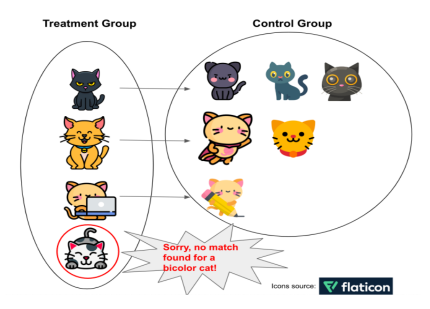
Causal Inference: Matching POLI 803 Research Methods in PS

Howard Liu 2025

The Idea of Matching



Estimators

- Now we will explore estimators that for lack of a better word "use covariates" to estimate aggregate causal parameters
- A few topics: subclassification, exact matching, inexact/approximate matching

Assumptions

- When attempting to estimate aggregate causal parameters using *covariate* adjustment, there are two assumptions
 - 1. **Unconfoundedness** (no ommitted X_i): fairly controversial to some because of what it implies about human behavior
 - 2. **Common support** (strong overlaps in X_i): since most people stop at assumption 1, there's much less attention to common support
- Today we will assume you satisfy unconfoundedness, and therefore our focus is on violations of assumption 2

Roadmap

1. Subclassification

- 2. Exact Matching
- Inexact Matching
 Nearest Neighbor Covariate Matching
 Propensity scores
 Coarsened Exact Matching (CEM)

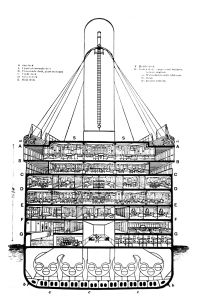
Subclassification method

 Subclassification: an early effort to tackle covariates, but which ultimately cannot handle large features due to common support issues

Subclassification method

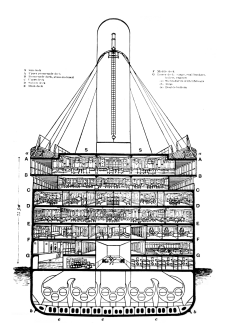
 Subclassification: an early effort to tackle covariates, but which ultimately cannot handle large features due to common support issues → no one uses it. shown for pedagogical purpose

The Titanic example



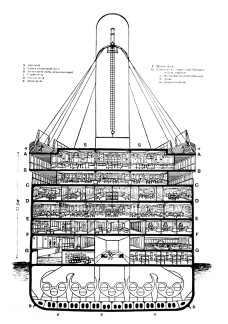
• Titanic sank: 2200 on board, but only 700 survived

The Titanic example



First Class: Decks B-C

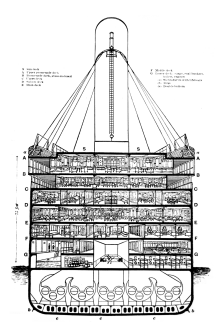
Titanic layout



Second Class: Decks D-E



Titanic layout



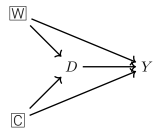
Third Class: Decks F and G

The Titanic example

- Women and children first was a maritime rule to ration lifeboats, but there were different cabins (1st class, 2nd class, etc.) on different levels with different proximity to boats
- Q: What was the causal effect of 1st class on survival (after adjusting for W and C)?
- Can you draw a DAG (Directed Ayclic graph)?

Exercise: Titanic DAG

Figure: Women W and children C first maritime rule is a confounder for estimating first class D effect on surviving Y



Backdoor criterion can be satisfied by blocking on W and C. These are our known confounders. Now we just need data to see if it's quantified.

 Stratify the confounders: Our age and sex variables are both binary, so we can only create four strata: male children, female children, male adults, female adults → to "control" the covariates

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- Calculate probability weights: Count the number of people in each strata and divide by the total number of souls aboard (crew and passengers)
- 4. Aggregate differences across strata using weights: Estimate the ATE by aggregating the difference in survival rates over the four strata with each strata-specific difference weighted by that strata's weight

Table: Counts and Titanic survival rates by strata and first class status.

