EC 425/525, Set 7

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Prologue

Schedule

Last time

- ullet The conditional independence assumption: $(\mathbf{Y}_{0i},\,\mathbf{Y}_{1i}) \perp \!\!\! \perp \mathbf{D}_i | \mathbf{X}_i$
- Omitted variable bias
- Good vs. bad controls

Today

- Return first round of project proposals.
- Matching estimators (MHE 3.2 and Cameron and Trivedi 25.4).

Upcoming

- Admin: Assignment and midterm
- Next round of the project proposal

Follow up

OLS weighting

At the beginning of the lecture, we discussed OLS weights—especially for heterogeneous treatment effects.

We should keep our questions clear.

- 1. Which weights on β_1 and β_2 recover β_{12} , where β_i comes from a regression using observations in group i?
- 2. What does β represent when the treatment effect is heterogeneous?

More soon.

The gist

Remember the **conditional independence assumption**[†] in a setting—*i.e.*, treatment is as-good-as random conditional on a known set of covariates?

Matching estimators take us at our word.

If we really believe $(Y_{1i}, Y_{0i}) \perp D_i | X_i$, then we can just calculate a bunch of treatment effects conditional on X_i , *i.e.*,

$$\tau(x) = E[\mathbf{Y}_{1i} - \mathbf{Y}_{0i} \mid \mathbf{X}_i = x]$$

The idea: Estimate a treatment effect only using observations with (nearly?) identical values of X_i . The CIA buys us causality within these groups.

Goals

Let's return to the fundamental problem of causal inference for a moment.

- 1. We want/need to know $au_i = Y_{1i} Y_{0i}$.
- 2. We cannot simultaneously observe both Y_{1i} and Y_{0i} .

Most empirical strategies boil to strategies to estimate \mathbf{Y}_{0i} for treated individuals—the unobservable counterfactual for the treatment group.

Matching is no different.

We match untreated observations to treated observations using X_i , i.e., calculate a \widehat{Y}_{0i} for each Y_{1i} , based upon "matched" untreated individuals.

More formally

We want to construct a counterfactual for each individual with $D_i = 1$.

The counterfactual for i should only use individuals that match X_i .

Let there be N_T treated individuals and N_C control individuals. We want

- N_T sets of weights
- ullet with N_C weights in each set : $w_i(j)$ $(i=1,\,\ldots,\,N_T;\,j=1,\,\ldots,\,N_C)$

Assume $\sum_j w_i(j) = 1$. Our estimate for the counterfactual of treated i is

$$\widehat{\mathrm{Y}_{0i}} = \sum_{j \in (D=0)} w_i(j) \mathrm{Y}_j$$

More formally

If our estimated counterfactual for treated individual i is

$$\widehat{\mathrm{Y}_{0i}} = \sum_{j} w_i(j) \mathrm{Y}_j$$

then our estimated treatment effect (for individual i) is

$$\hat{m{ au}}_i = \mathrm{Y}_{1i} - \widehat{\mathrm{Y}_{0i}} = \mathrm{Y}_{1i} - \sum_j w_i(j) \mathrm{Y}_j$$

... a generic matching estimator for the treatment effect on the treated is

$$\hat{ au}_M = rac{1}{N_T} \sum_{i \in (\mathrm{D}=1)} \left(\mathrm{Y}_{1i} - \widehat{\mathrm{Y}_{0i}}
ight) = rac{1}{N_T} \sum_{i \in (\mathrm{D}=1)} \left(\mathrm{Y}_{1i} - \sum_{j \in (D=0)} w_i(j) \mathrm{Y}_j
ight)$$

Weight for it[†]

So all we need is those weights and we're done. **

Q Where does one find these handy weights?

A You've got options, but you need to choose carefully/responsibly.

E.g., if $w_i(j)=\frac{1}{N_C}$ for all (i,j), then we're back to a difference in means. This weighting doesn't abide by our conditional independence assumption.

The plan Choose weights $w_i(j)$ that indicate **how close** X_j is to X_i .

† 👤 †† Plus an interesting, policy-relevant setting with valid conditional independence. And data.

Proximity

Our weights $w_i(j)$ should be a measure of **how close** X_i is to X_i .

If X is **discrete**, then we can consider equality, *i.e.*, $w_i(j) = \mathbb{I}(X_i = X_j)$, scaling as necessary to get $\sum_j w_i(j) = 1$.

Proximity

Our weights $w_i(j)$ should be a measure of **how close** X_j is to X_i .

If X is **continuous**, then we need *proximity* rather than *equality*.

Nearest-neighbor matching chooses the single closest control observation using the Euclidean distance between X_i and X_j , i.e.,

$$\mathrm{d}_{i,j} = \left(\mathrm{X}_i - \mathrm{X}_j
ight)'\left(\mathrm{X}_i - \mathrm{X}_j
ight)$$

- $\hat{ au}_i = \mathrm{Y}_{1i} \mathrm{Y}_{0j}^i$, where Y_{0j}^i is i's nearest neighbor in the control group.
- Estimator: $\hat{ au}_M = rac{1}{N_T} \sum_i \hat{ au}_i$
- Produces causal estimates if CIA is valid and we have sufficient overlap.
- Suffers from arbitrary choices of units.

