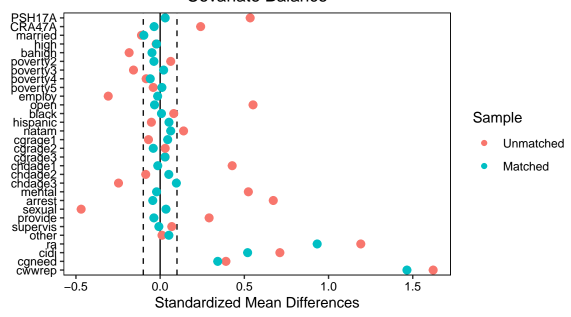
```
##
## Pearson's Chi-squared test with Yates' continuity correction
```

```
##
## data: gm_out_data$ra and gm_out_data$aodserv
## X-squared = 108.05, df = 1, p-value < 0.000000000000000022</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 the data argument.

```
cobalt::love.plot(gm_out,
  binary = "std", threshold = c(m = .1), drop.distance = T,
  addl = c("ra", "cidi", "cgneed", "cwwrep"), data = gm_df,
  sample.names = c("Unmatched", "Matched")
)
```

Covariate Balance



```
cobalt::bal.tab(gm_out,
  binary = "std", threshold = c(m = .1), un = T,
  addl = c("ra", "cidi", "cgneed", "cwwrep"), data = gm_df
)
```

```
## Balance Measures
##
               Type Diff.Un Diff.Adj
                                            M.Threshold
## distance Distance -1.3021
                              0.0068
                                         Balanced, <0.1
## PSH17A
          Binary 0.5347
                              0.0301
                                         Balanced, <0.1
             Binary 0.2406 -0.0359
## CRA47A
                                         Balanced, <0.1
## married
                                         Balanced, <0.1
             Binary -0.1108
                            -0.0969
             Binary -0.0236
                            -0.0214
## high
                                         Balanced, <0.1
## bahigh
             Binary -0.1840 -0.0485
                                         Balanced, <0.1
## poverty2
             Binary 0.0628 -0.0370
                                         Balanced, <0.1
## poverty3
             Binary -0.1584
                             0.0207
                                         Balanced, <0.1
## poverty4
             Binary -0.0829
                            -0.0587
                                         Balanced, <0.1
## poverty5
             Binary -0.0410
                             0.0105
                                         Balanced, <0.1
```

```
Balanced, <0.1
## employ
              Binary -0.3082 -0.0148
              Binary 0.5516
                             -0.0335
                                          Balanced, <0.1
## open
## black
              Binary 0.0807
                               0.0078
                                          Balanced, <0.1
              Binary -0.0519
                                          Balanced, <0.1
## hispanic
                               0.0524
## natam
              Binary 0.1392
                               0.0626
                                          Balanced, <0.1
## cgrage1
              Binary -0.0694
                               0.0448
                                          Balanced, <0.1
## cgrage2
              Binary 0.0300
                              -0.0409
                                          Balanced, <0.1
              Binary 0.0297
                                          Balanced, <0.1
## cgrage3
                               0.0289
              Binary 0.4279
## chdage1
                              -0.0142
                                          Balanced, <0.1
## chdage2
              Binary -0.0856
                              0.0513
                                          Balanced, <0.1
## chdage3
              Binary -0.2473
                               0.0963
                                          Balanced, <0.1
              Binary 0.5238
                                          Balanced, <0.1
## mental
                              -0.0212
## arrest
              Binary 0.6718
                              -0.0438
                                          Balanced, <0.1
                                          Balanced, <0.1
## sexual
              Binary -0.4690
                               0.0346
## provide
              Binary 0.2909
                              -0.0373
                                          Balanced, <0.1
## supervis
              Binary 0.0690
                              -0.0077
                                          Balanced, <0.1
## other
                               0.0518
              Binary 0.0110
                                          Balanced, <0.1
## ra
              Binary 1.1915
                               0.9327 Not Balanced, >0.1
## cidi
              Binary 0.7110
                               0.5195 Not Balanced, >0.1
## cgneed
              Binary 0.3907
                               0.3419 Not Balanced, >0.1
## cwwrep
              Binary 1.6213
                               1.4656 Not Balanced, >0.1
## 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
              1.4656 Not Balanced, >0.1
##
      cwwrep
##
## Sample sizes
##
             Control Treated
## All
                2460
                         298
## Matched
                 283
                         283
```

Unmatched

2177

15

Greedy Nearest Neighbor Mahalanobis Distance Matching Without Replacement

Here we perform Mahalanobis distance matching without replacement and without including estimated propensity scores by setting the distance argument to "mahalanobis." If propensity scores need to be estimated, the argument mahvars can be used to specify the variables used to create the Mahalanobis distance, while the distance argument should be either a vector of the propensity scores or whatever method is desired for estimating the propensity scores.

```
gm_f,
data = gm_df,
method = "nearest",
distance = "mahalanobis",
replace = F
))

## 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
cobalt::love.plot(gm_out2,
```

Check Balance

)

set.seed(1000)

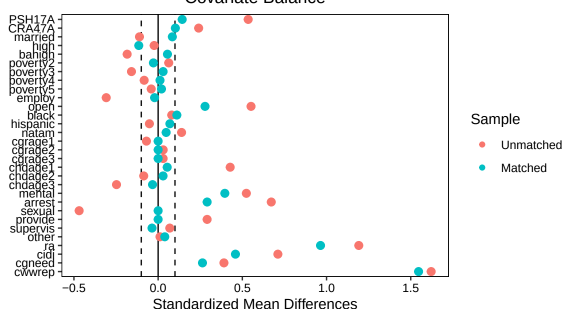
(gm out2 <- MatchIt::matchit(</pre>

binary = "std", threshold = c(m = .1),

sample.names = c("Unmatched", "Matched")

addl = c("ra", "cidi", "cgneed", "cwwrep"), data = gm_df,

Covariate Balance



```
binary = "std", threshold = c(m = .1),
  addl = c("ra", "cidi", "cgneed", "cwwrep"), data = gm_df
## Balance Measures
              Type Diff.Adj
                                    M.Threshold
## PSH17A
            Binary
                     0.1429 Not Balanced, >0.1
## CRA47A
            Binary
                     0.1022 Not Balanced, >0.1
## married Binary
                     0.0843
                                 Balanced, <0.1
            Binary -0.1150 Not Balanced, >0.1
## high
## bahigh
            Binary
                     0.0552
                                 Balanced, <0.1
## poverty2 Binary
                   -0.0281
                                 Balanced, <0.1
## poverty3 Binary
                     0.0295
                                 Balanced, <0.1
                                 Balanced, <0.1
## poverty4 Binary
                     0.0112
## poverty5 Binary
                     0.0199
                                 Balanced, <0.1
## employ
            Binary -0.0211
                                 Balanced, <0.1
## open
            Binary
                     0.2784 Not Balanced, >0.1
                     0.1111 Not Balanced, >0.1
## black
            Binary
                     0.0696
                                 Balanced, <0.1
## hispanic Binary
## natam
            Binary
                     0.0476
                                 Balanced, <0.1
## cgrage1 Binary
                     0.0000
                                Balanced, <0.1
## cgrage2 Binary
                     0.0000
                                Balanced, <0.1
                                Balanced, <0.1
## cgrage3 Binary
                     0.0000
## chdage1
            Binary
                     0.0538
                                 Balanced, <0.1
                                 Balanced, <0.1
## chdage2
           Binary
                     0.0292
## chdage3
           Binary -0.0333
                                 Balanced, <0.1
## mental
            Binary
                     0.3960 Not Balanced, >0.1
## arrest
                     0.2910 Not Balanced, >0.1
            Binary
                                 Balanced, <0.1
## sexual
            Binary
                     0.0000
## provide
            Binary
                     0.0000
                                 Balanced, <0.1
## supervis Binary -0.0364
                                 Balanced, <0.1
## other
            Binary
                     0.0394
                                 Balanced, <0.1
                     0.9643 Not Balanced, >0.1
## ra
            Binary
## cidi
                     0.4591 Not Balanced, >0.1
            Binary
## cgneed
            Binary
                     0.2633 Not Balanced, >0.1
## cwwrep
            Binary
                     1.5465 Not Balanced, >0.1
##
## Balance tally for mean differences
##
                      count
## Balanced, <0.1
                         19
## Not Balanced, >0.1
##
## Variable with the greatest mean difference
##
   Variable Diff.Adj
                             M.Threshold
##
      cwwrep
               1.5465 Not Balanced, >0.1
##
## Sample sizes
##
             Control Treated
## All
                2460
                         298
                         298
## Matched
                 298
## Unmatched
                2162
                           0
```

cobalt::bal.tab(gm_out2,

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." 14

Extract Matched Data After matching, the treatment effect can be estimated using the matched sample, which can be extracted using the MatchIt::match.data() function.

MatchIt::match.data(gm_out2)

 $^{^{14}\}mathrm{Stuart},$ 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

Propensity Score Estimation Using Generalized Boosted Regression

Generalized boosted regression is an iterative method for creating propensity scores. It uses an automated, data-adaptive algorithm that fits several models by way of a regression tree, and then merges the predictions produced by each model. The regression tree partitions the sample into small groups based on predictor variables.

GBR is a sum of regression trees. These trees are computationally fast to fit, and they are invariant to one-to-one transformations of the independent variables. Its advantage over logistic regression is that it doesn't require knowing the functional form of predictor variables.

Load Package

Generalized boosted regression (GBR) requires the gbm package.

```
pacman::p_load(gbm)
```

Load Data and Sort

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. To create reproducible results, we need to use the set.seed() function.