	First class		All other classes			
Strata	Obs	Mean	Obs	Mean	Total	
Male adult	175	0.326	1,492	0.188	1,667	
Female adult	144	0.972	281	0.626	425	
Male child	5	1	59	0.407	64	
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Strata	Differences in Survival Rates	$Weight_{k,ATE}$	$Weight_{k,ATT}$	$Weight_{k,ATU}$
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Female adult	0.346	0.19	0.44	0.15
Male child	0.593	0.03	0.02	0.03
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	No stratification	Stratif	fication weighted esti	imates ÂTU
Estimated coefficient	0.35	0.20	0.24	0.19

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Table 2: Estimates of aggregate parameters

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What if we drop the only female child in the 1st class (due to false data)?

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Table: Differences in survival rates, stratification weights, and estimates of parameters without perfect stratification

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Lack of Common Support (Empty cells)

- Stratification requires having units in both groups for every value of \boldsymbol{X} to get ATE
- If you want the ATT, you have to have units in the control group for every treated group based on its value of X (female children weren't treated after n/a, so didn't matter)

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- This has a technical word we are going to learn more about called a "lack of common support" (missings obs)

No overlap (common support violation)



No support is like an incomplete bridge which stops you from even being able to cross the moat even though the troops exist

The Curse of Dimensionality

 Dimensionality (k) usually means number of covariates. Think about our Titanic example?

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- Stratification methods break down in finite samples because as increase the number of covariates, the "dimension" grows even faster
- Assume we have k covariates and we divide each into 3 coarse categories (e.g., age: young, middle age, old; income: low, medium, high, etc.)
- The number of strata is 3^k . For k = 10, then it's $3^{10} = 59,049$
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- The curse of dimensionality is based on the slices of all *interactions* of the covariates, not just the covariates, and that explodes fast
 - \rightarrow we need a better method to create strata

Roadmap

1. Subclassification

2. Exact Matching

Inexact Matching
 Nearest Neighbor Covariate Matching
 Propensity scores
 Coarsened Exact Matching (CEM)

Exact matching



Exact Matching

 Subclassification: uses the difference between treatment and control group units and achieves covariate balance by using the k probability weights to weight the averages

Exact Matching

- Subclassification: uses the difference between treatment and control group units and achieves covariate balance by using the k probability weights to weight the averages
 - \rightarrow failed quickly when k goes large and many missing matches (common support failed)
- Exact Matching: Ultimately use N nearest neighbor to address the problem of infinite matches

Exact matching

- Exact matching finds a person in the control group whose value of X_j is exactly equal to each person in the treatment group i
- Will not work if the conditioning set includes a continuous variable
- Will also not work if K gets large (curse of dimensionality we discuss later)

The Training Example: Training → Earnings

• Take average with multiple exact matches $(\frac{1}{M})$

Table 29. Training example with exact matching (including matched sample).

			•					
Unit	Trainees t <mark>Age</mark> Earnings		Non-Trainees Unit Age Earnings		Matched Sample Unit Age Earnings			
1	18	9500	1	20	8500	14	18	8050
2	29	12250	2	27	10075	6	29	10525
3	24	11000	3	21	8725	9	24	9400
4	27	11750	4	39	12775	8	27	10075
5	33	13250	5	38	12550	11	33	11425
6	22	10500	6	29	10525	13	22	8950
7	19	9750	7	39	12775	17	19	8275
8	20	10000	8	33	11425	1	20	8500
9	21	10250	9	24	9400	3	21	8725
10	30	12500	10	30	10750	10,18	30	9875
			11	33	11425			
			12	36	12100			
			13	22	8950			
			14	18	8050			
			15	43	13675			
			16	39	12775			
			17	19	8275			
			18	30	9000			
			19	51	15475			
			20	48	14800			
Mean	24.3	\$11,075		31.95	\$11,101.25		24.3	\$9,380

The Training example: Age and Earnings

• How do we create matches/counterfactuals with the treated group?

Table 28. Training example with exact matching.

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1	18	9500	1	20	8500
2	29	12250	2	27	10075
3	24	11000	3	21	8725
4	27	11750	4	39	12775
5	33	13250	5	38	12550
6	22	10500	6	29	10525
7	19	9750	7	39	12775
8	20	10000	8	33	11425
9	21	10250	9	24	9400
10	30	12500	10	30	10750
			11	33	11425
			12	36	12100
			13	22	8950
			14	18	8050
			15	43	13675
			16	39	12775
			17	19	8275
			18	30	9000
			19	51	15475
			20	48	14800
Mean	24.3	\$11,075		31.95	\$11,101.25

ATT estimator: One match

We will focus on the ATT for the rest of today and the equation is:

$$\widehat{\delta}_{ATT} = \frac{1}{N_T} \sum_{D_i=1} (Y_i - Y_{j(i)}) \tag{1}$$

where $Y_{j(i)}$ is the j^{th} unit matched to the i^{th} unit based on the j^{th} being "exactly equal to" the i^{th} unit with respect to the X conditioning set

Multiple matches

Multiple matches: What if I find two or more M units with the identical X value? Then what?

