CAUSAL NETWORKS

POTENTIAL OUTCOMES

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POTENTIAL OUTCOMES

In this lecture you will learn about new concepts and notations needed to clearly describe causal concepts.

In particular, the lecture presents and discusses the following:

- Potential Outcomes and Individual Treatment Effects
- The Fundamental Problem of Causal Inference
 - Average Treatment Effects and Missing Data Interpretation
 - Ignorability Exchangeability
 - Conditional Exchangeability Unconfoundedness
 - Positivity Overlap Common Support and Extrapolation
 - No interference, Consistency, and SUTVA

POTENTIAL OUTCOMES

PART I

POTENTIAL OUTCOMES AND INDIVIDUAL TREATMENT EFFECTS

We will use the following notation:

- *X* denotes the random variable for **TREATMENT**,
- *Y* denotes the random variable for the **OUTCOME** of interest,
- Z denotes a set of random variables (COVARIATES),

In general, we will use uppercase letters Z to denote random variables and lowercase letters Z to denote values that random variables take on.

Much of what we consider will be settings where *X* and *Y* are binary.

In general, we can extend things to work in settings where *X* and *Y* can take on more than two values or where *X* and *Y* are continuous.

We have the following narration, you are unhappy (②), and consider whether or not to get a dog (②) to help make you happy (③).

To answer the question to the right we need to know more.

You get the dog and become happy!!!



The dog ($(\c led{m Q})$) **CAUSED** you to be happy ($(\c led{m Q})$)?

What if I told you "I'm certain you would have become happy also without getting the dog"?



The dog was not necessary to make you happy, so its claim to a causal effect on your happiness is weak.

What if instead I told you "I'm certain you would have remained unhappy without getting the dog"?



The dog has a pretty strong claim to a causal effect on your happiness.

We have just used the causal concept known as **POTENTIAL OUTCOMES**.

Happiness = outcome = Y

unhappy

Y = 0

OUTCOME

(Y)

happy

Y = 1

Dog = TREATMENT = X

do not get

X = 0

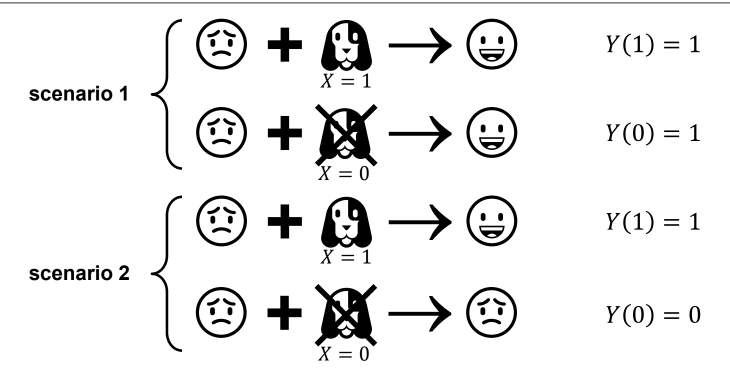
TREATMENT

(X)

get

X = 1

We let Y(1) be the **POTENTIAL OUTCOME OF HAPPINESS** you would observe if you were to get a dog (X = 1). We let Y(0) be the **POTENTIAL OUTCOME OF HAPPINESS** you would observe if you were to not get a dog (X = 0).



We let Y(1) be the **POTENTIAL OUTCOME OF HAPPINESS** you would observe if you were to get a dog (X = 1). We let Y(0) be the **POTENTIAL OUTCOME OF HAPPINESS** you would observe if you were to not get a dog (X = 0).

POTENTIAL OUTCOME

Y(x) denotes what your outcome would be, if you were to take treatment X = x.

- A POTENTIAL OUTCOME Y(x) is distinct from the OBSERVED OUTCOME Y in that not all potential outcomes are observed.
- All potential outcomes can potentially be observed.
- The actually observed potential outcome depends on the given value x of treatment X.

Up to now there is only a single individual in the whole population: you.

However, the **POPULATION** consists of many **INDIVIDUALS** or **UNITS**.

Each individual (unit) is tipically associated with one or more

you 🎵

population individuals units



individual (unit)



age gender weight

ith individual (unit)



 $X_i \mathbf{Z}_i Y_i$



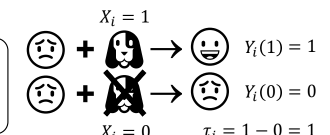
variables, referred to as **COVARIATES Z**.

individual (unit) as X_i , \mathbf{Z}_i and Y_i .