```
set.seed(1000)
gbr_df <- read_dta(here("data", "g3aca1.dta")) %>%
  haven::zap_formats() %>%
  sjlabelled::remove_all_labels() %>%
  as_tibble() %>%
  select(
    intbl, ageyc, fmale, blck, whit, hisp, pcedu, ipovl, pcemft, fthr, dicsagg2, dicsint2, dccereg2, dccscom2, dccpros2, draggr2
) %>%
  drop_na() %>%
  add_column(runif = runif(nrow(.))) %>%
  arrange(runif) %>%
  select(-runif)
```

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. For example, to reduce prediction error, the train.fraction argument can be set to use a subsample of the observations for the estimation process. Here we also use interaction.depth = 4 to specify a maximum of four splits for each sample tree used in the model, which allows all four-way interactions between all covariates to be considered for optimizing the likelihood function at each iteration. The shrinkage argument is also known as the learning rate or step-size reduction; we use a value of .0005 to ensure a smooth fit.

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 ouput 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)))

## intbl ~ ageyc + fmale + blck + whit + hisp + pcedu + ipovl +

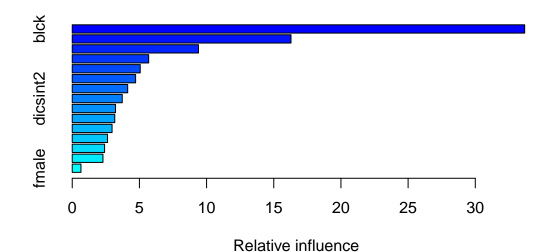
## pcemft + fthr + dicsagg2 + dicsint2 + dccereg2 + dccscom2 +

## dccpros2 + draggr2

## <environment: 0x000002037ea9b2a0>

set.seed(1000)
gbr_m1 <- gbm::gbm(
    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</pre>
```

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 16.2834103
## ageyc
            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
## dccereg2 dccereg2 2.9647178
## dccpros2 dccpros2 2.6269129
## hisp
               hisp 2.4079157
## fthr
               fthr
                     2.2874508
## fmale
              fmale 0.6494603
```

summary(gbr_m1)

Estimate Propensity Scores

After fitting the model, estimate propensity scores using the ${\tt predict.gbm}()$ function.

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

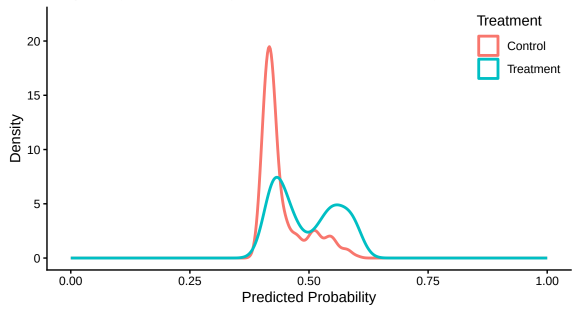
[1] 0.4441531 0.4405950 0.5288983 0.5958331 0.3985049 0.5562076

Plot Propensity Score Distributions

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

```
gbr_df %>%
  mutate(psb = psb, intbl = factor(intbl, labels = c("Control", "Treatment"))) %>%
  ggplot(aes(x = psb, color = intbl)) +
  theme_classic() +
  geom_density(linewidth = 1) +
  xlim(0, 1) +
  ylim(0, 22) +
  labs(
    x = "Predicted Probability", y = "Density",
    title = "Propensity Scores Using Generalized Boosted Regression",
    color = "Treatment"
  ) +
  theme(legend.position = c(0.9, 0.85))
```

Propensity Scores Using Generalized Boosted Regression



Summary Statistics of Propensity Scores

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

GBR Using the WeightIt Package

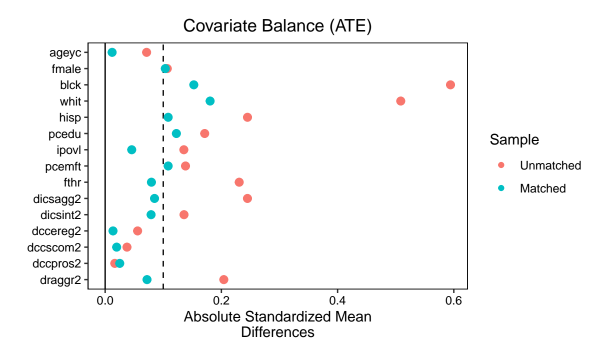
As an alternative to the gbm package, the WeightIt package can also fit GBR models. Note that the default maximum number of trees used (the n.trees argument) is 10000 for binary treatments.

```
set.seed(1000)
(gbr_m2 <- WeightIt::weightit(</pre>
  formula = gbr_f,
  data = gbr_df,
 method = "gbm",
  estimand = "ATE",
  distribution = "bernoulli", # binary treatment
  stop.method = "es.mean",
 n.trees = 10000,
 nTrain = 0.8 * nrow(gbr_df),
  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
# Propensity Scores
head(gbr_m2$ps)
```

[1] 0.3781517 0.6075811 0.6596871 0.7766516 0.2138153 0.7927817

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,
  sample.names = c("Unmatched", "Matched")
) +
  labs(title = "Covariate Balance (ATE)")
```



Other Packages

GBR is also implemented in the MatchIt::matchit() and twang::ps() functions. While the gbm package is used in both, each function uses a different set of default arguments. See their respective help files for more details.

4.7 Computer Lab: Imbalance Check and Outcome Analysis with R after optmatch

Optimal Matching

Load Packages

```
pacman::p_load(optmatch, knitr, broom, sandwich, rlang, marginaleffects)
select <- dplyr::select</pre>
```

Load Data

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

The dependent variable is lwss97, the age-normed "letter-word" identification score of the Woodcock-Johnson Revised Tests of Achievement. A high score on this measure indicates high achievement. The treatment variable is kuse or children who ever used Aid to Families With Dependent Children (AFDC). The covariates are:

- mratio96: Ratio of Family Income to Poverty Line in 1996
- pcged97: Caregiver's Education in 1997 (Years of Schooling)
- pcg_adc: Caregiver's History of Using Welfare (Number of Years; range: 0-7)
- black: Child's Race: African American (1 = African American; 0 = Other)
- age97: Child's Age in 1997
- male: Child's Gender: Male (1 = Male; 0 = Female)

```
d <- haven::read_dta("data/opt/chpt5_2_original.dta")
df <- haven::read_dta("data/opt/chpt5_2.dta")
cds <- haven::read_dta("data/opt/chpt5_2ps.dta")</pre>
```

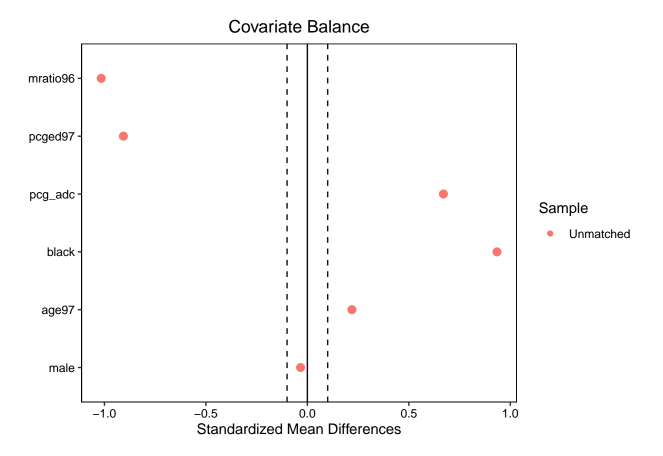
Bivariate Comparisons

With the exception of child's gender, the difference on each covariate between treated and control groups is statistically significant. Without controlling for these covariates, the study's estimate of treatment effect would be biased.

```
# Wilcoxon Rank-Sum (Mann-Whitney) test and t-test
d %>%
 select(mratio96, pcged97, pcg_adc, black, age97, male, kuse) %>%
 pivot_longer(-kuse, names_to = "variable") %>%
 group_by(variable) %>%
 nest() %>%
 mutate(wilcoxon = map(data, ~ wilcox.test(.$value ~ .$kuse, correct = F))) %>%
 mutate(wilcoxon.stat = map(wilcoxon, ~ round(.$statistic, 3))) %>%
 mutate(wilcoxon.pvalue = map(wilcoxon, ~ round(.$p.value, 3))) %>%
 unnest(cols = c(wilcoxon.stat, wilcoxon.pvalue)) %>%
 mutate(ttest = map(data, ~ t.test(.$value ~ .$kuse, var.equal = T))) %>%
 mutate(t.stat = map(ttest, ~ round(.$statistic, 3))) %>%
 mutate(t.pvalue = map(ttest, ~ round(.$p.value, 3))) %>%
 unnest(cols = c(t.stat, t.pvalue)) %>%
 select(-data, -wilcoxon, -ttest)
## # A tibble: 6 x 5
## # Groups: variable [6]
   variable wilcoxon.stat wilcoxon.pvalue t.stat t.pvalue
    <chr>
                    <dbl>
                                    <dbl> <dbl>
                                                     <dbl>
## 1 mratio96
                                    0
                                          12.3
                 172618.
                                                     Ω
## 2 pcged97
                 147410.
                                   0
                                          12.3
                                                     0
## 3 pcg_adc
                                   0
                                          -11.0
                                                     0
                  69728
                                   0 -12.7
## 4 black
                   58038
                                   0.002 - 3.11
## 5 age97
                  87129
                                                    0.002
                                   0.636 0.473
## 6 male
                   101544
                                                   0.637
# SMD Balance Checks
(c1 <- cobalt::bal.tab(</pre>
 x = d %>% select(mratio96, pcged97, pcg_adc, black, age97, male),
 treat = d$kuse,
 binary = "std",
 threshold = c(m = .1),
 s.d.denom = "pooled"
))
## Balance Measures
             Type Diff.Un
                             M.Threshold.Un
## mratio96 Contin. -1.0158 Not Balanced, >0.1
## pcged97 Contin. -0.9067 Not Balanced, >0.1
## pcg_adc Contin. 0.6703 Not Balanced, >0.1
           Binary 0.9343 Not Balanced, >0.1
## black
           Contin. 0.2196 Not Balanced, >0.1
## age97
## male
          Binary -0.0335
                               Balanced, <0.1
##
## Balance tally for mean differences
                     count
## Balanced, <0.1
## Not Balanced, >0.1
                         5
## Variable with the greatest mean difference
## Variable Diff.Un M.Threshold.Un
## mratio96 -1.0158 Not Balanced, >0.1
```

```
##
## Sample sizes
## Control Treated
## All 729 274

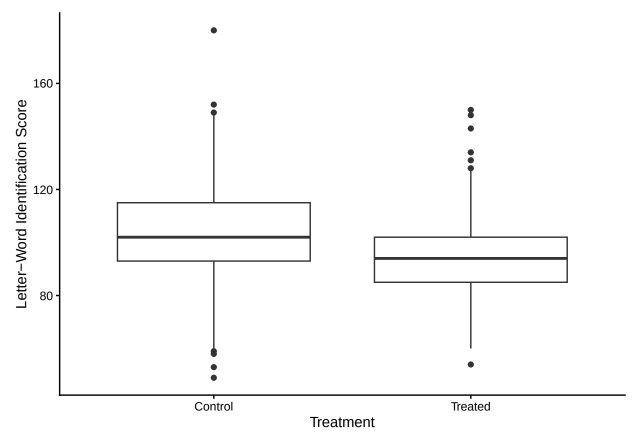
(cobalt::love.plot(c1, sample.names = c("Unmatched")))
```



Outcome Models Without Bias Control

We first observe that the control group had a higher letter-word identification score than the treatment group:

```
d %>%
  ggplot(aes(y = lwss97, x = factor(kuse))) +
  geom_boxplot() +
  labs(x = "Treatment", y = "Letter-Word Identification Score") +
  scale_x_discrete(labels = c("Control", "Treated")) +
  theme_classic()
```



