## MATCHING TECHNIQUES

## Technical Track Session VI

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The World Bank

## When can we use matching?

What if the assignment to the treatment is not done randomly or based on an eligibility index, but on the basis of observable variables?

This is when matching methods come in! Matching methods allow you to construct comparison groups when the assignment to the treatment (or the take-up of treatment) can be modeled on the basis of observable variables.

## When can we use matching?

Intuition: the comparison group needs to be as similar as possible to the treatment group, in terms of the observables before the start of the treatment. Avoids possible "confounders" from imbalance in observables.

- The method assumes there are no 'remaining' unobservable differences between treatment and comparison groups.


## Key Question

What is the effect of treatment when the assignment to the treatment is based on observable variables?

# Unconfoundedness \& Selection on observables 

- Let $X$ denote a matrix in which each row is a vector of pre-treatment observable variables for individual $i$.
- Unconfoundedness: Assignment to treatment is unconfounded given pre-treatment variables $X$ if

$$
Y_{1}, Y_{0} \perp D \mid X
$$

Unconfoundedness is equivalent to saying that: (1) within each cell defined by $X$ : treatment is random (2) the selection into treatment depends only on the observables $X$.

## Average effects of treatment Assuming unconfoundedness given $X$

- Intuition
- Estimate the treatment effect within each cell defined by $X$
- Take the weighted average over the different cells
- Maths

In your handouts: Annex 1

## Strategy for estimating average effect of treatment <br> Selection on observables

- Unconfoundedness suggests the following strategy for the estimation of the average treatment effect $\delta$
- Stratify the data into cells defined by each particular value of $X$
- Within each cell (i.e. conditioning on $X$ ) compute the difference between the average outcomes of the treated and the controls
- Average these differences with respect to the distribution of $X$ in the population of treated units.

Is this strategy feasible?

# Is our strategy feasible? The Dimensionality Problem 

- This may not be feasible when
- The sample is small
- The set of covariates is large
- Many of the covariates have many values or are continuous
- This is what we call...

The dimensionality problem

## The Dimensionality Problem

- Examples
- How many cells do we have with 2 binary $X$ variables? And with 3 binary $X$ variables? And with K binary $X$ variables?
- How about if we have 2 variables that take on 7 values each?
- As the number of cells grows, we'll get lack of common support
- cells containing only treated observations
- cells containing only controls


## An Alternative to solve the Dimensionality Problem

The propensity score allows to convert the multidimensional setup of matching into a onedimensional setup.
In that way, it allows to reduce the dimensionality problem.

## Rosenbaum and Rubin

Rosenbaum and Rubin (1983) propose an equivalent and feasible estimation strategy based on the concept of Propensity Score.

## Matching based on the Propensity Score

## Definition

The propensity score is the conditional probability of receiving the treatment given the pre-treatment variables:

$$
p(X)=\operatorname{Pr}\{D=1 \mid X\}=E_{X}\{D \mid X\}
$$

## Lemma 1

If $p(X)$ is the propensity score, then $D \perp X \mid p(X)$
"Given the propensity score, the pre-treatment variables are balanced between beneficiaries and non- beneficiaries"

## Lemma 2

$Y 1, Y 0 \perp D|X \quad=>Y 1, Y 0 \perp D| p(X)$
"Suppose that assignment to treatment is unconfounded given the pre-treatment variables $X$. Then assignment to treatment is unconfounded given the propensity score $p(X)$."

# Does the propensity score approach solve the dimensionality problem? 

## YES!

- The balancing property of the propensity score (Lemma 1) ensures that:
- Observations with the same propensity score have the same distribution of observable covariates independently of treatment status; and
- for a given propensity score, assignment to treatment is "random" and therefore treatment and control units are observationally identical on average.


## Implementation of the estimation strategy

This suggests the following strategy for the estimation of the average treatment effect on the treated, called $\delta$
Step 1
Estimate the propensity score (see Annex 3)
Step 2
Restrict the analysis to the region of common support (key source of bias in observational studies)
Step 3
Estimate the average treatment effect by matching based on the propensity score

## Implementation of the estimation strategy

## Step 1

Estimate the propensity score (see annex 3)
Estimate a logit (or probit) model of program participation. Predicted values are the "propensity scores".
E.g. With a logit function, see Annex 3.

This step is necessary because the "true" propensity score is unknown and therefore the propensity score has to be estimated.

# When is propensity score matching appropriate? 

- Idea behind propensity score matching: estimation of treatment effects requires a careful matching of treated and controls.
- If treated and controls are very different in terms of observables this matching is not sufficiently close and reliable or it may even be impossible.

The comparison of the estimated propensity scores across treated and controls provides a useful diagnostic tool to evaluate how similar are treated and controls, and therefore how reliable is the estimation strategy.