Proximity

Our weights $w_i(j)$ should be a measure of **how close** X_j is to X_i .

If X is **continuous**, then we need *proximity* rather than *equality*.

Nearest-neighbor matching with Mahalanobis distance chooses the single closest control using Mahalanobis distance between X_i and X_j , i.e.,

$$\mathrm{d}_{i,j} = \left(\mathrm{X}_i - \mathrm{X}_j
ight)' \Sigma_X^{-1} \left(\mathrm{X}_i - \mathrm{X}_j
ight)$$

where Σ_X^{-1} is the covariance matrix of X.

- **Estimator:** $\hat{ au}_M = rac{1}{N_T} \sum_i \hat{ au}_i$ where $\left(\hat{ au}_i = \mathrm{Y}_{1i} \mathrm{Y}_{0j}^i
 ight)$
- Produces causal estimates if CIA is valid and we have sufficient overlap.
- Does not suffer from arbitrary choices of units.

More neighbors?

Why limit ourselves to a **single** "best" match?

If we're going to let a function/algorithm choose the *nearest* match, can't we also let the function/algorithm choose *how many* matches?

Furthermore, if $N_C \gg N_T$, it we're throwing away *a lot* of information.

We could instead use this information and be more efficient.

More neighbors!

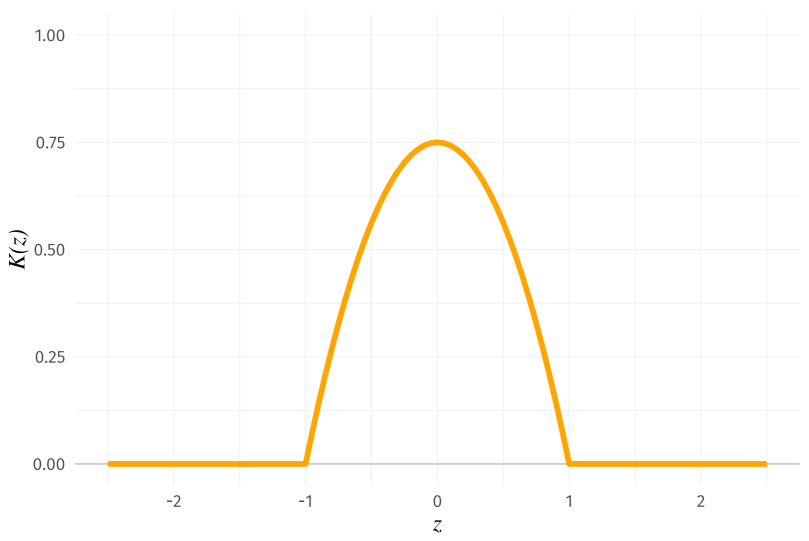
Kernel matching gives positive weight to all control observations within some **bandwidth** h, with higher weight for closer matches determined by some **kernel function** $K(\cdot)$,

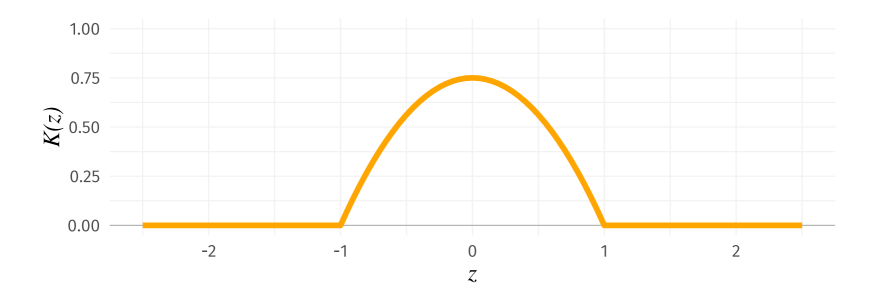
$$w_i(j) = rac{\mathit{K}\!\!\left(rac{\mathrm{X}_j - \mathrm{X}_i}{h}
ight)}{\sum_{j \in (D=0)} \mathit{K}\!\!\left(rac{\mathrm{X}_j - \mathrm{X}_i}{h}
ight)}$$

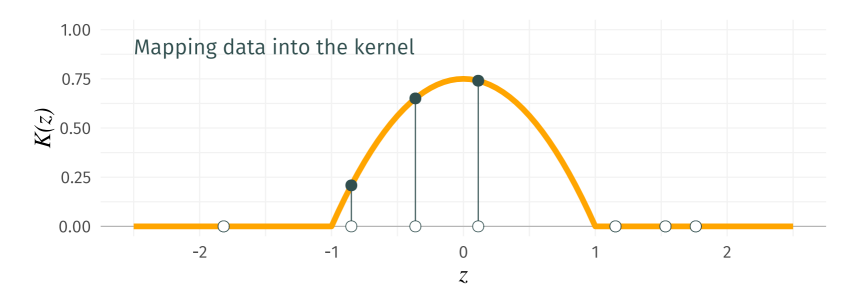
Example The Epanechnikov kernel is defined as