We can estimate the ATE using an independent samples t test. It shows that the treated group on average had a mean letter-word identification score that was 9.82 points lower than that of the control group (p < .001).

```
t.test(lwss97 ~ kuse, data = d, var.equal = T) %>%
  broom::tidy()
## # A tibble: 1 x 10
     estimate estimate1 estimate2 statistic p.value parameter conf.low conf.high
##
##
        <dbl>
                  <dbl>
                             <dbl>
                                       <dbl>
                                                 <dbl>
                                                           <dbl>
                                                                    <dbl>
                                                                               <dbl>
         9.82
                   104.
                              94.2
                                        8.51 6.27e-17
                                                            1001
                                                                     7.56
                                                                                12.1
## # i 2 more variables: method <chr>, alternative <chr>
```

We can also estimate the ATE using an OLS regression model. It shows that, controlling for covariates, the treated group on average had a letter-word identification score that is 4.73 points lower than that of the control group (p < .001).

```
reg <- lm(lwss97 ~ kuse + male + black + age97 + pcged97 + mratio96 + pcg_adc,
 data = d
)
broom::tidy(lmtest::coeftest(reg, vcov = sandwich::vcovCL, cluster = d$pcg_id))
## # A tibble: 8 x 5
##
     term
                 estimate std.error statistic p.value
##
     <chr>>
                    <dbl>
                              <dbl>
                                         <dbl>
                                                  <dbl>
## 1 (Intercept)
                   84.9
                              4.50
                                         18.9 4.30e-68
## 2 kuse
                   -4.73
                              1.39
                                         -3.40 7.00e- 4
```

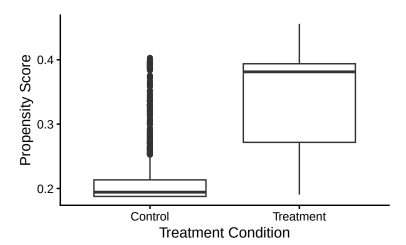
| ## 3 male | -2.00 | 0.987 | -2.02 4.34e- 2 |
|---------------|--------|-------|----------------|
| ## 4 black | -1.88 | 1.23 | -1.53 1.27e- 1 |
| ## 5 age97 | 0.873 | 0.170 | 5.14 3.30e- 7 |
| ## 6 pcged97 | 0.910 | 0.336 | 2.71 6.91e- 3 |
| ## 7 mratio96 | 1.13 | 0.332 | 3.41 6.67e- 4 |
| ## 8 pcg_adc | -0.758 | 0.311 | -2.44 1.50e- 2 |

Matching

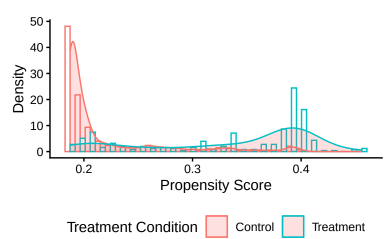
Note: To be consistent with the STATA result, we will use the propensity scores created by Stata.

Distribution of estimated propensity scores As seen in the figures below, there is no sizeable common support region of the estimated propensity scores by treatment status, so using greedy matching would produce a nontrivial loss of matched participants. Therefore, we decide to use optimal matching.

```
cds %>%
  ggplot(aes(x = factor(kuse), y = ps)) +
  geom_boxplot() +
  labs(y = "Propensity Score", x = "Treatment Condition") +
  scale_x_discrete(labels = c("Control", "Treatment")) +
  theme_classic()
```



```
cds %>%
  ggplot(aes(x = ps, color = factor(kuse))) +
  geom_histogram(aes(y = ..density..), fill = "white", position = "dodge") +
  geom_density(alpha = .2, fill = "#FF6666") +
  labs(y = "Density", x = "Propensity Score", color = "Treatment Condition") +
  scale_color_discrete(labels = c("Control", "Treatment")) +
  theme_classic() +
  theme(legend.position = "bottom")
```



We would lose 64.3% cases if we used nearest neighbor matching without replacement within specified caliper widths:

```
(nn_test <- matchit(kuse ~ mratio96 + pcged97 + pcg_adc + black + age97 + male,</pre>
  data = d,
 method = "nearest",
 distance = "glm",
 replace = F,
  caliper = .25
))
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred
## A 'matchit' object
## - method: 1:1 nearest neighbor matching without replacement
## - distance: Propensity score [caliper]
##
                - estimated with logistic regression
##
## - caliper: <distance> (0.07)
## - number of obs.: 1003 (original), 358 (matched)
## - target estimand: ATT
## - covariates: mratio96, pcged97, pcg_adc, black, age97, male
(nrow(d) - nrow(match.data(nn_test))) / nrow(d)
```

[1] 0.6430708

Optimal Matching The first argument that optmatch::fullmatch() requires is "a matrix of non-negative discrepancies, each indicating the permissibility and desirability of matching the unit corresponding to its row (treated) to the unit corresponding to its column (control)." First, we calculate the sample ranks of the propensity scores generated by generalized boosted regression using rank(ps). Second, we use outer(x, y, "-"), which will subtract y control units from x treated units, to form a matrix with rows equal to the number of treated subjects and columns equal to the number of control subjects. Finally, we take the absolute value of these differences.

By default, min.controls or the minimum ratio of controls to treatments that is to be permitted within a matched set is zero, and max.controls or the maximum ratio of controls to treatments is infinite.

Hansen found that in the context of a specific application, variable matching with a specific structure worked best; that is, each treated participant was matched to at least $\frac{.5(1-\hat{P})}{\hat{P}}$ controls and at most $\frac{2(1-\hat{P})}{\hat{P}}$ controls, where \hat{P} represents the proportion of treated participants in the sample. In this example, $\hat{P} = \frac{274}{729+274} = 27.32\%$, so the number of minimum controls is $\frac{.5(1-.2732)}{.2732} = 1.33$, and the number of maximum controls is $\frac{2(1-.2732)}{.2732} = 5.32$. There is a variance trade-off when selecting a variable number of controls, because the additional controls are generally less similar than the first closest match as the treated individual, which would increase bias, while the increase in sample size can decrease variance. ¹⁵

For optmatch::pairmatch(), we can set the number of controls to be matched to each treatment using the controls argument.

```
attach(cds)
prank <- rank(ps)</pre>
names(prank) <- kid</pre>
d1 <- outer(prank[kuse == 1], prank[kuse == 0], "-")
d1 \leftarrow abs(d1)
table(cds$kuse) # 729 control, 274 treated
dim(d1) # 274 x 729
# Full matching
fm <- fullmatch(d1)
(fm.d <- matched.distances(fm, d1, pres = TRUE))</pre>
max(unlist(fm.d))
# Variable Matching (at least 1, at most 4)
vm1 <- fullmatch(d1, min.controls = 1, max.controls = 4)</pre>
(vm1.d <- matched.distances(vm1, d1, pres = TRUE))</pre>
max(unlist(vm1.d))
# Variable Matching (at least 2, at most 4)
vm2 <- fullmatch(d1, min.controls = 2, max.controls = 4)</pre>
(vm2.d <- matched.distances(vm2, d1, pres = TRUE))</pre>
max(unlist(vm2.d))
# Variable Matching (using Hensen's equation)
vm3 <- fullmatch(d1, min.controls = 1.33, max.controls = 5.32)
(vm3.d <- matched.distances(vm3, d1, pres = TRUE))</pre>
max(unlist(vm3.d))
# Variable Matching (at least 2, at most 7)
vm4 <- fullmatch(d1, min.controls = 2, max.controls = 7)</pre>
```

 $^{^{15}\}mathrm{Stuart},$ 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

```
(vm4.d <- matched.distances(vm4, d1, pres = TRUE))
max(unlist(vm4.d))

# 1:1 Pair Matching
pm <- pairmatch(d1, controls = 1)
(pm.d <- matched.distances(pm, d1, pres = TRUE))
max(unlist(pm.d))</pre>
```

Optimal matching aims to minimize the total sample distance of propensity scores. We can extract the distances of matched units from their matched counterpairs using optmatch::matched.distances() and then calculate the total distance using sum(unlist()). The ratio of treatment to control and the number of strata for each structure can be inspected with optmatch::stratumStructure().

```
mean(unlist(fm.d))
## [1] 35.89389
sum(unlist(fm.d))
## [1] 31120
stratumStructure(fm)
         13:1 11:1 10:1
                              9:1
                                    7:1
                                                                                1:2
##
    25:1
                                          6:1
                                                 5:1
                                                       4:1
                                                             3:1
                                                                    2:1
                                                                          1:1
##
       1
             1
                   3
                          1
                                1
                                      1
                                            3
                                                   2
                                                         2
                                                               6
                                                                           47
                                                                                 12
                                                1:10 1:11 1:15
                                                                 1:17
                                                                         1:18
##
     1:3
           1:4
                 1:5
                        1:6
                              1:7
                                    1:8
                                           1:9
                                                                               1:21
                   7
                          5
                                2
                                      4
                                            3
                                                         3
                                                               1
##
       7
             6
                                                   2
         1:31 1:62 1:187
##
    1:24
##
             1
                    1
       1
mean(unlist(vm1.d))
## [1] 328.7579
sum(unlist(vm1.d))
## [1] 239664.5
stratumStructure(vm1)
## 1:1 1:3 1:4
## 122
         1 151
mean(unlist(vm2.d))
```

[1] 348.1852

```
sum(unlist(vm2.d))
## [1] 253827
stratumStructure(vm2)
## 1:2 1:3 1:4
## 183 1 90
mean(unlist(vm3.d))
## [1] 269.7222
sum(unlist(vm3.d))
## [1] 196627.5
stratumStructure(vm3)
## 1:1 1:6
## 183 91
mean(unlist(vm4.d))
## [1] 312.3162
sum(unlist(vm4.d))
## [1] 227678.5
stratumStructure(vm4)
## 1:2 1:3 1:7
## 237 1 36
mean(unlist(pm.d))
## [1] 143.2263
sum(unlist(pm.d))
## [1] 39244
stratumStructure(pm)
## 1:1 0:1
```

