<span id="page-0-0"></span>MY457/MY557: Causal Inference for Observational and Experimental Studies

Week 1: Causal Frameworks

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## A (Very) Simple Causal Model



## Effects of Causes

Causes and their effects have two properties: they are successive and can be reasoned about in counterfactual terms:

*[...] We may define a cause to be an object followed by another, [...] where, if the first object had not been, the second never had existed.*

*— Hume, 1748*

*[...] would not have died if he had not eaten of it, people would be apt to say that eating of that dish was the cause of his death.*

*— Mill, 1843*

One important implication is that causal variables must be manipulable:

*No causation without manipulation.*

*— Holland, 1986*

#### Good Causal Questions

Manipulability means we must think very carefully about causal questions...

- (Largely) immutable characteristics?
	- $\bullet$  Judges' sex assigned at birth  $\rightarrow$  decision making
	- Race and ethnicity  $\rightarrow$  employment outcomes
	- $\bullet$  Country of origin  $\rightarrow$  political beliefs
- Major global events?
	- Russian revolution  $\rightarrow$  Karl Marx's intellectual popularity
	- $\bullet$  9/11  $\rightarrow$  Arab Spring
- Non-successive chains?
	- Monthly expenditure  $\rightarrow$  monthly savings
	- $\bullet$  Holocaust  $\rightarrow$  modern AFD election support

## Neyman Urn Model



# Concepts: Treatment, Outcomes, and Potential Outcomes

#### Definition (Treatment)

*Di* : Indicator of treatment intake for *unit i*

$$
D_i = \left\{ \begin{array}{ll} 1 & \text{if unit } i \text{ received the treatment} \\ 0 & \text{otherwise.} \end{array} \right.
$$

#### Definition (Observed outcome)

*Yi* : Observed outcome variable of interest for unit *i*

#### Definition (Potential Outcome)

*Y*0*<sup>i</sup>* and *Y*1*<sup>i</sup>* : Potential outcomes for unit *i*:

- $Y_{1i}$  Outcome for unit *i* when  $D_i = 1$
- $Y_{0i}$  Outcome for unit *i* when  $D_i = 0$

(Alternative notation:  $Y_i(d)$ ,  $Y_i^d$ , etc.)

#### Concepts: Treatment, Outcomes, and Potential Outcomes

Under further assumptions ('SUTVA', more later), we can connect these three concepts mathematically:

$$
Y_{i} = D_{i} \cdot Y_{1i} + (1 - D_{i}) \cdot Y_{0i}
$$
  
i.e. 
$$
Y_{i} = \begin{cases} Y_{1i} & \text{if } D_{i} = 1 \\ Y_{0i} & \text{if } D_{i} = 0 \end{cases}
$$

- *A priori* each potential outcome could be observed (manipulability!)
- After treatment assignment, one is observed, the other is counterfactual

## Neyman Urn Model



## Neyman Urn Model



## Stable Unit Treatment Value Assumption (SUTVA)

- $\mathsf{Recall: Y_i = Y_{D_i}$ , or equivalently  $\mathsf{Y}_i = D_i \mathsf{Y}_{1i} + (1 D_i) \mathsf{Y}_{0i}$
- This notation implicitly makes the following assumption:

Assumption (SUTVA)

$$
Y_{(D_1, D_2, ..., D_N)i} = Y_{(D'_1, D'_2, ..., D'_N)i} \quad \text{if} \quad D_i = D'_i
$$

SUTVA comprises two sub-assumptions:

- **1** No interference between units
	- Potential outcomes for a unit not affected by treatment status of other units
	- Violations: spill-over effects, contagion, dilution
- 2 No different versions of treatment (stability, consistency)
	- Nominally identical treatments are in fact identical
	- Violations: variable levels of treatment, technical errors

#### Causal Inference Without SUTVA

Let  $\mathbf{D} = (D_1, D_2)$  be a vector of binary treatments for  $N = 2$ .

How many different values can **D** possibly take?

 $(D_1, D_2) = (0, 0)$  or  $(1, 0)$  or  $(0, 1)$  or  $(1, 1)$ 

How many potential outcomes for unit 1?

*Y*(0,0)<sup>1</sup> , *Y*(1,0)<sup>1</sup> , *Y*(0,1)<sup>1</sup> *Y*(1,1)<sup>1</sup> .