## Implementation of the estimation strategy

Step 2
Restrict the analysis to the region of common support (key source of bias in observational studies)

# So you want propensity score to be the "same" for treatments and controls... 

- The range of variation of propensity scores should be the same for treated and controls.
- Count how many controls have a propensity score lower than the minimum or higher than the maximum of the propensity scores of the treated
- and vice versa.
- Frequency of propensity scores is the same for treated and control.
- Draw histograms of the estimated propensity scores for the treated and controls.
- The bins correspond to the blocks constructed for the estimation of propensity scores.


## The issue of common support



Propensity score

## Step 3: Estimate the average treatment effect by matching based on the propensity score

## Step 3

Estimate the average treatment effect by matching based on the propensity score

- For each participant find a sample of non-participants that have "similar" propensity scores.
- Compare the outcome indicator for each participant and its comparison group.
- Calculate the mean of these individual gains to obtain the average overall gain (Heckman et al., 1998).

$$
A T T=\delta=\sum_{j=1}^{P}\left(Y_{j 1}-\sum_{i=1}^{N P} W_{i j} Y_{i j 0}\right) / P
$$

## Step 3: Estimate the average treatment effect by matching based on the propensity score

$\sigma$
"Similar" can be defined in many ways. These different weights correspond to different ways of doing matching:

- Stratification on the Score
- Nearest neighbor matching on the Score
- Radius matching on the Score
- Kernel matching on the Score
- Weighting on the basis of the Score


## To summarize:

- Matching is the observational analogue of an experiment in which placement is independent of outcomes
- The key difference is that a pure experiment does not require the essentially untestable assumption of independence conditional on observables.
- PSM requires
- No selection on unobservables
- Good data
- Common Support
- Matching is often combined with difference-in-differences methods
- Matching is performed on baseline characteristics
- DD controls for selection based on time-invariant unobserved characteristics
- Caution! Matching with ex post data is very risky


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## Thank You

$Q \& A$

## Annex 1: Average effects of treatment on the treated assuming unconfoundedness given $X$

If we are willing to assume unconfoundedness:

$$
\begin{aligned}
& E_{i}\left\{Y_{0}\left(u_{i}\right) \mid D_{i}=0, X\right\}=E_{i}\left\{Y_{0}\left(u_{i}\right) \mid D_{i}=1, X\right\}=E_{i}\left\{Y_{0}\left(u_{i}\right) \mid X\right\} \\
& E_{i}\left\{Y_{1}\left(u_{i}\right) \mid D_{i}=0, X\right\}=E_{i}\left\{Y_{1}\left(u_{i}\right) \mid D_{i}=1, X\right\}=E_{i}\left\{Y_{1}\left(u_{i}\right) \mid X\right\}
\end{aligned}
$$

Using these expressions, we can define for each cell defined by $X$
$\delta_{X}=$ average treatment effect on the treated in cell defined by $X$

$$
\begin{aligned}
& =E_{i}\left\{\Delta_{i} \mid D_{i}=1, X\right\} \\
& =E_{i}\left\{Y_{1}\left(u_{i}\right)-Y_{0}\left(u_{i}\right) \mid D_{i}=1, X\right\} \\
& =\underbrace{E_{i}\left\{Y_{1}\left(u_{i}\right) \mid D_{i}=1, X\right\}}_{\text {can measure sample analog }}-\underbrace{E\left\{Y_{0}\left(u_{i}\right) \mid D_{i}=1, X\right\}}_{\text {can NOT measure sample analog }}
\end{aligned}
$$

$$
=E_{i}\left\{Y_{1}\left(u_{i}\right) \mid D_{i}=1, X\right\}-\underbrace{E_{i}\left\{Y_{0}\left(u_{i}\right) \mid D_{i}=0, X\right\}}_{\text {can measure sample analog }}
$$

## Annex 1: Average effects of treatment on the treated assuming unconfoundedness given $X$

Now what is the relation between
$\delta$ "average treatment effect on the treated"... and....
$\delta_{X}$ "average treatment effect on the treated within cell defined by $X " ?$

$$
\begin{aligned}
\delta & =\text { average treatment effect on the treated } \\
& =E_{i}\left\{\Delta_{i} \mid D_{i}=1\right\}
\end{aligned}
$$

$\Downarrow$ by the law of iterated expectations
$=E_{i}\left\{E_{X}\left[\Delta_{i} \mid D_{i}=1, X\right]\right\}$
$=E_{X}\left\{E_{i}\left[\Delta_{i} \mid D_{i}=1, X\right]\right\}$
$=E_{X}\left\{\delta_{X}\right\}$
$=E_{X}\{$ average treatment effect on the treated within cell defined by $X\}$