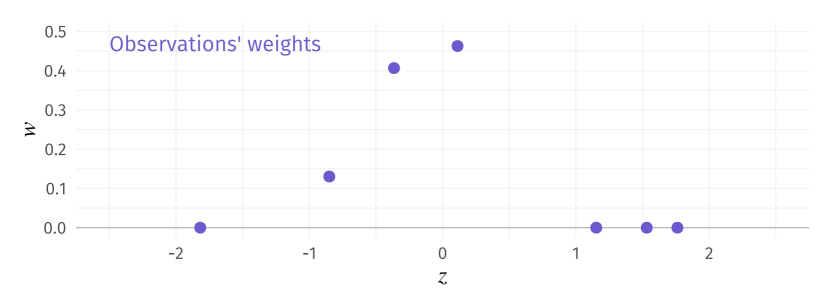
$$K(z) = rac{3}{4}ig(1-z^2ig) imes \mathbb{I}(|z|<1)$$

The Epanechnikov kernel $K(z)=rac{3}{4}ig(1-z^2ig) imes \mathbb{I}(|z|<1)$

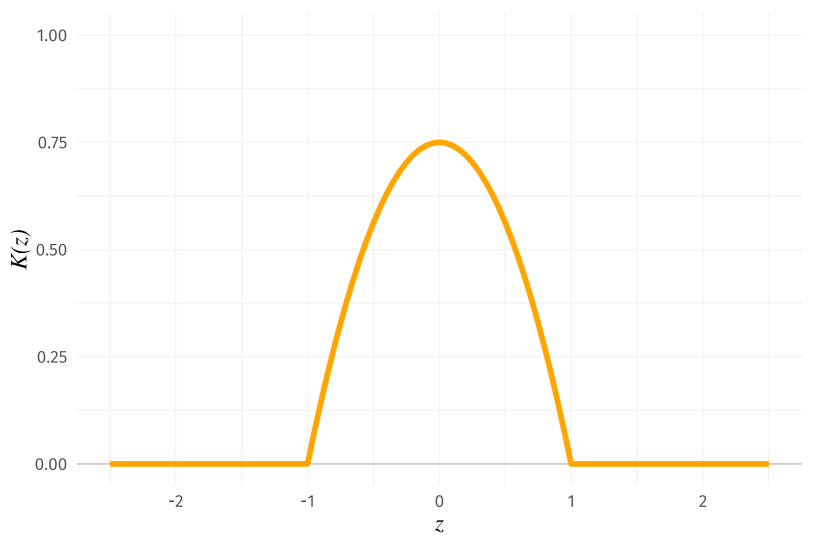




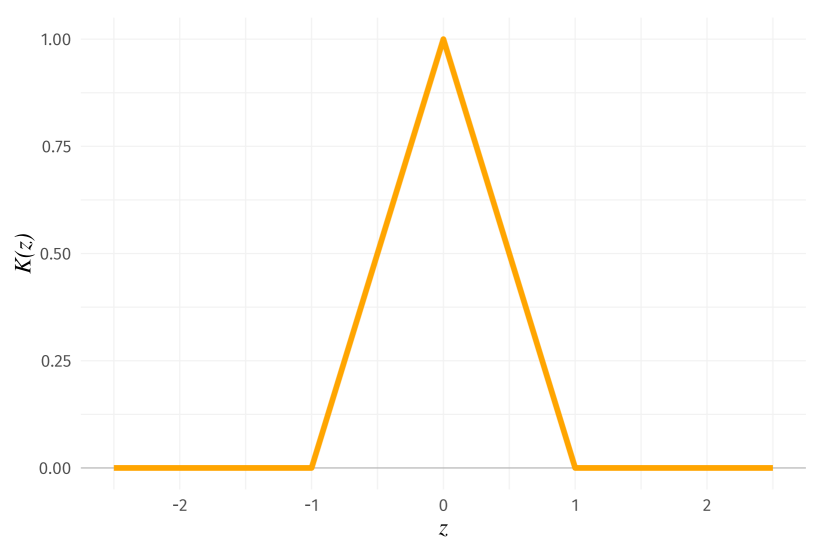




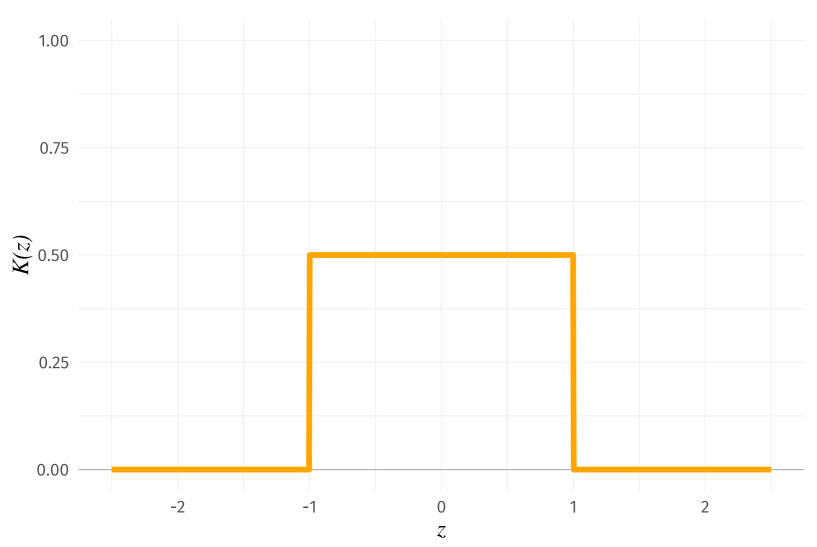
The Epanechnikov kernel $K(z)=rac{3}{4}ig(1-z^2ig) imes \mathbb{I}(|z|<1)$



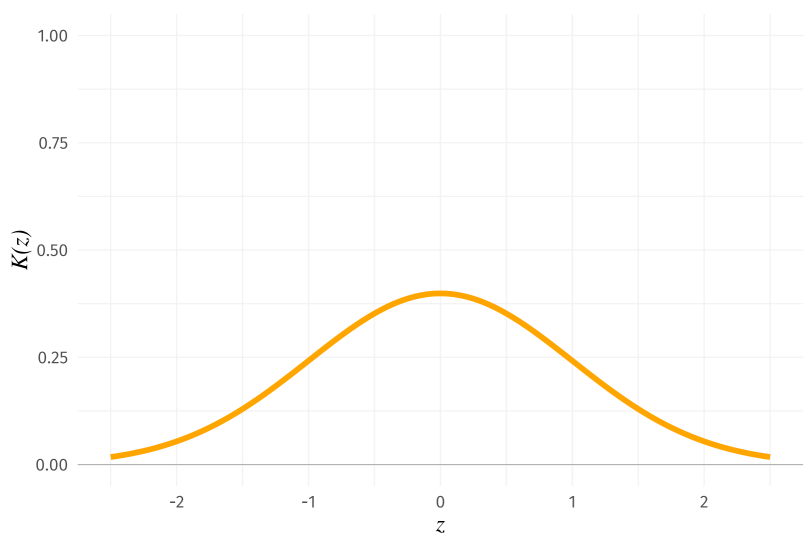
The Triangle kernel $K(z) = (1-|z|) imes \mathbb{I}(|z| < 1)$



The Uniform kernel $K(z) = rac{1}{2} imes \mathbb{I}(|z| < 1)$



The Gaussian kernel $K(z) = \left(2\pi ight)^{-1/2} \exp\left(-z^2/2 ight)$



Kernels

Aside

Kernel functions are good for more than just matching.

You will most commonly see/use them smoothing out densities—providing a smooth, moving-window average.

E.g., R's (ggplot2's) smooth, density-plotting function geom_density().

geom_density() defaults to kernel = "gaussian", but you can specify many
other kernel functions (including "epanechnikov").

You can also change the bandwidth argument. The default is a bandwidth-choosing function called bw.nrd0().

Adding neighbors

As we add more neighbors—either moving from 1 to n>1 or increasing our bandwidth—we potentially increase the efficiency of our estimator.