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$$\widehat{\delta}_{ATT} = \frac{1}{N_T} \sum_{D_i = 1} \left(Y_i - \left[\frac{1}{M} \sum_{m=1}^M Y_{j_m(1)} \right] \right) \tag{2}$$

Notice that we are only dealing with Y_i^0 (control) by matching; The Y_i^1 (treatment) is fine as is.

1. For each unit i in the treatment group with known and quantified confounder $X=x_i$, **find** all units j in the donor pool for whom $x_i=x_j$. These j units are our M matches and M can be one or it can be greater than one if you want it to be.

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- 3. For each unit i, calculate the difference between realized earnings and matched earnings, $\widehat{\delta}_i = Y_i \frac{1}{M} \sum Y_{j(i)}$.
- 4. Finally, estimate the sample ATT by averaging over all i differences in earnings from Step 3 as $\frac{1}{N_T}\sum \widehat{\delta}_i$, where N_T is the number of treatment units.
- Easy peasy!

• But what problem we would encounter here?

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- What if there are no exact matches?

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Inexact matching



Nearest Neighbor Covariate Matching (NN Matching)

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 What if you couldn't find another unit with that exact same value of X? → Then you do *approximate* matching

Nearest Neighbor Covariate Matching (NN Matching)

- What if you couldn't find another unit with that exact same value of X? → Then you do *approximate* matching
- Estimate $\widehat{\delta}_{ATT}$ by **imputing** the missing potential outcome of each treatment unit i using the observed outcome from that outcome's "nearest" neighbor j in the control set using X for the matching

$$\widehat{\delta}_{ATT} = \frac{1}{N_T} \sum_{D_i=1} (Y_i - Y_{j(i)})$$

where $Y_{j(i)}$ is the observed outcome of a control unit such that $X_{j(i)}$ is the **closest** value to X_i among all of the control observations (eg match on X)

Matching

 We could also use the average observed outcome over M closest matches:

$$\widehat{\delta}_{ATT} = \frac{1}{N_T} \sum_{D_i = 1} \left(Y_i - \left[\frac{1}{M} \sum_{m=1}^{M} Y_{j_m(1)} \right] \right)$$

• Works well when we can find good matches for each treatment group unit, so M is usually defined to be small (i.e., M=1 or M=2)

Example: Matching example with single covariate

i	Y_i^1	\mathbf{V}^0	D_i	X_i
$\overline{}^{\iota}$		Y_i^0	D_i	
1	6	?	1	3
2	1	?	1	1
3	0	?	1	10
4		0	0	2
4 5		9	0	3
6		1	0	-2
7		1	0	-4

Question: What is
$$\hat{\delta}_{ATT} = \frac{1}{N_T} \sum_{D_i=1} (Y_i - Y_{j(i)})$$
?

Example: Matching example with single covariate

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2	1	?	1	1
3	0	?	1	10
4		0	0	2
4 5 6		9	0	3
6		1	0	-2
7		1	0	-4

Question: What is
$$\hat{\delta}_{ATT} = \frac{1}{N_T} \sum_{D_i=1} (Y_i - Y_{j(i)})$$
?

Match and plug in!

Match on the X and plug in Y!

Matching example with single covariate

i	Y_i^1	Y_i^0	D_I	X_i
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ATT?

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ATT?