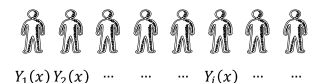
The individual treatment effect (ITE) for the i^{th} individual (unit) is defined as follows:

We denote **TREATMENT** X, **COVARIATES** Z and **OUTCOME** Y of the i^{th}

$$\tau_i \triangleq Y_i(1) - Y_i(0)$$



population individuals units



You

Individual treatment effects (ITEs) are some of the main quantities that we care about in causal inference.

INDIVIDUAL TREATMENT EFFECT - (ITE)

The individual treatment effect (ITE) for the i^{th} individual (unit) is defined as follows:

$$\tau_i \triangleq Y_i(1) - Y_i(0)$$

Y(x) is a random variable (different individuals have different potential outcomes).

 $Y_i(x)$ is treated as a non-random variable we are conditioning on so much individualized (and context-specific) information, that we restrict our focus to a single individual (in a specific context) whose potential outcomes are deterministic.

You would choose to get a dog because the **CAUSAL EFFECT (ITE)** $\tau_i \triangleq Y_i(1) - Y_i(0)$ of getting a dog on your happiness is positive $\tau_i = 1$.

population individuals units







Individual treatment effects (ITEs) are some of the main quantities that we care about in causal inference.

INDIVIDUAL TREATMENT EFFECT - (ITE)

 $Y_1(x) Y_2(x)$

The individual treatment effect (ITE) for the i^{th} individual (unit) is defined as follows:

$$\tau_i \triangleq Y_i(1) - Y_i(0)$$

Y(x) is a random variable (different individuals have different potential outcomes).

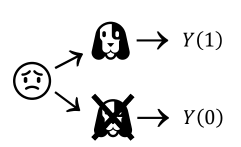
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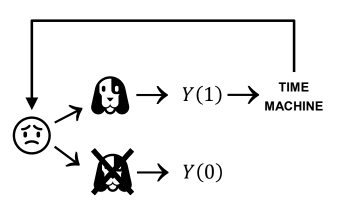
You would decide to not get a dog because there is **NO CAUSAL EFFECT** of getting a dog $\tau_i \triangleq Y_i(1) - Y_i(0) = 1 - 1 = 0.$

POTENTIAL OUTCOMES 11

PART II

THE FUNDAMENTAL PROBLEM OF CAUSAL INFERENCE

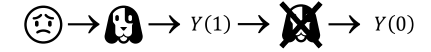




It is impossible to observe all potential outcomes for a given individual (unit).

In particular:

You cannot observe both Y(1) and Y(0), unless you have a **TIME MACHINE** that would allow you to go back in time and choose the version of treatment X that you didn't take the first time.



You cannot simply get a dog, observe Y(1), give the dog away, and then observe Y(0) because the second observation will be influenced by all the actions you took between the two observations and anything else that changed since the first observation.

THE FUNDAMENTAL PROBLEM OF CAUSAL INFERENCE

We cannot observe both $Y_i(1)$ and $Y_i(0)$, therefore we cannot observe the causal effect

$$\tau_i \triangleq Y_i(1) - Y_i(0)$$

Indeed, we care about making causal claims, which are defined in terms of potential outcomes.

In machine learning, we often only care about predicting the observed outcome *Y*, so there is no need for potential outcomes, which means machine learning does not have to deal with this fundamental problem that we must deal with in causal inference.

- The potential outcomes that you do not (and cannot) observe are known as COUNTERFACTUALS because they are counter to fact (reality).
- \longrightarrow \longrightarrow Y(1) COUNTERFACTUAL

- A potential outcome Y(x) does not become counter to fact (**COUNTERFACTUAL**) until another potential outcome Y(x') is observed.
- Note that there are no counterfactuals or factuals until the outcome is observed. Before that, there are only potential outcomes.

$$Y(0) = 1 \text{ FACTUAL}$$

$$X = 0$$

THE FUNDAMENTAL PROBLEM OF CAUSAL INFERENCE

We cannot observe both $Y_i(1)$ and $Y_i(0)$, therefore we cannot observe the causal effect

$$\tau_i \triangleq Y_i(1) - Y_i(0)$$

We know that we can't access individual treatment effects (ITE), due to the fundamental problem of causal inference.