We need to **be careful not to add too many controls** for each treated i.

CIA requires that we're actually conditioning on the observables—it does not allow us to take a simple average across all control observations.

The curse of dimensionality[†]

It turns out kernel- and bandwidth-selection are not our biggest enemies.

As the dimension of X expands (matching on more variables), it becomes harder and harder to find a nice, close control for each treated unit.

We need a way to shrink the dimensionality of X.

Setup

Let's begin with two assumptions—one old and one new.

- 1. Conditional independence: $(Y_{0i}, Y_{1i}) \perp \!\!\! \perp D_i | X_i$
- 2. **Overlap:** $0 < \Pr(D_i = 1 \mid X_i) < 1$

We can estimate an average treatment effect by conditioning on X_i .

However, overlap may fail if the dimensions of X are large and N is finite.

Propensity scores provide a solution to this mess.

The magic

It turns out that if $(Y_{0i}, Y_{1i}) \perp D_i | X_i$, then we actually only need to match/condition on $p(X_i) = E[D_i | X_i]$.

 $p(X_i)$ is the **propensity score**, the probability of treatment given X_i .

Propensity-score theorem If $(Y_{0i}, Y_{1i}) \perp \!\!\! \perp D_i | X_i$, then $(Y_{0i}, Y_{1i}) \perp \!\!\! \perp D_i | p(X_i)$.