274 455

```
# Define covariates
cov_labels <- tibble(</pre>
  cov = c(
    "mratio96", "pcged97", "pcg_adc",
    "black", "age97", "male"
  ),
  new = c(
    "Ratio of family income to poverty line in 1996",
    "Caregiver's education in 1997 (years of schooling)",
    "Caregiver's number of years using AFDC in childhood",
    "Child's race: African American (reference: other)",
    "Child's age in 1997", "Child's gender: male (reference: female)"
  order = c(1, 2, 3, 4, 5, 6)
# Calculate dx and dxm using imbalance()
arg1 <- rep(c("mratio96", "pcged97", "pcg_adc", "black", "age97", "male"), each = 7)</pre>
arg2 <- rep(c("before", "fm", "vm1", "vm2", "vm3", "vm4", "pm"), length(arg1) / 7)</pre>
table_5.10 <- map2_dfr(arg1, arg2, imbalance) %>%
  mutate(method = recode(method,
    `before` = "Before Matching",
    `fm` = "Full Matching",
    `vm1` = "Variable Matching 1 (at least 1, at most 4)",
    `vm2` = "Variable Matching 2 (at least 2, at most 4)",
    `vm3` = "Variable Matching 3 (Hansen's equation)",
    `vm4` = "Variable Matching 4 (at least 2, at most 7)",
    `pm` = "Pair matching"
  )) %>%
  left_join(cov_labels, by = "cov") %>%
  arrange(order) %>%
  select(cov, matching_scheme = method, dx, dxm) # or replace cov with new
# Print Table
print(table_5.10, n = 42)
```

Covariate Imbalance Before and After Matching by Matching Scheme

```
## # A tibble: 42 x 4
##
                                                                dx
                                                                        dxm
      COV
              matching_scheme
##
      <chr>
              <chr>
                                                             <dbl>
                                                                      <dbl>
## 1 mratio96 Before Matching
                                                            1.02
                                                                  NΑ
## 2 mratio96 Full Matching
                                                                    0.0408
## 3 mratio96 Variable Matching 1 (at least 1, at most 4) NA
                                                                    0.790
## 4 mratio96 Variable Matching 2 (at least 2, at most 4) NA
                                                                    0.940
## 5 mratio96 Variable Matching 3 (Hansen's equation)
                                                                    0.736
## 6 mratio96 Variable Matching 4 (at least 2, at most 7) NA
                                                                    0.869
## 7 mratio96 Pair matching
                                                           NΑ
                                                                    0.253
## 8 pcged97 Before Matching
                                                            0.907 NA
## 9 pcged97 Full Matching
                                                                    0.168
## 10 pcged97 Variable Matching 1 (at least 1, at most 4) NA
                                                                    0.722
## 11 pcged97 Variable Matching 2 (at least 2, at most 4) NA
                                                                    0.867
```

```
Variable Matching 3 (Hansen's equation)
                                                                      0.752
## 12 pcged97
               Variable Matching 4 (at least 2, at most 7)
## 13 pcged97
                                                             NΑ
                                                                      0.814
## 14 pcged97
               Pair matching
                                                             NA
                                                                      0.340
                                                              0.670
## 15 pcg_adc
               Before Matching
                                                                     NΑ
## 16 pcg_adc
               Full Matching
                                                             NΑ
                                                                      0.00737
## 17 pcg adc
               Variable Matching 1 (at least 1, at most 4)
                                                                      0.560
                                                             NA
               Variable Matching 2 (at least 2, at most 4)
## 18 pcg adc
                                                                      0.649
               Variable Matching 3 (Hansen's equation)
## 19 pcg_adc
                                                                      0.551
## 20 pcg_adc
               Variable Matching 4 (at least 2, at most 7)
                                                                      0.626
## 21 pcg_adc
               Pair matching
                                                             NA
                                                                      0.403
## 22 black
               Before Matching
                                                              0.933
                                                                     NA
## 23 black
               Full Matching
                                                             NA
                                                                      0.0106
## 24 black
               Variable Matching 1 (at least 1, at most 4)
                                                                      0.788
                                                             NΑ
## 25 black
               Variable Matching 2 (at least 2, at most 4)
                                                                      0.919
               Variable Matching 3 (Hansen's equation)
## 26 black
                                                                      0.732
## 27 black
               Variable Matching 4 (at least 2, at most 7)
                                                                      0.848
## 28 black
               Pair matching
                                                             NA
                                                                      0.442
## 29 age97
               Before Matching
                                                              0.220
                                                                     NA
                                                                      0.0690
## 30 age97
               Full Matching
                                                             NA
## 31 age97
               Variable Matching 1 (at least 1, at most 4)
                                                                      0.215
## 32 age97
               Variable Matching 2 (at least 2, at most 4)
                                                                      0.219
               Variable Matching 3 (Hansen's equation)
## 33 age97
                                                                      0.190
               Variable Matching 4 (at least 2, at most 7) NA
## 34 age97
                                                                      0.238
## 35 age97
               Pair matching
                                                             NA
                                                                      0.137
## 36 male
               Before Matching
                                                              0.0335 NA
## 37 male
               Full Matching
                                                             NA
                                                                      0.0532
               Variable Matching 1 (at least 1, at most 4)
                                                                      0.0219
## 38 male
## 39 male
               Variable Matching 2 (at least 2, at most 4)
                                                                      0.0487
               Variable Matching 3 (Hansen's equation)
## 40 male
                                                                      0.00974
## 41 male
               Variable Matching 4 (at least 2, at most 7)
                                                                      0.0325
                                                             NA
## 42 male
               Pair matching
                                                             NA
                                                                      0.0873
```

Post-Full-Matching Analysis of Outcome

Hodges-Lehmann Aligned Rank Test The Hodges-Lehmann aligned rank test can be used on matched samples created by full matching or variable matching. We used full matching and found that children who used AFDC had a letter-word identification score in 1997 that was on average 1.97 points lower than those who had never used AFDC; the difference was statistically significant at a .05 level (one-tailed). We used the Hodges-Lehmann test to gauge the statistical significance. The study also detected an effect size of .19, which is a small effect size in terms of Cohen's (1988) criteria.

```
hodgesl(df, lwss97, fm, kuse)
```

```
## # A tibble: 1 x 6
## HL_mean HL_se z p i tx_effect
## <dbl> <dbl> <dbl> <int> <dbl> +int> <dbl> = -7039. 3247. -2.17 0.0151 1 -1.97
```

```
imbalance2(df, lwss97, kuse, fm)$dxm # dxm for the outcome variable is Cohen's d
```

[1] 0.1896308

Post-Matching Analysis Using Regression of Difference Scores

A regression of difference scores may be performed on matched samples created by optimal pair matching. Based on the pair-matched sample, we first calculated difference scores between treated and control cases for each pair on all study variables (i.e., on the outcome variable and all covariates). We then regressed the difference score of the outcome on the difference scores of covariates. In addition, note that our model includes a correction for clustering effects (i.e., children are nested within caregivers) that we accomplished by using robust estimates of standard errors. The intercept of a difference score regression indicates the ATE of the sample. The estimated intercept from this model is -3.17 (p < .05). Thus, using pair matching and regression adjustment, the study found that, on average, children who used AFDC had a letter-word identification score in 1997 that was 3.17 points lower than do children who never used AFDC; this finding was statistically significant.

```
# kuse == 1
df1 <- df %>%
  select(pm, kuse,
   y1 = lwss97, male1 = male, black1 = black, age971 = age97,
   pcged971 = pcged97, mratio961 = mratio96, pcg_id
  ) %>%
  filter(kuse == 1)
# kuse == 0
df0 <- df %>%
  select(pm, kuse,
   y0 = lwss97, male0 = male, black0 = black, age970 = age97,
   pcged970 = pcged97, mratio960 = mratio96, pcg_id
  ) %>%
 filter(kuse == 0)
# Merge Data
df2 <- left join(df0, df1, "pm")
df2 <- df2 %>%
  mutate(
   y = y1 - y0,
   male = male1 - male0,
   black = black1 - black0,
   age97 = age971 - age970,
   pcged97 = pcged971 - pcged970,
   mratio96 = mratio961 - mratio960
  )
# Regression of Difference Scores (one-tailed test)
reg1 <- lm(y ~ male + black + age97 + pcged97 + mratio96, data = df2)
broom::tidy(lmtest::coeftest(reg1, vcov = vcovCL, cluster = df2$pcg_id.x)) %>%
 mutate(p.value.one.tailed = p.value / 2)
```

```
## 1 (Intercept) -3.17
                              1.73
                                      -1.83
                                               0.0682
                                                                  0.0341
## 2 male
                  -1.21
                              1.84
                                      -0.657
                                               0.512
                                                                  0.256
## 3 black
                  -3.34
                              2.33
                                      -1.43
                                               0.153
                                                                  0.0765
                                       0.695
## 4 age97
                   0.229
                              0.329
                                                                  0.244
                                               0.487
## 5 pcged97
                   0.0111
                              0.508
                                       0.0219 0.983
                                                                  0.491
## 6 mratio96
                                                                  0.0301
                   3.15
                              1.67
                                       1.89
                                               0.0602
```

Optimal (Pair) Matching Using the MatchIt Package

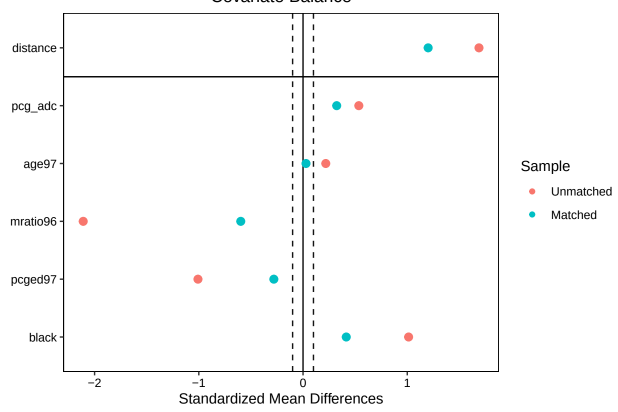
For an easier interface for the optmatch package, the MatchIt::matchit() function also implements optimal full matching and optimal pair matching with the method = "full" and method = "optimal" arguments, respectively. Variable matching can be set using the ratio, min.controls, and max.controls arguments.