How many causal effects for unit 1?

$$
Y_{(1,1)1} - Y_{(0,0)1}, \t Y_{(1,1)1} - Y_{(0,1)1},
$$
  
\n
$$
Y_{(1,0)1} - Y_{(0,0)1}, \t Y_{(1,0)1} - Y_{(0,1)1},
$$
  
\n
$$
Y_{(1,1)1} - Y_{(1,0)1}, \t Y_{(0,1)1} - Y_{(0,0)1}.
$$

How many observed outcomes for unit 1? Only one:  $Y_1 = Y_{(D_1,D_2)1}$ Without SUTVA, causal inference is exponentially more difficult as *n* ↑.

#### Causal Inference as a Missing Data Problem

Imagine a population with 4 units:



- We take the values of both *Y*1*<sup>i</sup>* and *Y*0*<sup>i</sup>* to be real and fixed for all *i*
- But we can only observe one of them for any *i* ...
- This is known as the fundamental problem of causal inference (FPCI)

#### Causal Inference as a Missing Data Problem

... because of the FPCI we see only this:



Our goal:

- define causal estimands in terms of potential outcomes (previous table)
- **e** estimate them using observable data on this slide (previous table)
- **e** essentially: fill in the missing counterfactuals as best as possible!

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#### Esti-what?

#### **Estimand**:

 $\rightarrow$  Unobserved population parameter or function.

#### **Estimator**:

 $\rightarrow$  A function that can be applied to observed data.

#### **Estimate**:

 $\rightarrow$  A specific output of said function.

#### Unit-Level Causal Estimands

#### Definition (Individual Treatment Effect)

Causal effect of the treatment on the outcome for unit *i*, defined by the comparison of two potential outcomes:

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

This cannot be observed, and is also very hard to estimate:

- We cannot observe both potential outcomes *Y*1*<sup>i</sup>* and *Y*0*<sup>i</sup>* for the same unit *i*.
- Hard to reliably fill in the missing potential outcome for any one unit *i*.

#### Group-Level Causal Estimands

- Consider a fixed group (population) of units  $i = 1, \ldots, N$
- Values of the potential outcomes for this population can be represented as two vectors:

$$
\mathbf{Y}_1 = (Y_{11}, Y_{12}, \ldots, Y_{1N})
$$

$$
\mathbf{Y}_0 = (Y_{01}, Y_{02}, \ldots, Y_{0N})
$$

- **■** A population causal estimand is a comparison of **Y**<sub>1</sub> and **Y**<sub>0</sub>
- A common choice is a difference of their expected values (means).

## Causal Estimand: The ATE

Definition (Average treatment effect, ATE)

$$
\tau_{ATE} = \frac{1}{N} \sum_{i=1}^N (Y_{1i} - Y_{0i})
$$

or equivalently

$$
\tau_{ATE} = \mathbb{E}[Y_{1i} - Y_{0i}]
$$

- **•** In the rest of this course, we will consider various assumptions under which τ*ATE* can be identified from observed information
- Note on notation: We represent the estimand as a greek letter (in this case  $\tau$ , but could be anything). We typically represent an estimator for that estimand as a greek letter with something on top (e.g.  $\tilde{\tau}$  or  $\hat{\tau}$ ). An estimate will be a realised number (interval, etc.).

#### Causal Estimand: The ATT

Definition (Average treatment effect on the treated, ATT)

$$
\tau_{ATT} = \frac{1}{N_1} \sum_{i=1}^{N} D_i (Y_{1i} - Y_{0i}) \text{ where } N_1 = \sum_{i=1}^{N} D_i
$$

or equivalently

$$
\tau_{ATT} = \mathbb{E}[Y_{1i} - Y_{0i} | D_i = 1]
$$

(Note: The mathematical symbol | means "conditional on".)

- **In words, N<sub>1</sub> equals the number of treated units.**
- When would  $\tau_{ATT} \neq \tau_{ATF}$ ? When  $D_i$  and  $Y_{di}$  are associated.
- Exercise: Define τ*ATU*, ATE on the untreated (control) units, also called the ATU.  $\tau_{ATU} = \mathbb{E}[Y_{1i} - Y_{0i} | D_i = 0]$

# Causal Estimand: The CATE

Definition (Conditional average treatment effects, CATE)

```
\tau_{CATE}(x) = \mathbb{E}[Y_{1i} - Y_{0i}|X_i = x]
```
where *X<sup>i</sup>* is a pre-treatment covariate for unit *i*

- **•** In words,  $τ_{CATE}(x)$  is a subgroup effect, treatment effect on units who have particular characteristics *x*.
- This estimand sometimes goes by other names (e.g. local average treatment effect or LATE).
- This is an increasing important area for causal inference (e.g. optimal policy targeting), and we will return to it later!