Question: What is
$$\widehat{\delta_{ATT}} = \frac{1}{N_T} \sum_{D_i=1} (Y_i - Y_{j(i)})$$
?

$$\hat{\delta}_{ATT} = \frac{1}{3} \cdot (6-9) + \frac{1}{3} \cdot (1-0) + \frac{1}{3} \cdot (0-9) = -3.7$$

Measuring the matching discrepancy

- What does it mean to be close when I am working with a large number of covariates k > 1?
- Need a way of measuring a match in terms of how "close" each unit's X_i value was to the matched X_i

Measuring the matching discrepancy

- What does it mean to be close when I am working with a large number of covariates k > 1?
- Need a way of measuring a match in terms of how "close" each unit's X_i value was to the matched X_i
 - \to Let's do that and use the square root of the sum of all squared differences in each unit's $X_i-X_{j(i)}$ as a measure of how bad the match is
 - → This is called the Euclidean distance

Euclidean distance

Definition: Euclidean distance

$$||X_i - X_j|| = \sqrt{(X_i - X_j)'(X_i - X_j)}$$

= $\sqrt{\sum_{n=1}^k (X_{ni} - X_{nj})^2}$

Minimizing the Euclidean distance

- Abadie and Imbens (2006) show that there exists a unique solution to the matching problem that minimizes a given distance metric
- Matching in R and teffects in Stata (not sure in python)
- But the idea here is that any other match will always have a higher Euclidean distance

Other distance metrics

- Our example treated a one unit difference in age and one unit difference in GPA as the same, but those scales are different and matter a lot
- The Euclidean distance is *not* invariant to changes in the scale of the X's.
- Alternative distance metrics that are invariant to changes in scale are more commonly used
- Normalized Euclidean distance and Mahalanobis distance both try to normalize it so that scale doesn't matter

Normalized Euclidean distance

Definition: Normalized Euclidean distance

A commonly used distance is the normalized Euclidean distance:

$$||X_i - X_j|| = \sqrt{(X_i - X_j)'\hat{V}^{-1}(X_i - X_j)}$$

where

$$\widehat{V}^{-1} = \operatorname{diag}(\widehat{\sigma}_1^2, \widehat{\sigma}_2^2, \dots, \widehat{\sigma}_k^2)$$

Normalized Euclidean distance

Notice that the normalized Euclidean distance is equal to:

$$||X_i - X_j|| = \sqrt{\sum_{n=1}^k \frac{(X_{ni} - X_{nj})^2}{\widehat{\sigma}_n^2}}$$

• Thus, if there are changes in the scale of X_{ni} , these changes also affect $\widehat{\sigma}_{n}^{2}$, and the normalized Euclidean distance does not change

Mahalanobis distance

Definition: Mahalanobis distance

The Mahalanobis distance is the **scale-invariant** distance metric:

$$||X_i - X_j|| = \sqrt{(X_i - X_j)'\widehat{\Sigma}_X^{-1}(X_i - X_j)}$$

where $\widehat{\Sigma}_X$ is the sample variance-covariance matrix of X.

Matching and the Curse of Dimensionality

Common support and the dimensality curse?

Matching and the Curse of Dimensionality

- Common support and the dimensality curse? The larger the dimensions of the conditioning set, the less likely common support holds, and you can't do it because you need these covariate dimensions to satisfy weak unconfoundedness!
- This problem is caused by the finite dataset, and it introduces a particular type of selection bias
- Curses are only overcome with new spells!

Matching and the Curse of Dimensionality

- Common support and the dimensality curse? The larger the dimensions of the conditioning set, the less likely common support holds, and you can't do it because you need these covariate dimensions to satisfy weak unconfoundedness!
- This problem is caused by the finite dataset, and it introduces a particular type of selection bias
- Curses are only overcome with new spells! → Abadie and Imbens (2011) derived a way to reduce the bias (bias adjustment or bias correction) → in the matchIt package

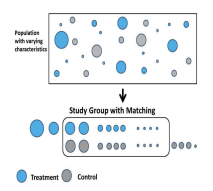
Curse of dimensionality, bias and heterogeneous treatment effects

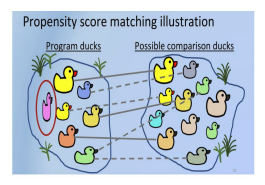
- Recall the problem of many covariates for exact matching the curse of dimensionality makes matching on K covariates implausible as the dimensions grow exponentially with K
- This is problem because recall there are two assumptions needed to match
 - 1. Unconfoundedness: this gives you the right to match
 - 2. Common support: this gives you the ability to match
- Without both, then depending on the amount of hetergeneity in the treatment effects, matching will be biased

Curse of dimensionality, bias and heterogeneous treatment effects

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 - \rightarrow We introduce a different technique to address this curse of dimension

Propensity score as dimension reduction





Propensity score as dimension reduction

- Rubin (1977) and Rosenbaum and Rubin (1983) developed the propensity score method
- Propensity score theorum: They show that if treatment is independent of potential outcomes conditional on K covariates, then it will be independent of potential outcomes conditional on propensity score

Propensity score as dimension reduction

- Rubin (1977) and Rosenbaum and Rubin (1983) developed the propensity score method
- Propensity score theorum: They show that if treatment is independent of potential outcomes conditional on K covariates, then it will be independent of potential outcomes conditional on propensity score
- Main value of the propensity score is dimension reduction to reduce K covariates into a single scalar without loss of information
- Variety of ways to incorporate the propensity score stratification, weighting and matching. We focus on propensity score matching here.