Below is an illustration of optimal pair matching (by default, 1:1 without replacement) using generalized boosted model to generate the propensity scores and then a regression model for the outcome that incorporates the matching weights. Here we find that on average children who used AFDC had a letter-word identification score in 1997 that was 4.13 points lower than do children who never used AFDC (p < .001).

```
# Optimal 1:1 Pair Matching Without Replacement Using GBM (ATT)
set.seed(1000)
(pm2 <- matchit(</pre>
  formula = kuse ~ pcg_adc + age97 + mratio96 + pcged97 + black,
  data = d,
  method = "optimal",
 replace = F,
  ratio = 1,
  distance = "gbm",
  estimand = "ATT"
))
## A 'matchit' object
## - method: 1:1 optimal pair matching
   - distance: Propensity score
##
                - estimated with GBM
##
  - number of obs.: 1003 (original), 548 (matched)
## - target estimand: ATT
## - covariates: pcg_adc, age97, mratio96, pcged97, black
# Balance Checks
cobalt::bal.tab(pm2, binary = "std", un = T, threshold = c(m = .1))
## Balance Measures
##
                                             M.Threshold
                Type Diff.Un Diff.Adj
## distance Distance 1.6880
                               1.2004
             Contin. 0.5352
                               0.3230 Not Balanced, >0.1
## pcg_adc
             Contin. 0.2181
                               0.0285
                                          Balanced, <0.1
## age97
## mratio96 Contin. -2.1090 -0.5976 Not Balanced, >0.1
## pcged97
             Contin. -1.0082
                              -0.2802 Not Balanced, >0.1
              Binary 1.0129
## black
                              0.4148 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 -0.5976 Not Balanced, >0.1
##
##
## Sample sizes
             Control Treated
##
## All
                 729
## Matched
                 274
                         274
## Unmatched
                 455
cobalt::love.plot(pm2, un = T, binary = "std", threshold = c(m = .1),
                  sample.names = c("Unmatched", "Matched"))
```

Covariate Balance



```
broom::tidy(lmtest::coeftest(pm2.lm1, vcov = vcovCL, cluster = md$pcg_id))
## # A tibble: 2 x 5
##
    term
               estimate std.error statistic p.value
    <chr>>
                <dbl> <dbl> <dbl>
## 1 (Intercept)
               100.
                          1.09
                                  91.5 0
## 2 kuse
                 -5.84
                           1.47
                                  -3.98 0.0000774
broom::tidy(lmtest::coeftest(pm2.lm2, vcov = vcovCL, cluster = md$pcg_id))
## # A tibble: 12 x 5
##
     term
                 estimate std.error statistic p.value
                   <dbl> <dbl> <dbl>
##
     <chr>
                                              <dbl>
## 1 (Intercept)
                   81.5
                             8.55
                                     9.54 5.01e-20
                           11.5
                                     0.321 7.48e- 1
## 2 kuse
                    3.68
## 3 male
                           1.87
                   -4.47
                                     -2.39 1.72e- 2
## 4 black
                             2.36
                                   -0.731 4.65e- 1
                   -1.72
## 5 age97
                   0.792 0.335
                                   2.37 1.82e- 2
                                     1.39 1.65e- 1
## 6 pcged97
                   0.977 0.703
## 7 mratio96
                   2.83
                           1.05
                                      2.71 7.04e- 3
## 8 kuse:male
                   3.96
                           2.66
                                     1.49 1.37e- 1
                   -1.02
                            3.44
                                     -0.296 7.68e- 1
## 9 kuse:black
                   -0.475 0.458
                                     -1.04 3.00e- 1
## 10 kuse:age97
## 11 kuse:pcged97
                                     -0.366 7.15e- 1
                   -0.343 0.937
## 12 kuse:mratio96 -1.18 1.44
                                     -0.819 4.13e- 1
broom::tidy(lmtest::coeftest(pm2.lm3, vcov = vcovCL, cluster = md$pcg_id))
## # A tibble: 8 x 5
##
               estimate std.error statistic p.value
    term
    <chr>
                <dbl>
                          <dbl>
                                  <dbl>
                                           <dbl>
                                  12.8 7.32e-33
## 1 (Intercept) 89.5
                          7.00
## 2 kuse
               -3.11
                         1.90
                                  -1.64 1.02e- 1
               -2.55
                                  -1.98 4.86e- 2
## 3 male
                          1.29
                                 -0.638 5.24e- 1
## 4 black
                -1.21
                          1.89
## 5 age97
                0.576 0.229
                                  2.51 1.23e- 2
## 6 pcged97
                0.595 0.484
                                  1.23 2.19e- 1
## 7 mratio96
                1.59
                          1.10
                                   1.45 1.46e- 1
## 8 distance
                -4.15
                          4.85
                                  -0.856 3.92e- 1
broom::tidy(lmtest::coeftest(pm2.lm4, vcov = vcovCL, cluster = md$pcg_id))
## # A tibble: 7 x 5
   term
              estimate std.error statistic p.value
    <chr>>
                <dbl> <dbl> <dbl>
                                           <dbl>
                          5.83
                                  14.7 2.35e-41
## 1 (Intercept) 85.6
## 2 kuse
               -3.98
                          1.53
                                  -2.60 9.68e- 3
## 3 male
               -2.56
                          1.29
                                  -1.99 4.73e- 2
## 4 black
                -1.90
                          1.72
                                   -1.11 2.69e- 1
                                   2.47 1.37e- 2
## 5 age97
                0.563 0.228
## 6 pcged97
                0.778
                         0.463
                                   1.68 9.39e- 2
## 7 mratio96
                                   3.12 1.93e- 3
                2.23
                          0.717
```

```
# Estimate ATT with marginaleffects::avg_comparisons() (G-computation)
marginaleffects::avg_comparisons(pm2.lm4,
    variables = "kuse", # treatment
    vcov = ~subclass + pcg_id, # cluster-robust SEs
    newdata = subset(md, kuse == 1), # ATT
    wts = "weights"
)

##
## Estimate Std. Error    z Pr(>|z|)    S 2.5 % 97.5 %
##    -3.98    1.55 -2.57    0.0101 6.6 -7.01 -0.949
##
## Term: kuse
```

Type: response
Comparison: 1 - 0

5.5 Computer Lab: Running the IPTW Estimator with R

Load Packages

Propensity score weighting can be accomplished with base R functions. 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.

pacman::p_load(lmtest, sandwich)

Description of Dataset

This dataset is from a study that investigates intergenerational dependence on welfare and its relation to child academic achievement. 16

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.

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). Propensity score weighting for estimating ATE is generally referred to as the inverse probability of treatment weights (IPTW) estimator.

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:

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

¹⁶Hofferth, 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.

Load Data with Propensity Scores and Calculate Weights

```
psw_df <- read_dta("data/chpt5_2_original.dta") %>%
  haven::zap_formats() %>%
  sjlabelled::remove_all_labels() %>%
  as_tibble() %>%
  mutate(
   ate_w = ifelse(kuse == 0, 1 / (1 - ps), 1 / ps),
   att_w = ifelse(kuse == 0, ps / (1 - ps), 1)
)
```

Check Balance

In addition to standardized mean differences, variance ratios can be requested. Common thresholds for balanced groups are .5 and 2, but it will be close to 1 when group variances are similar. Remember to use 's.d.denom = "pooled" for the ATE and 's.d.denom = "treated" for the ATT.

```
# ATE
cobalt::bal.tab(
    x = select(psw_df, male, black, age97, pcged97, mratio96),
    treat = psw_df$kuse,
    binary = "std",
    continuous = "std",
    s.d.denom = "pooled",
    un = T,
    stats = c("mean.diffs", "variance.ratios"),
    thresholds = c(m = .1, v = 2)
)
```

```
## Balance Measures
##
              Type Diff.Un
                               M.Threshold.Un V.Ratio.Un V.Threshold.Un
            Binary -0.0335
## male
                               Balanced, <0.1
           Binary 0.9343 Not Balanced, >0.1
## black
## age97
           Contin. 0.2196 Not Balanced, >0.1
                                                  1.0266
                                                             Balanced, <2
## pcged97 Contin. -0.9067 Not Balanced, >0.1 0.6789
                                                             Balanced, <2
## mratio96 Contin. -1.0158 Not Balanced, >0.1 0.1312 Not Balanced, >2
## Balance tally for mean differences
                     count
## Balanced, <0.1
                         1
## Not Balanced, >0.1
##
## Variable with the greatest mean difference
## Variable Diff.Un
                        M.Threshold.Un
## mratio96 -1.0158 Not Balanced, >0.1
##
## Balance tally for variance ratios
##
                   count
## Balanced, <2
## Not Balanced, >2
## Variable with the greatest variance ratio
```

```
## Variable V.Ratio.Un V.Threshold.Un
## mratio96 0.1312 Not Balanced, >2
##
## Sample sizes
## Control Treated
## All 729
                    274
# ATT
cobalt::bal.tab(
  x = select(psw_df, male, black, age97, pcged97, mratio96),
  treat = psw df$kuse,
  binary = "std",
  continuous = "std",
  s.d.denom = "treated",
  un = T,
  stats = c("mean.diffs", "variance.ratios"),
  thresholds = c(m = .1, v = 2)
## Balance Measures
        Type Diff.Un
                                  M.Threshold.Un V.Ratio.Un V.Threshold.Un
## male
            Binary -0.0335
                                  Balanced, <0.1
## black
            Binary 1.0129 Not Balanced, >0.1
## age97 Contin. 0.2181 Not Balanced, >0.1 1.0266 Balanced, <2
## pcged97 Contin. -1.0082 Not Balanced, >0.1 0.6789 Balanced, <2
## mratio96 Contin. -2.1090 Not Balanced, >0.1 0.1312 Not Balanced, >2
## Balance tally for mean differences
                       count
## Balanced, <0.1
                        1
## Not Balanced, >0.1
##
## Variable with the greatest mean difference
## Variable Diff.Un M.Threshold.Un
## mratio96 -2.109 Not Balanced, >0.1
## Balance tally for variance ratios
##
                   count
## Balanced, <2
                         2
## Not Balanced, >2
##
## Variable with the greatest variance ratio
## Variable V.Ratio.Un V.Threshold.Un
## mratio96
                0.1312 Not Balanced, >2
##
## Sample sizes
##
       Control Treated
## All
         729
                    274
```

Calculate Weights with the WeightIt Package

```
# Load Package
pacman::p_load(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))

##
## FALSE TRUE
## 327 676</pre>
```

Outcome Analysis

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.

```
# ATE
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))
##
## t test of coefficients:
##
##
            Estimate Std. Error t value
                                                Pr(>|t|)
## kuse
            -5.16399
                       1.42438 -3.6254
                                               0.0003031 ***
            -1.62201
                       1.09186 -1.4855
                                               0.1377180
## male
                      1.34670 -1.8556
## black
            -2.49898
                                               0.0638009 .
            0.73868 0.18075 4.0867
                                              0.00004727 ***
## age97
## pcged97
            0.99264
                       0.35596 2.7886
                                               0.0053938 **
## mratio96 1.13856
                       0.32220 3.5337
                                               0.0004286 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

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.