## Annex 2: Average effects of treatment and the propensity score

$$
\begin{aligned}
& \text { So let's match treatments and controls } \\
& \text { on the basis of the propensity score } \mathrm{p}(\mathrm{X}) \text { instead of } \mathrm{X} \text {. } \\
& E_{i}\left\{Y_{0}\left(u_{i}\right) \mid D_{i}=0, p\left(X_{i}\right)\right\}=E_{i}\left\{Y_{0}\left(u_{i}\right) \mid D_{i}=1, p\left(X_{i}\right)\right\}=E_{i}\left\{Y_{0}\left(u_{i}\right) \mid p\left(X_{i}\right)\right\} \\
& E_{i}\left\{Y_{1}\left(u_{i}\right) \mid D_{i}=0, p\left(X_{i}\right)\right\}=E_{i}\left\{Y_{1}\left(u_{i}\right) \mid D_{i}=1, p\left(X_{i}\right)\right\}=E_{i}\left\{Y_{1}\left(u_{i}\right) \mid p\left(X_{i}\right)\right\}
\end{aligned}
$$

$$
\begin{aligned}
& \text { Using these expressions, we can define f cell defined by } p(X) \\
\delta_{p(X)} & =\text { average treatment effect on the treated in cell defined by } p(X) \\
& =E_{i}\left\{\Delta_{i} \mid D_{i}=1, p(X)\right\} \\
& =E_{i}\left\{Y_{1}\left(u_{i}\right)-Y_{0}\left(u_{i}\right) \mid D_{i}=1, p(X)\right\} \\
& =\underbrace{E_{i}\left\{Y_{1}\left(u_{i}\right) \mid D_{i}=1, p(X)\right\}}_{\text {can measure sample analog }}-\underbrace{E\left\{Y_{0}\left(u_{i}\right) \mid D_{i}=1, p(X)\right\}}_{\text {can NOT measure sample analog }} \\
& =E_{i}\left\{Y_{1}\left(u_{i}\right) \mid D_{i}=1, p(X)\right\}-\underbrace{E_{i}\left\{Y_{0}\left(u_{i}\right) \mid D_{i}=0, p(X)\right\}}_{\text {can measure sample analog }}
\end{aligned}
$$

## Annex 2: Average effects of treatment and the propensity score

Now what is the relation between
$\delta$ "average treatment effect on the treated"... and....
$\delta_{p(X)}$ "average treatment effect on the treated within cell defined by $p(X)$ "?

$$
\begin{aligned}
\delta & =\text { average treatment effect on the treated } \\
& =E_{i}\left\{\Delta_{i} \mid D_{i}=1\right\} \\
& \Downarrow \text { by the law of iterated expectations } \\
& =E_{i}\left\{E_{p(X)}\left[\Delta_{i} \mid D_{i}=1, p(X)\right]\right\} \\
& =E_{p(X)}\left\{E_{i}\left[\Delta_{i} \mid D_{i}=1, p(X)\right]\right\} \\
& =E_{p(X)}\left\{\delta_{p(X)}\right\} \\
& =E_{p(X)}\{\text { treatment effect on the treated within cell defined by } p(X)\}
\end{aligned}
$$

## Annex 3: Estimation of the propensity score

Any standard probability model can be used to estimate the propensity score, e.g. a logit model:

$$
\begin{equation*}
\operatorname{Pr}\left\{D_{i} \mid X_{i}\right\}=\frac{e^{\lambda h\left(X_{i}\right)}}{1+e^{\operatorname{\lambda h}\left(X_{i}\right)}} \tag{16}
\end{equation*}
$$

where $h(X i)$ is a function of covariates with linear and higher order terms.

## Estimation of the propensity score

- Which higher order terms do you include in $\mathrm{h}(\mathrm{Xi})$ ?
This is determined solely by the need to obtain an estimate of the propensity score that satisfies the balancing property.
- The specification of $h(X i)$ is (1) more parsimonious than the full set of interactions between observables $X$ (2) though not too parsimonious: it still needs to satisfy the balancing property.

Note: the estimation of the propensity scores does not need a behavioral interpretation.

## An algorithm for estimating the propensity score

1. Start with a parsimonious logit or probit function to estimate the score.
2. Sort the data according to the estimated propensity score (from lowest to highest).
3. Stratify all observations in blocks such that in each block the estimated propensity scores for the treated and the controls are not statistically different:
a) start with five blocks of equal score range $\{0-0.2, \ldots, 0.8-1\}$
b) test whether the means of the scores for the treated and the controls are statistically different in each block
c) if yes, increase the number of blocks and test again
d) if no, go to next step.

## An algorithm for estimating the propensity score (continued)

4. Test that the balancing property holds in all blocks for all covariates:
a) for each covariate, test whether the means (and possibly higher order moments) for the treated and for the controls are statistically different in all blocks;
b) if one covariate is not balanced in one block, split the block and test again within each finer block;
c) if one covariate is not balanced in all blocks, modify the logit estimation of the propensity score adding more interaction and higher order terms and then test again.

Note: In all this procedure the outcome has no role.
Use the STATA program pscore.ado, psmatch2.ado, match.ado
(from STATA type "findit 'name ado')