## Illustration: Average Treatment Effect

Let's return to our population of 4 units:



Why does  $\tau_{\text{ATE}} \neq \tilde{\tau}$ ? When would they be equal?

## Illustration: Average Treatment Effect on the Treated

Again suppose we observe a population of 4 units:



$$
\tau_{ATT} = \mathbb{E}[Y_{1i} - Y_{0i} | D_i = 1] = \mathbb{E}[\tau_i | D_i = 1] = \frac{3+0}{2} = 1.5.
$$

#### Average Treatment Effect on the Treated

Why does  $\tau_{ATT} \neq \tau_{ATF}$ ?

Because  $\mathbb{E}[Y_{1i}] \neq \mathbb{E}[Y_{1i}|D_i = 1]$  (and likewise for  $\mathbb{E}[Y_{0i}])$ 

That is, *D<sup>i</sup>* and *Ydi* are associated

## Internal and External Validity

- Note that when we talk about the 'population' in causal inference settings we often mean *only* to the *N* units for whom we have observed data (i.e. what we would typically call the 'sample')
- The estimands considered on this course are defined and estimated for this population (not for some super-population from which a sample was drawn)
- Internal validity refers to the validity of our estimates of these effects. This class is focused only on internal validity.

#### Internal and External Validity

- External validity refers to the validity generalising our estimates of causal effects from the 'population' of *N* units to any other population (note, this could include generalising from a realised sample to a population)
- If claimed, external validity has to be justified by different kinds of arguments, e.g.
	- Representative sampling (ideally probability sampling) of the *N* units from a larger population. This is a population inference task, as in survey research (see MY456).)
	- Some re-weighting strategy designed to adjust the observed sample. Again, this is a population inference task.
	- Substantive theory / assumptions / wishful thinking about why a causal effect for these *N* units would also apply elsewhere

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## Identification Problem for Causal Inference

- In statistics, an estimand (parameter) is identified if its value can be uniquely estimated based on the observed data and unidentified if not
- Recall that in causal inference, estimands are population causal effects but the FPCI tells us that at least half of the potential outcomes are always missing
- An identification strategy is a combination of data and assumptions which allows us to identify a causal estimand by estimating ("filling in") the missing potential outcomes (usually at a group level)

#### Selection Bias

Consider again the naïve difference of observed means in the treatment groups:

$$
\underbrace{E(Y_i|D_i=1)-E(Y_i|D_i=0)}_{\text{Observed difference in average}} = E(Y_{1i}|D_i=1)-E(Y_{0i}|D_i=0)
$$
\n
$$
= \underbrace{E(Y_{1i}|D_i=1)-E(Y_{0i}|D_i=1)}_{\text{ATT}} + \underbrace{E(Y_{0i}|D_i=1)-E(Y_{0i}|D_i=0)}_{\text{Selection bias}}
$$

- The same observed mean difference could be due to different combinations of the ATT (estimand!) and selection bias terms. We might say the causal effect of *D* on *Y* is confounded.
- Thus ATT is not identified from the naïve observed mean difference: it is not uniquely mapped from the observed data. We need more assumptions.
- Correlation [association, here observed mean difference] is not necessarily causation.

#### Selection Bias

$$
E(Y_i|D_i = 1) - E(Y_i|D_i = 0)
$$
  
= 
$$
[E(Y_{1i}|D_i = 1) - E(Y_{0i}|D_i = 1)] + [E(Y_{0i}|D_i = 1) - E(Y_{0i}|D_i = 0)]
$$

- $E(Y_{0i}|D_i=1)-E(Y_{0i}|D_i=0)$  is referred to as selection bias because if it is not 0, it implies treatment and control groups are systematically different in potential outcome  $Y_{0i}$ .
- Canonical example: Job training program
	- participants are self-selected from a population of individuals in difficult labor situations
	- perhaps better resourced or more motivated individuals decide to take part
	- $\bullet$  even in the absence of the program, post-training period earnings for those people would then have been higher than those for those who did not opt in  $(E[Y_0|D = 1] - E[Y_0|D = 0] > 0)$

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# Causal Diagrams  $U_{Z}$

So far we have reasoned about causal effects using potential outcomes. An alternative (but intimately connected) framework is the graphical approach.