This theorem extends our CIA to a one-dimensional score, avoiding the curse of dimensionality.

Theorem If $(Y_{0i}, Y_{1i}) \perp D_i | X_i$, then $(Y_{0i}, Y_{1i}) \perp D_i | p(X_i)$.

Proof

To prove this theorem, we will show $\Pr(D_i = 1 \mid Y_{0i}, Y_{1i}, p(X_i)) = p(X_i)$, i.e., D_i is independent of (Y_{0i}, Y_{1i}) after conditioning on $p(X_i)$.

Theorem If $(Y_{0i}, Y_{1i}) \perp \!\!\! \perp D_i | X_i$, then $(Y_{0i}, Y_{1i}) \perp \!\!\! \perp D_i | p(X_i)$.

Proof

$$egin{aligned} & \Priggl[\mathrm{D}_i = 1 \Big| \mathrm{Y}_{0i}, \, \mathrm{Y}_{1i}, \, p(\mathrm{X}_i) iggr] \ & = Eiggl[\mathrm{D}_i \Big| \mathrm{Y}_{0i}, \, \mathrm{Y}_{1i}, \, p(\mathrm{X}_i) iggr] \ & = Eiggl[Eiggl(\mathrm{D}_i \Big| \mathrm{Y}_{0i}, \, \mathrm{Y}_{1i}, \, p(\mathrm{X}_i), \, \mathrm{X}_i iggr) \Big| \mathrm{Y}_{0i}, \, \mathrm{Y}_{1i}, \, p(\mathrm{X}_i) iggr] \ & = Eiggl[Eiggl(\mathrm{D}_i \Big| \mathrm{Y}_{0i}, \, \mathrm{Y}_{1i}, \, \mathrm{X}_i iggr) \Big| \mathrm{Y}_{0i}, \, \mathrm{Y}_{1i}, \, p(\mathrm{X}_i) iggr] \end{aligned}$$

Theorem If $(Y_{0i}, Y_{1i}) \perp D_i | X_i$, then $(Y_{0i}, Y_{1i}) \perp D_i | p(X_i)$.

Proof

$$\begin{split} \Pr \bigg[\mathrm{D}_i &= 1 \Big| \mathrm{Y}_{0i}, \, \mathrm{Y}_{1i}, \, p(\mathrm{X}_i) \bigg] = \dots = E \bigg[E \bigg(\mathrm{D}_i \Big| \mathrm{Y}_{0i}, \, \mathrm{Y}_{1i}, \, \mathrm{X}_i \bigg) \Big| \mathrm{Y}_{0i}, \, \mathrm{Y}_{1i}, \, p(\mathrm{X}_i) \bigg] \\ &= E \bigg[E \bigg(\mathrm{D}_i \Big| \mathrm{X}_i \bigg) \Big| \mathrm{Y}_{0i}, \, \mathrm{Y}_{1i}, \, p(\mathrm{X}_i) \bigg] \\ &= E \bigg[p(\mathrm{X}_i) \Big| \mathrm{Y}_{0i}, \, \mathrm{Y}_{1i}, \, p(\mathrm{X}_i) \bigg] \\ &= p(\mathrm{X}_i) \\ & \therefore \, (\mathrm{Y}_{0i}, \, \mathrm{Y}_{1i}) \perp \!\!\! \perp \mathrm{D}_i | \mathrm{X}_i \implies (\mathrm{Y}_{0i}, \, \mathrm{Y}_{1i}) \perp \!\!\! \perp \mathrm{D}_i | p(\mathrm{X}_i) \quad \checkmark \end{split}$$

Intuition

Q What's going on here?

 X_i carries way more information than $p(X_i)$, so how can we still get conditional independence of treatment by only conditioning on $p(X_i)$?

 A_1 Conditional independence of treatment isn't about extracting all of the information possible from X_i . We actually only care about creating a situation in which D_i something is independent of (Y_{0i}, Y_{1i}) .

 A_2 Back to our main concern: **selection bias**. People select into treatment. If X says two people were equally likely to be treated, and if X_i explains all of selection (CIA), then there cannot be selection between these two people.

Estimation

So where do propensity scores come from?

We estimate them—and there are a lot of ways to do that.

- 1. Flexible (i.e., interactions) logit specification
- 2. Kernel regression (remember kernel functions?)
- 3. Many others—machine learning, series-logit estimator, etc.
- Q Can we just use plain OLS (linear probability model)?
- A Sort of. Think about FWL. This route is going to be the same as a regression conditioning on X_i .

Estimation

From MHE (p. 83)

Question

A big question here is how to best model and estimate $p(X_i)$...

Answer

The answer to this is inherently application-specific. A growing empirical literature suggests that a logit model for the propensity score with a few polynomial terms in continuous covariates works well in practice...

Application

So you have some estimated propensity scores $\hat{p}(X_i)$. What next?