Caveat: **Propensity score** matching

- King and Nielson (2019) "Why propensity scores should not be used for matching"
- "The more balanced the data (or the more balanced it comes by trimming some of the observations through matching), the more likely propensity score matching will degrade inferences."

Caveat: **Propensity score** matching

- King and Nielson (2019) "Why propensity scores should not be used for matching"
- "The more balanced the data (or the more balanced it comes by trimming some of the observations through matching), the more likely propensity score matching will degrade inferences."
- → caution against assuming that more balance always improves causal inference
- Reduced sample size
- Bias from exclusion of extreme values
- Overfitting to the matched sample
- The author thinks its fine. But I see less work on p-score matching these days

Propensity score matching: three core steps

1. Takes necessary covariates

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- 2. Estimates a maximum likelihood model of the conditional probability of treatment (usually a logit or probit so as to ensure that the fitted values are bounded between 0 and 1),

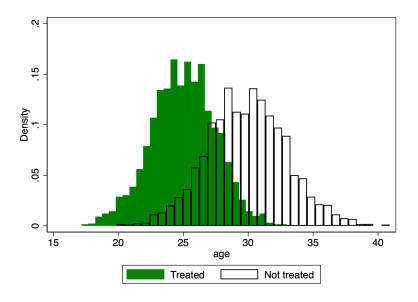
Propensity score matching: three core steps

- 1. Takes necessary covariates
- 2. Estimates a maximum likelihood model of the conditional probability of treatment (usually a logit or probit so as to ensure that the fitted values are bounded between 0 and 1),
- Uses the predicted values from that estimation to collapse those covariates into a single scalar called the propensity score. All comparisons between the treatment and control group are then based on that value (propensity).

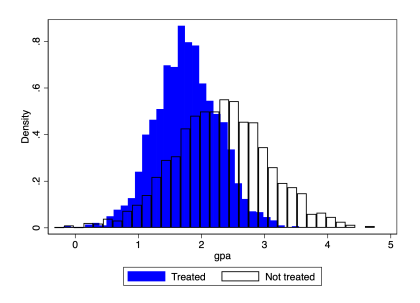
Investigating overlap (common support)

- Once you obtain the propensity score, you can use it for estimation, but you can also use it for evaluating covariate balance
- It's an easy way to do it with histograms, and since the propensity score theorem holds for the dimensions of X, there's no loss of generality in investigating overlap that way versus one by one
- Remember: you need common support, not on X individual covariates alone, but within the K dimensions, so investigating with the propensity score is easier to do

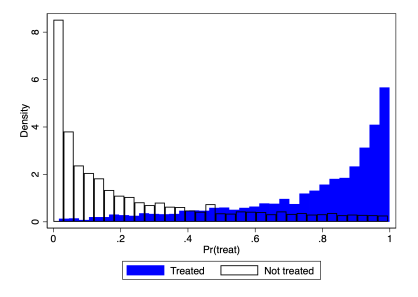
Covariate 1 histograms: Age



Covariate 2 histograms: GPA



Summarizing both with propensity score histogram



Navigating the vastness of estimation

- Estimators abound and can be a little bewildering so to summarize them:
 - 1. Match units from one group to another **using the propensity score** (with various rules for finding how close to be)
 - 2. Weighting by the inverse propensity score
- Variety of techniques to derive standard errors from parametric methods to bootstrapping
- Can even introduce "doubly robust" methods to deal with matching bias like we did with nearest neighbor bias correction
- But all these estimators assume unconfoundedness, so really it's all about addressing the lack of overlap; that's the only bias that exists when you have unconfoundedness

