Section 8

Matching and Weighting Estimators

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Overview

- Logistics:
 - Pset 7 released! Due at 11:59 pm (ET) on Nov 10
- Today's topics:
 - 1. Matching estimators
 - 2. Weighting estimators

[Review] Observational studies

Identification

- Most common observational assumptions:
 - No unmeasured confounders: $D_i \perp (Y_i(0), Y_i(1)) \mid \mathbf{X}_i$
 - Overlap/positivity: $0 < \mathbb{P}(D_i = 1 \mid \mathbf{X}_i = x) < 1$
- Estimand:
 - ATE = $\mathbb{E}[Y_i(1) Y_i(0)]$ (We identified this in Module 5)
 - ATT = $\mathbb{E}[Y_i(1) Y_i(0) \mid D_i = 1]$
 - ATC = $\mathbb{E}[Y_i(1) Y_i(0) \mid D_i = 0]$

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Estimation

- Regression estimators: $\widehat{\mu}_1(\mathbf{x})$ and $\widehat{\mu}_0(\mathbf{x})$ (Module 5)
- Matching estimator (for ATT):

$$\widehat{\tau}_m = \frac{1}{n_1} \sum_{i=1}^n D_i \left(Y_i - \frac{1}{M} \sum_{j \in \mathcal{J}_M(i)} Y_j \right)$$

- Weighting estimators:
 - Horvitz-Thompson estimator (= IPW estimator)
 - Hajek estimator (normalized weights)

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- Mahalanobis distance matching: use distance metrics in case of high dimensional X;
- Propensity score matching:

$$\left(Y_i(0),Y_i(1)\right) \perp D_i \mid \mathbf{X}_i \Rightarrow \left(Y_i(0),Y_i(1)\right) \perp D_i \mid \pi(\mathbf{X}_i)$$

- This holds under **true** propensity score $\pi(\mathbf{X}_i)$.
- We need to estimate it $(\widehat{\pi}(\mathbf{X}_i))$: e.g., using logistic regression (can add interactions) or machine learning.
- Have to check if \mathbf{X}_i is actually balanced.

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 - Optimal matching:
 - **D**: $n \times n$ matrix of pairwise distance or a cost matrix
 - Select n elements of D such that there is only one element in each row and one element in each column and the sum of pairwise distances is minimized → linear sum assingment problem

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- Assessing balance
 - standardized mean differences
 - Kolmogorov–Smirnov statistic (comparing distributions)

Matching estimators

- Workflow (in general):
 - 1. Check the balance before the matching
 - Choose matching type (compute/estimate the distance/balancing score if necessary)
 - 3. Conduct matching (check the matched dataset)
 - 4. Check the balance after the matching
 - 5. Estimate ATT using matching estimator $(\widehat{\tau}_m)$
 - 6. Estimate the standard error
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- Useful packages:
 - cobalt: for balance check (bal.tab() and love.plot())
 - MatchIt: for matching
 - Matching: for matching + estimating
 - Machine learning packages for estimating $\widehat{\pi}(\mathbf{X}_i)$: e.g., randomForest
 - Optimal matching: clue::solve_LSAP()

Example: LaLonde dataset

- The effectiveness of a job training program (National Supported Work Demonstration; NSW) on wage increases.
- The federal government instituted a randomized evaluation of this program
- How well the result may be recovered when the experimental controls are replaced with a set of observational controls (Population Survey of Income Dynamics; PSID)?
- **Problem**: Imbalances between the experimental and observational data → use matching

Example: LaLonde dataset

Data:

Treated: 185 units from NSW
 Control: 2490 units from PSID

Treatment: Participation in the job training program (nsw)

Outcome: 1978 earnings (in dollars; re78)

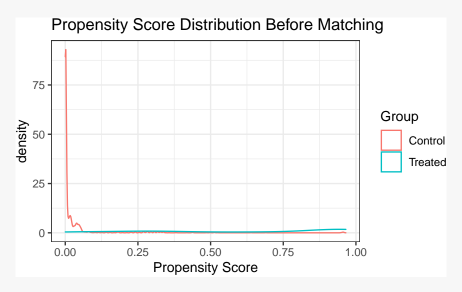
 Pre-treatment covariates: age, race, marriage, past earnings, past employment