Option 1 Conditioning via regression

Option 1a Use a **regression to condition** on $p(X_i)$, i.e.,

$$Y_i = \alpha + \delta D_i + \beta p(X_i) + u_i$$
 (1a)

Option 1b If we think treatment effects are heterogeneous and may covary with X, then we might want to also **interact** treatment with $p(X_i)$, i.e.,

$$Y_i = \alpha + \delta_1 D_i + \delta_2 D_i p(X_i) + \beta p(X_i) + u_i$$
 (1b)

Heterogeneity with regression

Let's think a bit more about heterogeneous treatment effects in this setting.

$$\mathbf{Y}_{0i} = \alpha + \beta \mathbf{X}_i + u_i$$

 $\mathbf{Y}_{1i} = \mathbf{Y}_{0i} + \delta_1 + \delta_2 \mathbf{X}_i$

i.e., the treatment effect depends upon X_i .

$$egin{aligned} \mathbf{Y}_i &= \mathbf{D}_i \mathbf{Y}_{1i} + \left(1 - \mathbf{D}_i
ight) \mathbf{Y}_{0i} \ &= \mathbf{D}_i igg(\mathbf{Y}_{0i} + \delta_1 + \delta_2 \mathbf{X}_iigg) + \left(1 - \mathbf{D}_i
ight) \mathbf{Y}_{0i} \ &= \mathbf{Y}_{0i} + \delta_1 \mathbf{D}_i + \delta_2 \mathbf{D}_i \mathbf{X}_i \ &= lpha + \delta_1 \mathbf{D}_i + \delta_2 \mathbf{D}_i \mathbf{X}_i + eta \mathbf{X}_i + u_i \end{aligned}$$

Heterogeneity

This final equation

$$\mathbf{Y}_i = \alpha + \delta_1 \mathbf{D}_i + \delta_2 \mathbf{D}_i \mathbf{X}_i + \beta \mathbf{X}_i + u_i$$

suggests that we want $p(X_i)$ and $D_i p(X_i)$, i.e.,

$$Y_i = \alpha + \delta_1 D_i + \delta_2 D_i p(X_i) + \beta p(X_i) + u_i$$
 (1b)

which yields

- 1. a **group-specific treatment effect** $\delta_1 + \delta_2 \mathbf{X}_i$ for each \mathbf{X}_i
- 2. an **average treatment effect** $\delta_1 + \delta_2 \overline{p}(\mathbf{X}_i)$

More flexibility

We motivated propensity scores with a desire to reduce dimensionality and estimate/choose/assume fewer parameters.

Adding $p(X_i)$ and $D_i p(X_i)$ as covariates in a linear regression doesn't quite exhaust our potential for flexible/nonparametric estimation.

Blocking

Option 2 Block (stratify) on propensity scores.

- 1. Divide the range of $\hat{p}(\mathbf{X}_i)$ into K blocks (e.g., 0.05-wide blocks).
- 2. Place each observation into a block via its $\hat{p}(\mathbf{X}_i)$.
- 3. Calculate $\hat{\tau}_k$ for each block via difference in means.
- 4. Average the $\hat{\tau}_k$ using their shares of the sample, i.e.,

$$\hat{ au}_{ ext{Block}} = \sum_{k=1}^K \hat{ au}_k rac{N_{1k} + N_{0k}}{N}$$

Note Blocking is similar to NN/kernel matching using $p(X_i)$ as distance.

Choosing blocks

Blocking on propensity scores requires defining defining blocks.

One common route involves some iteration.

- 1. Choose blocks.
- 2. Check the **balance of the covariates** within each block.[†]
 - If covariates are not balanced, then split your blocks and repeat.
 - If covariates are balanced, then stop.

[†] Keep multiple-hypothesis testing in mind. With many covariates and many blocks, you are bound to find statistically significant relationships—even if you are balanced in truth.

Overlap

Blocking emphasizes our overlap assumption, i.e., $0 < \Pr(D_i|X_i) < 1$.

If a block contains zero treated/control units, we cannot calculate $\hat{\tau}_k$.

Caution Logit can hide violations—it forces $0 < \hat{p}(X_i) < 1$.

Common practice Empirically enforce overlap:

- Drop control units with $\hat{p}(\mathbf{X}_i)$ below the minimum propensity score in the treatment group.
- Drop treated units with $\hat{p}(\mathbf{X}_i)$ above the maximum propensity score in the control group.

Weighting

Option 3 Weight observations by the inverse propensity score.

Q How does weighting by $1/\hat{p}(\mathbf{X}_i)$ make sense?