Step 1: Pick your parameter ATE vs ATT vs ATU

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- Which population are you studying? Only those who were discriminated against? That's the ATT
- Do you want to imagine "what if blacks and whites were both discriminated against?" That's the ATE
- The more you can focus on one particular causal parameter, the easier and more justified it gets as it weakens both assumptions (unconfoundedness and common support)

Step 2: Estimate the propensity score

 Estimate the conditional probability of treatment using probit or logit model (or ML)

$$Pr(D_i = 1|X_i) = F(\beta X_i)$$

- Note: don't use OLS because while it will get the mean right, it will
 not get correct values in the tails because of its linear projections
- OLS will give propensity scores outside the [0,1] bounds and probabilities cannot be negative or greater then one

Step 2: Estimate the propensity score

 Use the estimated coefficients to predict the propensity score for each unit i

$$\widehat{\rho}_i(X_i) = \widehat{\beta}X_i$$

 Note that each unit i now has a predicted probability of treatment given the values of their covariates relative to everyone else's

Step 2: Estimate the propensity score

- Think of the propensity score as a frequentist concept of probability
- "If I drew someone from the sample with these characteristics, then how many of those are in the treatment group divided by the total with those characteristics"
- Or for each dimension of X, a ratio of $\frac{N_T}{(N_T+N_C)}$

Step 3a: Propensity Score Estimation with matching

- Most common method is to use matching (propensity score matching or matching on propensity scores)
- Matching finds a unit in the comparison group with a similar $\widehat{\rho}_i(X)$ to service as counterfactual for the unit

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- · Lack of overlap creates issues for matching which we'll note later

Step 3b: Propensity Score Estimation with stratification

- Rare to see this done anymore, though it was one of the methods that Dehejia and Wahba (2002) tried
- Stratification is a kind of weighting method similar to Cochran's subclassification method where weights are group shares within certain ranges of the propensity score
- Weighting on the propensity score: most common is the inverse probability weighting (IPW)

Step 3c: Estimation with inverse probability weighting

• Estimate the propensity score:

$$E[X_i] = Pr(D_i = 1|X_i)$$

- Inverse: Assign weights based on how "rare" each observation's treatment is:
- For treated:

$$w = \frac{1}{E[X_i]}$$

For control:

$$w = \frac{1}{1 - E[X_i]}$$

Step 3c: Estimation with inverse probability weighting

Intuition:

- Units who are unlikely to receive treatment (low E[X])) but did are given more weight, since they provide rare, informative comparisons.
- Units who were very likely to receive treatment (high E[X])) are given less weight, since they are common and less informative.

Step 3c: Estimation with inverse probability weighting

- IPW uses the estimated propensity score to reweight the outcomes (making treated and control groups more comparable)
- IPW is non-parametric you are just taking averages and multiplying by the inverse of the propensity score weights depending on which parameter you want to estimate
- There are fewer implementation choices than in matching (i.e., no choice over distance, number of neighbors)
- There are bias adjustment methods called double robust where you combine imputing counterfactuals with weighting by the propensity score

Step 3d: Coarsened Exact Matching

- Also called weighting on the propensity score
- lacus et al. (2012) introducted it
- Very simple idea: it's possible to do exact matching once we coarsen the data enough.
- E.g., 0- to 10-year-olds, 11- to 20-year olds, then oftentimes we can find exact matches

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 automatic binning algorithm. For instance, schooling becomes less
 than high school, high school only, some college, college graduate,
 post college
- 3. Then we create one stratum per unique observation of X* and place each observation in a stratum. Assign these strata to the original and uncoarsened data, X, and drop any observation whose stratum doesn't contain at least one treated and control unit.
- 4. Then add weights for stratum size and analyze without matching (similar to what we did in the subclassfication example)

Step 4: Standard Errors

Standard errors can be constructed a few different ways:

- We need to adjust the standard errors for first-step estimation of $\rho(X)$
 - → Parameteric first step: Newey and McFadden (1994)
 - → Non-parametric first step: Newey (1994)

Reminder: Check for common support assumption using histograms

- Assessing whether there are units it both groups for whichever parameter you're focused on is simple with propensity score as shown earlier using histograms of the propensity score for treated and control → little overlaps mean bad matching
- Crump, et al. (2009) suggest keeping propensity scores within the interval [0.1,0.9] ("trimming") but any trimming will drop units and dropping units means moving away from the parameter
- Let's look at a picture again just to remind ourselves

Assessing overlap