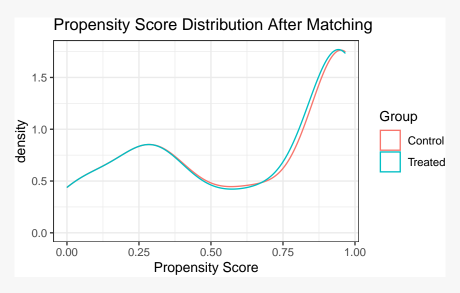
Example: Balance before matching

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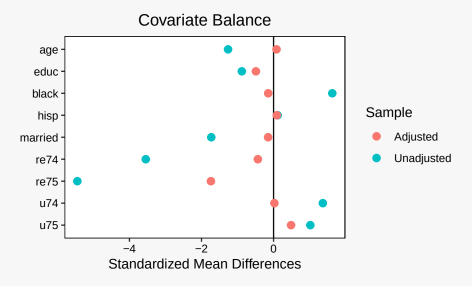
```
## Balance Measures
##
             Type Diff.Un
          Contin. -1.0094
## age
## educ Contin. -0.6805
## black Binary 1.4816
## hisp Binary 0.1288
## married Binary -1.8453
## re74
          Contin. -1.7178
## re75
          Contin. -1.7744
## u74
           Binary 1.6454
## u75
           Binary 1.2309
##
## Sample sizes
      Control Treated
##
## All
         2490
                  185
```

```
##
## Estimate... -941.15
## SE...... 903.91
## T-stat.... -1.0412
## p.val.... 0.29779
##
## Original number of observations...... 2675
## Original number of treated obs...... 185
## Matched number of observations (unweighted). 185
```





```
library(cobalt)
love.plot(nsw ~ age+educ+black+hisp+married+re74+re75+u74+u75,
  data = dat,
  stats = "mean.diffs",
  weights = data.frame(Matched = get.w(match_ps)),
  method = c("matching"), binary = "std")
```



Weighting estimators

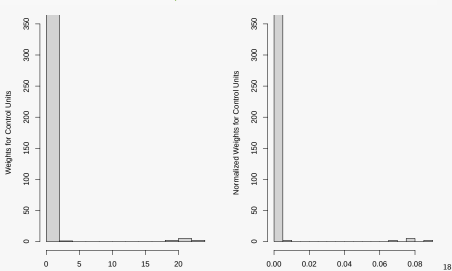
- Matching is actually a special case of a weighting estimator
- Horvitz-Thompson estimator: weight by inverse propensity score.

$$\widehat{\mathsf{ATE}} = \widehat{\tau}_{ipw} = \frac{1}{n} \sum_{i=1}^{n} \left(\frac{D_i Y_i}{\widehat{\pi}(\mathbf{X}_i)} - \frac{(1 - D_i) Y_i}{1 - \widehat{\pi}(\mathbf{X}_i)} \right)$$

- Would be unbiased if we knew the true propensity scores, $\pi(\mathbf{X}_i)$ (Pset 7 Q2 Bonus)
- Under no unmeasured confounders, $\widehat{\tau}_{ipw} \stackrel{p}{\to} \tau$ (consistent)
- Hajek estimator: normalizes the weights
- Potential of extreme weights due to lack of overlap: $\pi(\mathbf{X}_i)$ close to 0 or 1
 - Windsorizing: trim weights beyond 5th and 95th percentile

Example

Generating propensity score weights for the ATT



Estimating ATT with Weights

```
## append estimated weights
dat <- dat %>% mutate(weights = W.out$weights)
## ATT
att_ipw <- function(dat, indices = NULL) {</pre>
  if (is.null(indices)) indices <- 1:nrow(dat)</pre>
  dat <- dat[indices,]</pre>
  weights <- dat %>% filter(treat == 0) %>% pull(weights)
  reweights <- weights / sum(weights)</pre>
  Y1 <- dat %>% filter(treat == 1) %>% pull(re78)
  Y0 <- dat %>% filter(treat == 0) %>% pull(re78)
  att_ht <- sum(Y1 - Y0 * weights) / nobs
  att_hjk <- mean(Y1) - sum(reweights * Y0)
  return(c(att_ht, att_hjk))
}
## Use bootsrap for estimating SE
```