A Consider our old (likely biased) friend the difference in means, i.e.,

$$\hat{ au}_{ ext{Diff}} = \overline{ ext{Y}}_{ ext{T}} - \overline{ ext{Y}}_{ ext{C}} = rac{\sum_{i} ext{D}_{i} ext{Y}_{i}}{\sum_{i} ext{D}_{i}} - rac{\sum_{i} \left(1 - ext{D}_{i}
ight) ext{Y}_{i}}{\sum_{i} \left(1 - ext{D}_{i}
ight)}$$

which we've discussed is biased due to selection into treatment, i.e.,

$$E[\mathrm{Y}_{0i}|\mathrm{D}_i=1]
eq E[\mathrm{Y}_{0i}]$$

Weighting, justified

Suppose we know $p(\mathrm{X}_i)$ and we weight each **treated** individual by $1/p(\mathrm{X}_i)$

$$Eigg[rac{\mathrm{D}_i\mathrm{Y}_i}{p(\mathrm{X}_i)}igg]$$

$$= Eigg[rac{\mathrm{D}_i\left(\mathrm{D}_i\mathrm{Y}_{1i} + (1-\mathrm{D}_i)\mathrm{Y}_{0i}
ight)}{p(\mathrm{X}_i)}igg]$$

$$= Eigg[rac{\mathrm{D}_i \mathrm{Y}_{1i}}{p(\mathrm{X}_i)}igg]$$

$$= Eigg(Eigg[rac{\mathrm{D}_i \mathrm{Y}_{1i}}{p(\mathrm{X}_i)} \ igg| \mathrm{X}_i igg] igg)$$

$$\int E[D_i \mid X_i] E[Y_{1i} \mid X_i] \setminus$$

Weighting: The estimator

Thus, we can estimate an unbiased treatment effect via

$$\hat{ au}_{p ext{Weight}} = rac{1}{N} \sum_{i=1}^{N} \left[rac{ ext{D}_{i} ext{Y}_{i}}{p(ext{X}_{i})} - rac{(1- ext{D}_{i} ext{Y}_{i})}{1-p(ext{X}_{i})}
ight]$$

Intuition We're trying to overcome selection bias, i.e., treated individuals were more likely to be treated as a function of X_i —producing higher $p(X_i)$.

We want to get back to as-good-as random variation in treatment.

So we upweight (1) **treated** individuals with low $p(X_i)$ and (2) **control** observations with high $p(X_i)$.

Weighting: The example

Suppose for some individual i, $p(X_i) = 0.80$.

This propensity score says someone with this set of X_i was four-times more likely to be **treated** than **control**.

Our weights fix this imbalance for each X_i .

- If i is **treated**, then her weight is $1/p(\mathrm{X}_i) = 1/0.80 = 1.25$
- If i is **control**, then her weight is $1/(1-p(\mathbf{X}_i))=1/(1-0.80)=5$

And guess what 5/1.25 is...

4!

This weighting scheme gets us back to equal representation for each set $q_{3/52}^{
m f}$

Weighting: Last issue

Practical issue Nothing guarantees $\sum_i \hat{p}(\mathbf{X}_i) = 1$.

Solution Normalize weights by their total sum.

Applying the normalized (and estimated) propensity scores

$$\hat{ au}_{p ext{Weight}} = \sum_{i=1}^N rac{rac{\mathrm{D}_i \mathrm{Y}_i}{\hat{p}(\mathrm{X}_i)}}{\sum_i rac{\mathrm{D}_i}{\hat{p}(\mathrm{X}_i)}} - \sum_{i=1}^N rac{rac{(1-\mathrm{D}_i)\mathrm{Y}_i}{1-\hat{p}(\mathrm{X}_i)}}{\sum_i rac{(1-\mathrm{D}_i)}{1-\hat{p}(\mathrm{X}_i)}}$$

Hirano, Imbens, and Ridder (2003) suggests this estimator is efficient.

Why choose one?

There's nothing special about weighted averages—regression can weight.

Thus, a regression-based estimate

$$\mathbf{Y}_i = \alpha + \mathbf{X}_i \boldsymbol{\beta} + \tau \mathbf{D}_i + u_i$$

with weights

$$w_i = \sqrt{rac{\mathrm{D}_i}{\hat{p}(\mathrm{X}_i)} + rac{(1-\mathrm{D}_i)}{1-\hat{p}(\mathrm{X}_i)}}$$

offers a doubly robust property—you have two chances to be right: $p(X_i)$ or the regression specification.

Why choose one? Part two

An alternative, doubly robust method combines propensity-score blocking with regression.

Step 1 For each block k, we run the regression

$$\mathbf{Y}_i = lpha_k + \mathbf{X}_i eta_k + au_k \mathbf{D}_i + u_i$$

Step 2 Aggregate block-level treatment-effect estimates

$$\hat{ au} = \sum_{k=1}^K \hat{ au}_k rac{N_{1k} + N_{0k}}{N}$$

Major requirements

Don't get (too) caught up in the bells and whistles.

We still have two **major** requirements for any of these methods to work?