```
# ATT
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|)
                          5.05331 16.8790 < 0.000000000000000022 ***
## (Intercept) 85.29467
## kuse
               -4.62058
                           1.41182 -3.2728
                                                        0.001102 **
## male
              -1.58995
                          1.14705 -1.3861
                                                        0.166020
## black
               -2.74605
                           1.41605 -1.9392
                                                        0.052756 .
                           0.20001 3.0787
## age97
               0.61577
                                                        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 '** 0.01 '* 0.05 '.' 0.1 ' 1
```

Check Balance

To assess balance before and after propensity score weighting, we can use weighted logistic regression for the dummy covariates and weighted simple regression for the 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.

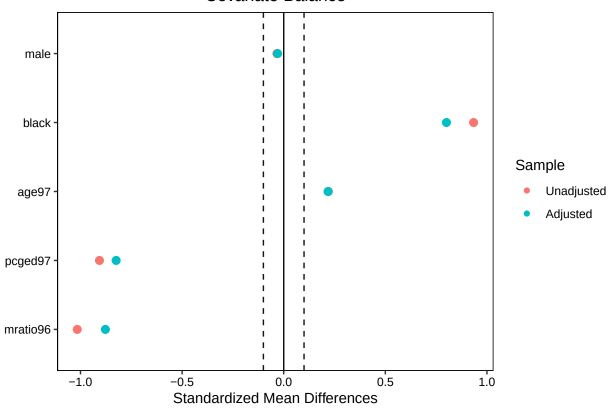
```
psw_c1 <- glm(male ~ kuse, family = quasibinomial, data = psw_df, weights = ate_w)</pre>
lmtest::coeftest(psw_c1, vcov. = vcovCL(psw_c1, cluster = psw_df$pcg_id))
##
## z test of coefficients:
##
##
                Estimate Std. Error z value Pr(>|z|)
## (Intercept) 0.142911
                    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

```
##
## z test of coefficients:
##
               Estimate Std. Error z value Pr(>|z|)
##
## (Intercept) 0.152522 0.077410 1.9703 0.0488 *
              -0.079497 0.147480 -0.5390 0.5899
## kuse
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 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$pcg_id))
## t test of coefficients:
##
##
              Estimate Std. Error t value
                                                       Pr(>|t|)
## (Intercept) 6.51389 0.10900 59.7615 < 0.00000000000000022 ***
              0.61064
                          0.21883 2.7905
                                                       0.005362 **
## kuse
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
psw_c4 <- lm(age97 ~ kuse, weights = att_w, data = psw_df)</pre>
lmtest::coeftest(psw_c4, vcov. = vcovCL(psw_c4, cluster = psw_df$pcg_id))
##
## t test of coefficients:
##
              Estimate Std. Error t value
##
                                                      Pr(>|t|)
## (Intercept) 6.55501 0.11499 57.0057 < 0.00000000000000002 ***
               0.56178
                          0.21939 2.5606
## kuse
                                                       0.01059 *
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
Alternatively, balance can be assessed using the cobalt package.
# ATE
(psw_b1 <- cobalt::bal.tab(</pre>
 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", "variance.ratios"),
 thresholds = c(m = .1, v = 2)
))
## Balance Measures
##
              Type Diff.Un V.Ratio.Un Diff.Adj
                                                      M.Threshold V.Ratio.Adj
           Binary -0.0335 . -0.0301
                                                   Balanced, <0.1
## male
## black
           Binary 0.9343
                                 . 0.8006 Not Balanced, >0.1
```

```
## age97 Contin. 0.2196 1.0266 0.2181 Not Balanced, >0.1 ## pcged97 Contin. -0.9067 0.6789 -0.8244 Not Balanced, >0.1
                                                                          1.0089
                                                                         0.6940
                                                                           0.1781
## mratio96 Contin. -1.0158 0.1312 -0.8775 Not Balanced, >0.1
               V.Threshold
## male
## black
## age97
                Balanced, <2
              Balanced, <2
## pcged97
## mratio96 Not Balanced, >2
##
## Balance tally for mean differences
                      count
## Balanced, <0.1
                          1
## Not Balanced, >0.1
##
## Variable with the greatest mean difference
## Variable Diff.Adj
                              M.Threshold
## mratio96 -0.8775 Not Balanced, >0.1
## Balance tally for variance ratios
                   count
##
## Balanced, <2
## Not Balanced, >2
                         1
## Variable with the greatest variance ratio
## Variable V.Ratio.Adj
                             V.Threshold
## mratio96
                  0.1781 Not Balanced, >2
## Effective sample sizes
              Control Treated
## Unadjusted 729.
                        274.
## Adjusted
               725.43 257.71
```

cobalt::love.plot(psw_b1)



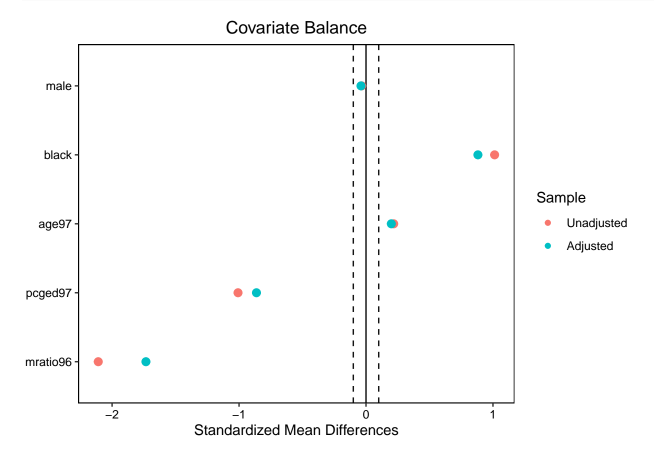


```
# ATT
(psw_b2 <- cobalt::bal.tab(
    x = select(psw_df, male, black, age97, pcged97, mratio96),
    treat = psw_df$kuse,
    weights = psw_df$att_w,
    binary = "std",
    continuous = "std",
    s.d.denom = "treated",
    un = T,
    stats = c("mean.diffs", "variance.ratios"),
    thresholds = c(m = .1, v = 2)
))</pre>
```

```
## Balance Measures
             Type Diff.Un V.Ratio.Un Diff.Adj
##
                                                  M.Threshold V.Ratio.Adj
## male
          Binary -0.0335 . -0.0396
                                               Balanced, <0.1
          Binary 1.0129
                                . 0.8814 Not Balanced, >0.1
## black
                           1.0266 0.1993 Not Balanced, >0.1
          Contin. 0.2181
## age97
                                                                  1.0085
## pcged97 Contin. -1.0082 0.6789 -0.8629 Not Balanced, >0.1
                                                                  0.6694
## mratio96 Contin. -2.1090 0.1312 -1.7335 Not Balanced, >0.1
                                                                  0.1463
##
               V.Threshold
## male
## black
## age97
              Balanced, <2
## pcged97
              Balanced, <2
## mratio96 Not Balanced, >2
```

```
##
## 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 -1.7335 Not Balanced, >0.1
##
## Balance tally for variance ratios
##
                    count
## Balanced, <2
## Not Balanced, >2
##
## Variable with the greatest variance ratio
   Variable V.Ratio.Adj
                              V.Threshold
   mratio96
                  0.1463 Not Balanced, >2
##
## Effective sample sizes
##
              Control Treated
## Unadjusted 729.
                          274
## Adjusted
               663.65
                          274
```

cobalt::love.plot(psw_b2)



Using WeightIt

The WeightIt package can generate propensity scores and the appropriate weights in a single step.

```
# Estimate propensity scores and generate ATT weights
psw_g1 <- WeightIt::weightit(</pre>
 kuse ~ male + black + age97 + pcged97 + mratio96,
 data = psw_df, estimand = "ATT", method = "glm"
)
## Warning: Propensity scores numerically equal to 0 or 1 were estimated,
## indicating perfect separation and infinite parameter estimates. These
## may yield problems with inference. Consider trying a different 'link'.
## See 'help("method_glm", package = "WeightIt")' for details.
# Check balance
cobalt::bal.tab(
 psw_g1,
 binary = "std",
 continuous = "std",
 s.d.denom = "treated", # att
 un = T,
 stats = c("m"),
 thresholds = c(m = .1)
## Balance Measures
                                              M.Threshold
                 Type Diff.Un Diff.Adj
## prop.score Distance 1.5861 0.1758
## male Binary -0.0335 -0.0208
                                           Balanced, <0.1
             Binary 1.0129 0.0628
## black
                                           Balanced, <0.1
## age97
            Contin. 0.2181 0.0363
                                           Balanced, <0.1
## pcged97 Contin. -1.0082 0.0587
                                           Balanced, <0.1
## mratio96 Contin. -2.1090 -0.1460 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 -0.146 Not Balanced, >0.1
## Effective sample sizes
             Control Treated
                         274
## Unadjusted 729.
## Adjusted
               141.1
# Attach ATT weights to dataframe
psw_df2 <- psw_df %>% mutate(att_weights = psw_g1$weights)
# Fit outcome model and estimate treatment effect with cluster-robust SEs
```

```
psw_g2 \leftarrow lm(lwss97 \sim kuse + male + black + age97 + pcged97 + mratio96,
data = psw_df2, weights = att_weights
marginaleffects::avg_comparisons(
 psw_g2,
 variables = "kuse",
vcov = ~pcg_id,
newdata = subset(psw_df2, kuse == 1), # att
 wts = "att_weights"
##
## Estimate Std. Error z Pr(>|z|) S 2.5 % 97.5 %
## -3.94 1.67 -2.36 0.0182 5.8 -7.2 -0.671
##
## Term: kuse
## Type: response
## Comparison: 1 - 0
# How would you calculate the ATE weights using this approach?
```

6.9 Computer Lab: Running the Matching Estimators with R

Matching Estimators

Load Packages

A variety of matching estimators are implemented in the Matching package. Unlike the MatchIt package, which uses "subset selection" to arrive at a weighted subset of the units from the dataset, the Matching package uses "matching imputation" to impute potential outcomes using the observed outcomes of paired units.¹⁷

Because the assumptions about constant treatment effect and homoskedasticity may not be valid for certain types of data, we will use the lmtest package for the Breusch-Pagan Test to check this assumption.

```
pacman::p_load(Matching)
```

Note that the Matching::Match() function is intended to be used in conjunction with the MatchBalance() function. However, functions from the cobalt package also work well and tend to be cleaner in presentation:¹⁸

```
data(lalonde, package = "Matching")
ex_f <- as.formula(treat ~ age + I(age^2) + educ + I(educ^2) + black +
 hisp + married + nodegr + re74 + I(re74^2) + re75 + I(re75^2) +
  u74 + u75
ex_m1 <- glm(ex_f, family = binomial, data = lalonde)</pre>
ex_p <- ex_m1$fitted.values</pre>
ex_X <- ex_m1\fitted
ex_Y <- lalonde$re78</pre>
ex_Tr <- lalonde$treat
ex_rr \leftarrow Match(Y = ex_Y, Tr = ex_Tr, X = ex_X, M = 1)
summary(ex rr)
ex_mb <- MatchBalance(treat ~ age + I(age^2) + educ + I(educ^2) + black +
 hisp + married + nodegr + re74 + I(re74^2) + re75 + I(re75^2) +
  u74 + u75, data = lalonde, match.out = ex rr, nboots = 10)
(ex_bal <- cobalt::bal.tab(ex_rr, ex_f,</pre>
  data = lalonde, distance = ~ex p, un = T,
  binary = "std", threshold = .1
cobalt::love.plot(ex_bal, sample.names = c("Unmatched", "Matched"))
```

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:

- pcg_adc: Caregiver's History of Using Welfare (Number of Years; range: 0-7)
- age97: Child's Age in 1997