This uses causal diagrams, tools that allow us to:

- 1. Specify the variables (observed and unobserved) we care about
- 2. Specify how those variables are connected
- 3. See what we can learn about causal effects, and with what assumptions.

This can help us to:

- 1. Study how conditioning affects our research designs
- 2. Create new research designs and methodologies.

#### Causal Diagrams as Directed Acyclic Graphs



Components of a causal diagram as a Directed Acyclic Graphs (DAG):

- Nodes: Representing "variables" (also called vertices)
- Directed Edges: Encoding one-way (causal) relationships
	- $\rightarrow$  This implies nodes are ordered (each pair: head and tail)
	- $\rightarrow$  These connections can be observed (solid) or unobserved (dashed)

Features of a DAG:

- Acyclic: No directed cycles (e.g. A does not terminate A)
- Non-connections: The absence of relationships between variables

## Directed Acyclic Graphs: Example





"These aren't the DAGs you're looking for"

## Directed Acyclic Graphs: Example



What can we learn from this DAG?

- $\bullet$  *Z*  $\rightarrow$  *Y* is confounded by *W*
- $\bullet$  *D*  $\rightarrow$  *Y* is confounded by *Q*
- $\bullet$  *Z*  $\rightarrow$  *D* is identified

But only if our DAG is correct!

#### Representing Interventions



Treatments (interventions) are represented by the *do*() operator. For example,  $do(d_0)$  holds  $D = d_0$  exogenously.

#### Identification



ATE of *D* on *Y* defined as the average difference in *Y* between two interventions:

```
\mathbb{E}[Y \mid do(d_1)] - \mathbb{E}[Y \mid do(d_0)]
```
Problem: Can this be estimated without an explicit intervention (identification)? Insight: If the DAG is equivalent with and without *do*(), yes.

Generally: We can identify the effect of *D* on *Y* if all back-door paths are blocked.

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## Assignment Mechanism

#### Definition (Assignment Mechanism)

The assignment mechanism is the procedure that determines the treatment status of each unit.

- Most causal inference methods achieve identification by restricting the (assumed) assignment mechanism
- For example, if we are willing to assume that treatment assignment is independent of potential outcomes under no treatment, then:

$$
E(Y_{0i}|D_i=1)-E(Y_{0i}|D_i=0)=0
$$

i.e. selection bias is zero and the observed mean difference is (in expectation) equal to ATT (and also ATE in that case)

#### Different Assignment Mechanisms

Imbens and Rubin (2015, Ch. 3) present three assumptions about assignment mechanisms (for each unit) that provide the grounds for identification:

- 1. *Individualistic*: Assignment does not depend on the covariates or potential outcomes for other units.
- 2. *Probabilistic*: There is a nonzero probability of each treatment value, for every unit.
- 3. *Unconfounded*: Assignment does not depend on potential outcomes.

Assuming the above, we can distinguish:

- Experiments: The assignment mechanism is both known and controlled by the researcher, and
- Observational studies: The assignment mechanism is not known to, or not under the control of, the researcher

# Our Key Assignment Mechanisms

#### Randomised Experiments:

- These come in many flavours, only a few of which we will discuss!
	- $\rightarrow$  Within designs, between designs
	- $\rightarrow$  Unit-randomized, cluster-randomized, dynamic randomization
	- $\rightarrow$  Crossover designs, stepped-wedge designs, etc. etc. etc.

#### Observational Studies:

- Adjustment: Selection on observables with regression, matching, etc.
- Temporal: Diff-in-diff, event studies, synthetic control methods
- **.** Instrumental variables, shift-share designs, etc.
- Sharp and fuzzy regression discontinuity designs

# The Continuum of Credibility**TM**



Key point: The art (and science) of applied causal inference is making defensible assumptions. There is no 'magic' solution to the fundamental problem of causal inference, only assumptions all the way down!

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#### <span id="page-45-0"></span>Key ideas from this week

- Learned to think about causal effects in terms of potential outcomes, not realized (observed) outcomes
- $\bullet$  Observed association is neither necessary nor sufficient for causality  $\sim$ focused on one big problem, selection bias
- $\bullet$  Introduced an alternative framework for thinking about causal models the graphical approach
- Learning about causal effects should start from understanding the assignment mechanism for treatment
- **E** Evaluate the plausibility of your assumptions to understand the credibility of your conclusions