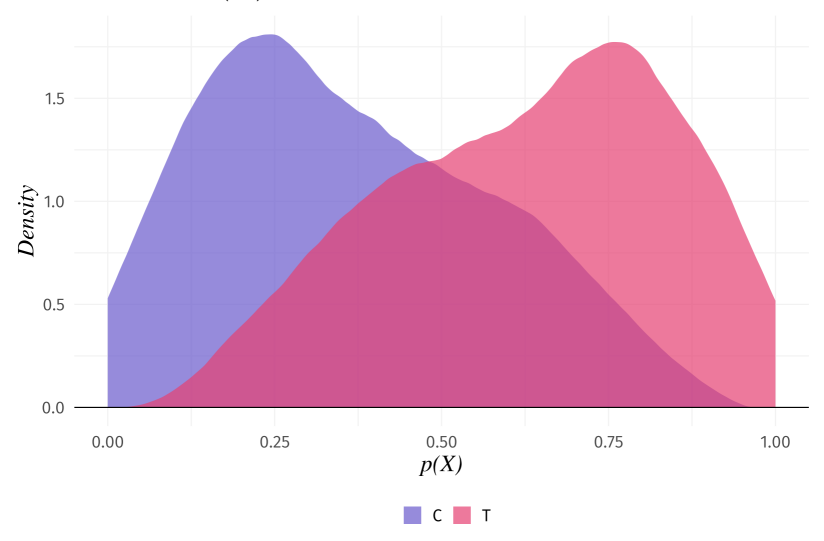
- 1. Is the conditional-independence assumption true?
- 1. Do we have **overlap** between treatment and control units.

We can look for evidence of (2) in the data—particularly if we're using propensity-score methods.[†]

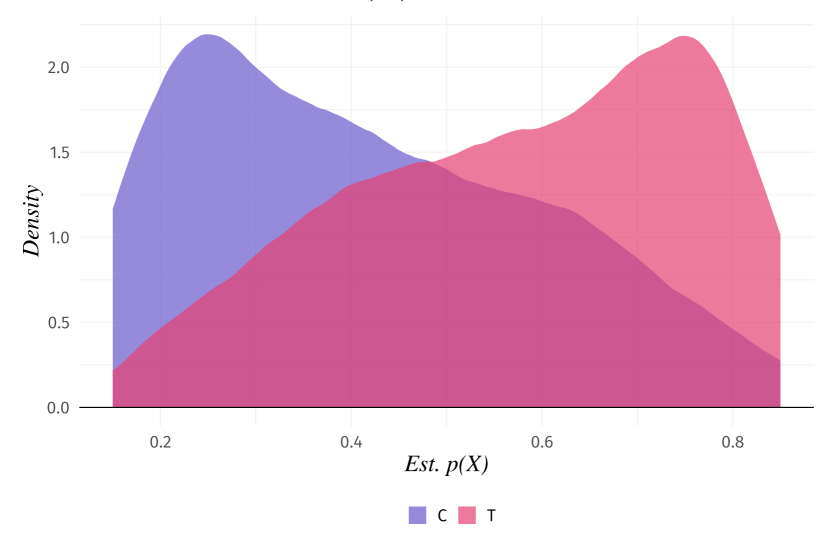
How? Plot the distributions of $p(X_i)$ for **T** and **C**.

 $[\]dagger$ Checking for overlap in X-space, can be tough as the dimensions of X expand.

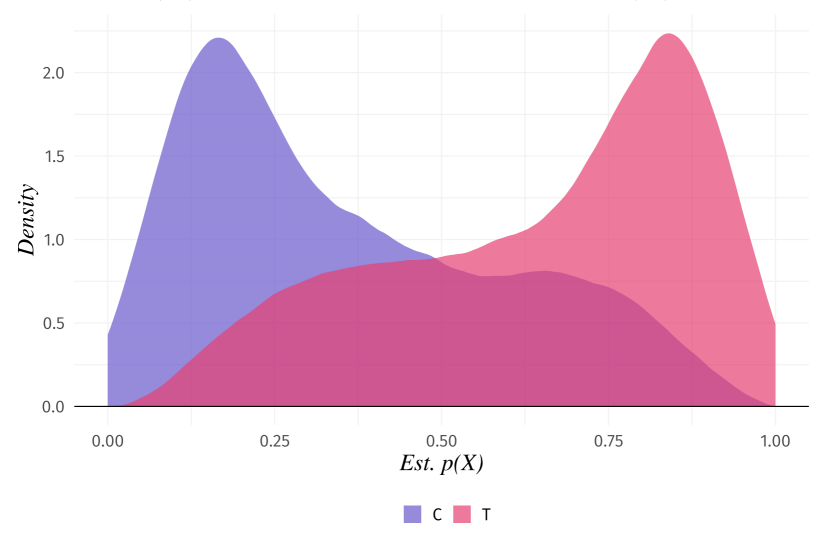
Missing overlap in $p(\mathbf{X}_i)$



Authentic (enforced) overlap in $p(X_i)$



Logit-based $\hat{p}(\mathbf{X}_i)$ hiding some of the missing overlap in $p(\mathbf{X}_i)$



Overlap in one dimension does not guarantee in two dimensions.

Note Shading denotes **share of treatment:** white =0% and **pink**=100%.



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