 $^{^{17}} https://kosukeimai.github.io/MatchIt/articles/matching-methods.html\\$

 $^{^{18} \}rm https://ngreifer.github.io/cobalt/articles/other-packages.html$

- mratio96: Ratio of Family Income to Poverty Line in 1996
- pcged97: Caregiver's Education in 1997 (Years of Schooling)
- male: Child's Gender: Male (1 = Male; 0 = Female)
- black: Child's Race: African American (1 = African American; 0 = Other)

Load Data

```
me_df <- haven::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 | pcged97 | pcg_adc | mratio96 | 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 |

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.

To carry out the Breusch-Pagan Test, first we regress the outcome variable on the matching variables using OLS:

```
# Regress outcome on treatment and matching variables using OLS
me_m1 <- lm(pcss97 ~ kuse + male + black + age97 + pcged97 + mratio96 + pcg_adc,
    data = me_df
)</pre>
```

Next, we can run the Breusch-Pagan test for each matching variable:

```
lmtest::bptest(me_m1, ~kuse, data = me_df, studentize = F)
lmtest::bptest(me_m1, ~male, data = me_df, studentize = F)
lmtest::bptest(me_m1, ~black, data = me_df, studentize = F)
lmtest::bptest(me_m1, ~age97, data = me_df, studentize = F) # significant
lmtest::bptest(me_m1, ~pcged97, data = me_df, studentize = F)
lmtest::bptest(me_m1, ~mratio96, data = me_df, studentize = F)
lmtest::bptest(me_m1, ~pcg_adc, data = me_df, studentize = F)
```

Or use a function to test every variable:

```
bp <- function(var, df, md) {
   lmtest::bptest(md, as.formula(paste0("~", var)), data = df, studentize = F) %>%
        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 | 0.92 |
| male | 3.07 | 0.08 |
| black | 1.43 | 0.23 |
| age97 | 25.27 | 0.00 |
| pcged97 | 0.19 | 0.66 |
| mratio 96 | 0.62 | 0.43 |
| pcg_adc | 0.29 | 0.59 |

Results from the Breusch-Pagan tests showed that the homoskedasticity assumption is not valid for child's age (age97) (p < .05) and indicated that the conditional variance of the outcome variable was not constant across levels of child's age. Based on this finding, we should use the robust variance estimator that allows for heteroskedasticity (i.e., the Var.calc argument in the Matching::Match() function).

Matching Estimators

Of the six matching variables in this example, 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. (Tip: When you have one or more continuous covariate in your matching, always use the bias-corrected matching estimator.)

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).

The M argument specifies the number of matches which should be found. The default is one-to-one matching (i.e., M=1). Abadie and Imbens suggest that M be small, and M=4 typically performs well in terms of mean-squared error.

If Var.calc = 0, then homoskedasticity is assumed. Use Var.calc = 4 to request the robust variance estimator using four matches. This algorithm developed by Abadie and Imbens (2002) includes a second matching procedure such that it matches treated units to treated units and control units to controls.

The estimand argument is by default "ATT" but can be set to "ATE" or "ATC". Typically, we are interested in the ATE or ATT. The ATT seeks to answer questions such as "How would treated patients' outcomes

differ, on average, had they not received treatment?"; the ATE is the treatment effect for both control and treated units; and the ATC, the average treatment effect for the controls, investigates the effect of the treatment for a population who did not receive treatment.¹⁹

The sample argument is a logical flag indicating whether the population or sample variance should be estimated. 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." In other words, the sample effect shows whether the program is successful in the sample at hand, while the population effect shows whether the same program would be successful in a second sample from the population.

Results from the Matching::Match() function are identical to Stata's nnmatch program.

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

Use the Y, Tr, and X, arguments in the Match() function to specify the outcome vector, the treatment vector, and the matrix of variables to match on, respectively. The X matrix may or may not contain the propensity score.

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

Get Estimators Individually

Note that by default matching is done with replacement. However, this can be changed with the replace argument. A matched dataset can be recovered by using the index.treated and index.control vectors. Further, index.dropped contains observations that have been dropped (e.g., due to a caliper setting).

```
# Sample Average Treatment Effect (SATE)
me1 <- Match(
 Y = me_Y, Tr = me_Tr, X = me_X, M = 4, BiasAdjust = T, Var.calc = 4,
  estimand = "ATE", sample = T
)
summary(me1)
head(me1$index.treated)
head(me1$index.control)
head(me1$index.dropped)
head(me1$se) # Abadie-Imbens standard error
# 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
```

¹⁹https://arxiv.org/abs/2106.10577

 $^{^{20}} https://journals.sagepub.com/doi/pdf/10.1177/1536867X0400400307$

```
))
# 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
))
```

Get All Estimators

```
# Function for extracting estimate, SE, t-stat, and p-value from Match()
get_match <- function(estimand, sample, Y, Tr, X) {</pre>
  m <- Matching::Match(</pre>
    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
  ))
}
# 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, me_Y, me_Tr, me_X))) %>%
  unnest_wider(match) %>%
  select(-estimand, -sample) %>%
    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 |

The results suggest that childhood poverty strongly affected children's academic achievement.

ATE: On average, children who used AFDC in childhood had a passage comprehension score 4.7 units lower than that of children who had never used AFDC in childhood. This effect is statistically significant in the sample at hand (SATE p < .05) as well as in a second sample drawn from the same population (PATE p < .05).

ATT: With regard to the subpopulation of treated participants, on average, children who used AFDC in childhood had a passage comprehension score 5.2 units lower than that of children who had never used AFDC in childhood. This effect is statistically significant in the sample at hand (SATT p < .05) as well as in a second sample drawn from the same population (PATT p < .05).

ATC: Had all controls (i.e., children who never used AFDC) used AFDC and all treated children had not used AFDC, then on average, the control children would have a passage comprehension score 4.5 units lower than their counterparts (SATC p < .05; PATC p < .05).

Additional observations:

- 1. A population effect indicates whether the tested intervention will be effective in a second sample taken from the same population. Taking SATT and PATT as examples, the study indicated that the treatment effect for the treated group was statistically significant in the sample at a level of .01. If we take a second sample from the population, we are likely to observe the same level of treatment effect for the treated, and the effect should remain statistically significant at a level of .01. The point estimate of the population effect is identical to the point estimate of its corresponding sample effect. A population effect differs from its corresponding sample effect on variance, and thus significance test on a population effect may have a different conclusion than that on its corresponding sample effect. Note that if treated units are discarded (e.g., due to a caliper), the estimand no longer estimates a particular population but rather only the average treatment effect on the remaining matched units (ATM).
- 2. Note that in this study, SATT = -5.23 and SATC = -4.47, or a difference of 0.76 units. This difference is attributable either to additional selection bias that was not accounted for in the study or to study data that violated assumptions of matching estimators, which suggests the need for further scrutiny.
- 3. All six treatment effects were statistically significant (p < .05). Thus, we can conclude that the study data could not reject a null hypothesis of a zero treatment effect, and childhood poverty appears to be an important factor causing children's poor achievement in passage comprehension.

Specify Variables in the Bias-Corrected Matching

The Z argument can be used to specify the covariates for which we wish to make bias adjustments.

```
# Sample Average Treatment Effect (SATE)
me_Z <- select(me_df, age97, pcged97, mratio96, pcg_adc)</pre>
summary(Matching::Match(
 Y = me_Y, Tr = me_Tr, X = me_X, Z = me_Z, M = 4,
 BiasAdjust = T, Var.calc = 4, estimand = "ATE", sample = T
))
##
## Estimate... -4.4867
## AI SE..... 1.7697
## T-stat.... -2.5353
## p.val..... 0.011235
##
## Original number of observations..... 606
## Original number of treated obs...... 188
## Matched number of observations..... 606
## Matched number of observations (unweighted). 2441
```

Check Balance

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

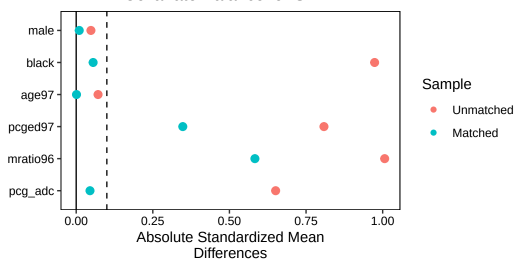
By default, the denominator for standardized mean differences uses a pooled estimate (square root of the average of the group variances) for ATE and the standard deviation of the treated group for ATT, and both standard deviations are computed using the sample before matching. This option can also be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

```
# Example of Checking Balance for SATE
me_SATE <- Match(
    Y = me_Y, Tr = me_Tr, X = me_X, M = 4, BiasAdjust = T, Var.calc = 4,
    estimand = "ATE", sample = T
)
(me_SATE_bal <- cobalt::bal.tab(
    me_SATE, kuse ~ male + black + age97 + pcged97 + mratio96 + pcg_adc,
    data = me_df,
    abs = T,
    un = T,
    binary = "std",
    thresholds = c(m = .1),
    s.d.denom = "pooled"
))</pre>
```

```
## Balance Measures
##
               Type Diff.Un Diff.Adj
                                            M.Threshold
## male
             Binary 0.0480
                              0.0099
                                         Balanced, <0.1
             Binary 0.9738
## black
                              0.0553
                                         Balanced, <0.1
## age97
            Contin.
                     0.0713
                              0.0010
                                         Balanced, <0.1
## pcged97 Contin. 0.8084
                              0.3478 Not Balanced, >0.1
                              0.5831 Not Balanced, >0.1
## mratio96 Contin. 1.0066
## pcg_adc Contin. 0.6508
                              0.0450
                                         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
             0.5831 Not Balanced, >0.1
##
## Sample sizes
##
                        Control Treated
## All
                         418.
                                 188.
## Matched (ESS)
                                  85.38
                         318.46
## Matched (Unweighted)
                         418.
                                 188.
```

```
cobalt::love.plot(me_SATE_bal, sample.names = c("Unmatched", "Matched")) +
labs(title = "Covariate Balance for SATE")
```

Covariate Balance for SATE



Outcome Analysis

```
# Recovering a matched dataset
me_out <- Match(</pre>
 Y = me_Y, Tr = me_Tr, X = me_X, M = 1, Var.calc = 4,
  estimand = "ATT", sample = T, replace = F
summary(me_out)
##
## Estimate... -7.4149
## SE..... 1.4634
## T-stat.... -5.0669
## p.val..... 0.0000040443
##
## Original number of observations.....
## Original number of treated obs.....
## Matched number of observations.....
## Matched number of observations (unweighted). 188
me_out_md <- rbind(me_df[me_out$index.treated,], me_df[me_out$index.control,])</pre>
```

Bonus: Nearest Neighbor Matching with the Matching Package

```
# Calculate propensity scores
me_glm <- glm(kuse ~ male + black + age97 + pcged97 + mratio96 + pcg_adc,</pre>
 family = binomial, data = me_df
# Nearest neighbor matching with MatchIt::matchit()
me_MatchIt <- MatchIt::matchit(</pre>
 kuse ~ male + black + age97 + pcged97 + mratio96 + pcg_adc,
 method = "nearest",
 distance = "glm",
 estimand = "ATT",
 data = me_df,
 m.order = "data",
 replace = F
)
# Nearest neighbor matching with Matching::Match()
me_Matching <- Matching::Match(</pre>
 Y = me_df$pcss97,
 Tr = me_df$kuse,
 X = me glmfitted,
 M = 1
 replace = F,
 estimand = "ATT",
 distance.tolerance = 0,
 ties = F,
  sample = F
# Verify that the results are identical
cobalt::bal.tab(me_MatchIt, weights = me_Matching)
## Balance Measures
               Type Diff.matchit Diff.Match
## distance Distance 0.9028 0.9028
## male Binary
                         -0.0213 -0.0213
           Binary
## black
## age97
## black
                        0.1755
                                    0.1755
            Contin.
                         0.0642
                                    0.0642
## pcged97 Contin.
                         -0.4191 -0.4191
## mratio96 Contin.
                         -0.5793
                                    -0.5793
## pcg_adc Contin.
                         0.3581
                                    0.3581
##
## Effective sample sizes
##
        Control Treated
         418 188
## All
## matchit 188 188
## Match 188 188
```

Appendix A: IMBALANCE Stata Module in R

```
imbalance <- function(df, varname, treatname, blockname) {</pre>
  \# dx
  df2 <- df %>%
    group_by({{ treatname }}) %>%
    summarise(
      m_x = mean(\{\{ varname \}\}),
      sd_x = sd(\{\{ varname \}\}),
      .groups = "drop"
    )
  mxt < -df2[2, 2]
  mxc \leftarrow df2[1, 2]
  s2xt <- df2[2, 3]^2
  s2xc \leftarrow df2[1, 3]^2
  sx \leftarrow sqrt((s2xt + s2xc) / 2)
  dx <- as.numeric(abs(mxt - mxc) / sx)</pre>
  \# dx
  df3 <- df %>%
    group_by({{ blockname }}, {{ treatname }}) %>%
    summarise(
      m x = mean(\{\{ varname \}\}),
      sd_x = sd(\{\{ varname \}\}),
      n = n(),
      .groups = "drop"
  mxc <- as.numeric(mean(filter(df3, {{ treatname }}} == 0)$m_x))</pre>
  mxt <- as.numeric(mean(filter(df3, {{ treatname }} == 1)$m_x))</pre>
  dxm_num <- abs(mxt - mxc)</pre>
  dxm <- as.numeric(dxm_num / sx)</pre>
  return(list(dx = dx, dxm = dxm))
}
```

Example Usage

```
imbalance(df, mratio96, kuse, fm)
```

Appendix B: HODGESL Stata Module in R

```
hodgesl <- function(dataname, varname, blockname, treatname) {</pre>
  blockname_str <- deparse(substitute(blockname))</pre>
  set.seed(1000)
  renamed_file <- dataname %>%
    filter(!is.na({{ blockname }}))
  r1 <- renamed_file %>%
    group_by({{ blockname }}) %>%
    summarise(m_y = mean({{ varname }}), .groups = "drop") %>%
    arrange({{ blockname }})
  r2 <- renamed file %>%
    group_by({{ blockname }}, {{ treatname }}) %>%
    summarise(
      mean_y = mean({{ varname }}),
     n = n(), .groups = "drop"
   ) %>%
   mutate(mean_diff = ifelse({{ treatname }} == 1,
      ((n + lag(n)) / sum(n)) * (mean_y - lag(mean_y)), NA
   )) %>%
   mutate(tx_effect = sum(mean_diff, na.rm = T)) %>%
   mutate(i = row_number()) %>%
    slice(1) %>%
    select(tx_effect, i)
  fm_results <- renamed_file %>%
    group_by({{ blockname }}, {{ treatname }}) %>%
    summarise(
      mean_y = mean({{ varname }}),
      n = n(), .groups = "drop"
  r3 <- renamed file %>%
    group_by({{ blockname }}, {{ treatname }}) %>%
    summarise(m_or_n = n(), .groups = "drop") %>%
    arrange({{ blockname }}, {{ treatname }}) %>%
   mutate(mi = ifelse({{ treatname }} == 0, m_or_n,
      ifelse({{ treatname }} == 1, NA, NA)
    mutate(ni = ifelse({{ treatname }} == 1, m_or_n,
      ifelse({{ treatname }} == 0, NA, NA)
   )) %>%
   mutate(Ni = ni + lag(mi)) %>%
   mutate(mi = ifelse(is.na(mi), lag(mi), mi)) %>%
   filter(!is.na(Ni)) %>%
   mutate(factor = (mi * ni) / (Ni * (Ni - 1))) %>%
   select({{ blockname }}, factor) %>%
    arrange({{ blockname }})
  r4 <- renamed file %>%
   arrange({{ blockname }}) %>%
   left_join(r1, by = blockname_str) %>%
   mutate(dy = {{ varname }} - m_y) %>%
   arrange(dy) %>%
   mutate(rk = row_number()) %>%
   arrange({{ blockname }})
```

```
r4a <- r4 %>%
   filter({{ treatname }} != 0) %>%
   group_by({{ blockname }}) %>%
    summarise(wsi = sum(rk), .groups = "drop")
 r5 <- r4 %>%
   group_by({{ blockname }}) %>%
    summarise(ki_ = mean(rk), .groups = "drop")
 r6 <- r4 %>%
   filter({{ treatname }} != 0) %>%
   group_by({{ blockname }}) %>%
    summarise(ni = n(), .groups = "drop") %>%
   arrange({{ blockname }}) %>%
   left_join(r5, by = blockname_str) %>%
   mutate(E_wsi = ni * ki_) %>%
   arrange({{ blockname }})
 r7 <- r4 %>%
    arrange({{ blockname }}) %>%
   left_join(r5, by = blockname_str) %>%
   mutate(k = (rk - ki_)^2) \%
    group_by({{ blockname }}) %>%
    summarise(ss_kd_i = sum(k), .groups = "drop") %>%
   arrange({{ blockname }})
  results <- r3 %>%
    arrange({{ blockname }}) %>%
   left_join(r7, by = blockname_str) %>%
   left_join(r6, by = blockname_str) %>%
   left_join(r4a, by = blockname_str) %>%
   mutate(
     var_wsi = factor * ss_kd_i,
     var = sum(var_wsi),
     sum_Ewsi = sum(E_wsi),
     ws = sum(wsi),
     HL_mean = ws - sum_Ewsi,
     HL_se = sqrt(var),
     z = HL_mean / HL_se,
      p = 1 - pnorm(abs(z))
   ) %>%
   select(HL_mean, HL_se, z, p) %>%
   slice(1) %>%
   mutate(i = row_number()) %>%
   left_join(r2, by = "i")
 return(results)
}
```

Session Info

sessionInfo()

```
## R version 4.5.1 (2025-06-13 ucrt)
## Platform: x86_64-w64-mingw32/x64
## Running under: Windows 11 x64 (build 26100)
##
## Matrix products: default
    LAPACK version 3.12.1
##
## locale:
## [1] LC_COLLATE=English_United States.utf8
## [2] LC_CTYPE=English_United States.utf8
## [3] LC_MONETARY=English_United States.utf8
## [4] LC NUMERIC=C
## [5] LC_TIME=English_United States.utf8
## time zone: Asia/Taipei
## tzcode source: internal
##
## attached base packages:
                 graphics grDevices utils
## [1] stats
                                                datasets methods
                                                                    base
## other attached packages:
## [1] Matching_4.10-15
                               MASS_7.3-65
                                                       WeightIt_1.5.0
## [4] lmtest_0.9-40
                               zoo_1.8-14
                                                       marginaleffects_0.30.0
## [7] rlang_1.1.6
                               sandwich 3.1-1
                                                       broom 1.0.10
## [10] knitr_1.50
                               optmatch_0.10.8
                                                       gbm_2.2.2
## [13] here_1.0.2
                               psych_2.5.6
                                                       survival_3.8-3
## [16] kableExtra_1.4.0
                               lubridate_1.9.4
                                                       forcats_1.0.0
## [19] stringr_1.5.2
                               dplyr_1.1.4
                                                       purrr_1.1.0
## [22] readr 2.1.5
                               tidyr 1.3.1
                                                       tibble 3.3.0
                                                       MatchIt_4.7.2
## [25] ggplot2_4.0.0
                               tidyverse_2.0.0
## [28] cobalt_4.6.1
                               sjlabelled_1.2.0
                                                       haven_2.5.5
## [31] formatR_1.14
                               tictoc_1.2.1
                                                       pacman_0.5.1
##
## loaded via a namespace (and not attached):
## [1] tidyselect_1.2.1
                           viridisLite_0.4.2
                                               farver_2.1.2
                                                                  S7_0.2.0
## [5] fastmap_1.2.0
                           digest_0.6.37
                                               timechange_0.3.0
                                                                  lifecycle_1.0.4
## [9] magrittr_2.0.4
                           compiler_4.5.1
                                               tools_4.5.1
                                                                  utf8_1.2.6
## [13] yaml_2.3.10
                           data.table_1.17.8 labeling_0.4.3
                                                                  mnormt_2.1.1
## [17] xml2_1.4.0
                           RColorBrewer_1.1-3 withr_3.0.2
                                                                  grid_4.5.1
## [21] scales_1.4.0
                           tinytex_0.57
                                               insight_1.4.2
                                                                  cli_3.6.5
## [25] rmarkdown 2.29
                           crayon_1.5.3
                                               generics 0.1.4
                                                                  rstudioapi_0.17.1
## [29] tzdb_0.5.0
                                               splines_4.5.1
                           rlemon_0.2.1
                                                                  parallel_4.5.1
## [33] vctrs_0.6.5
                           Matrix_1.7-3
                                               hms_1.1.3
                                                                  systemfonts_1.2.3
## [37] glue_1.8.0
                           chk_0.10.0
                                               stringi_1.8.7
                                                                  gtable_0.3.6
## [41] pillar_1.11.1
                           htmltools_0.5.8.1 R6_2.6.1
                                                                  textshaping_1.0.3
## [45] rprojroot_2.1.1
                           evaluate_1.0.5
                                               lattice_0.22-7
                                                                  backports 1.5.0
## [49] Rcpp_1.1.0
                           svglite_2.2.1
                                               nlme_3.1-168
                                                                  checkmate_2.3.3
## [53] xfun_0.53
                           pkgconfig_2.0.3
```