However, what about **AVERAGE TREATMENT EFFECTS?**

AVERAGE TREATMENT EFFECT - ATE

The average treatment effect (ATE) is obtained by taking an average over the ITEs:

$$\tau \triangleq \mathbb{E}[Y_i(1) - Y_i(0)] = \mathbb{E}[Y(1) - Y(0)]$$

where we recall that the average is over the individuals "i" if $Y_i(x)$ is deterministic.

How would we actually compute the ATE?

Let us assume that data in the right table (Table 2.1) represent the whole population of interest.

The **FUNDAMENTAL PROBLEM OF CAUSAL INFERENCE** can be seen as a **MISSING DATA PROBLEM**, i.e., all question marks (?) in Table 2.1 mean that we do not observe the value for the corresponding cell.

Therefore, we can not compute directly the average treatment effect (ATE) or average causal effect (ACE):

$$\tau \triangleq \mathbb{E}[Y_i(1) - Y_i(0)] = ?$$

However, we know the following:

$$\tau \triangleq \mathbb{E}[Y_i(1) - Y_i(0)] = \mathbb{E}[Y(1) - Y(0)]$$

and linearity of expectation $\mathbb{E}[\cdot]$ gives:

Table 2.1

ASSOCIATIONAL DIFFERENCE

We could be tempted to use the associational difference

$$\mathbb{E}[Y|X=1] - \mathbb{E}[Y|X=0]$$

$$\tau \triangleq \mathbb{E}[Y(1) - Y(0)] = \mathbb{E}[Y(1)] - \mathbb{E}[Y(0)] \stackrel{!}{=} \mathbb{E}[Y|X=1] - \mathbb{E}[Y|X=0]$$

Unfortunately, this is **not true in general**. If it were, that would mean that causation is simply association.

$$\mathbb{E}[Y|X=1] - \mathbb{E}[Y|X=0]$$

is an associational quantity, while

$$\mathbb{E}[Y(1)] - \mathbb{E}[Y(0)]$$

is a causal quantity.

In general, they are not equal due to **CONFOUNDING**.

The graphical representation of such a difference is depicted in Figure 2.1.

In particular, we say that the covariate Z confounds the effect of X on Y, because the of the following path

$$X \leftarrow Z \rightarrow Y$$

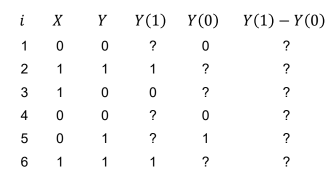
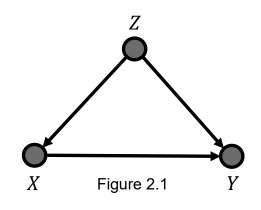


Table 2.1



But ... under which assumptions the following equality holds true?

$$\tau \triangleq \mathbb{E}[Y(1)] - \mathbb{E}[Y(0)] = \mathbb{E}[Y|X=1] - \mathbb{E}[Y|X=0]$$

In other terms, when ATE is equal to the associational difference?

What legitimates us to calculate the ATE by taking the average of the Y(0) column, ignoring the question marks, and subtracting that from the average of the Y(1) column, ignoring the question marks?

Ignoring question marks (missing data) is known as:

IGNORABILITY — EXCHANGEABILITY

$$(Y(1), Y(0)) \perp \!\!\! \perp X$$

Assuming

IGNORABILITY — EXCHANGEABILITY

means we can ignore how a

patient ended up selecting the
treatment *X* she selected and
just assuming she was randomly
assigned her treatment *X*.

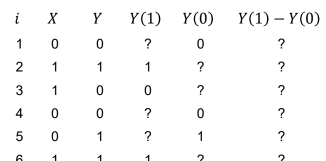
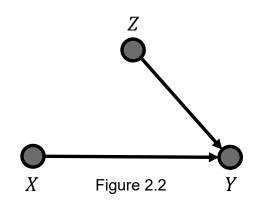


Table 2.1



IGNORABILITY — **EXCHANGEABILITY** is fundamental because it allows us to reduce the ATE to the associational difference:

$$\tau \triangleq \mathbb{E}[Y(1)] - \mathbb{E}[Y(0)]$$

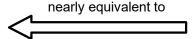
$$= \mathbb{E}[Y(1)|X = 1] - \mathbb{E}[Y(0)|X = 0]$$

$$= \mathbb{E}[Y|X = 1] - \mathbb{E}[Y|X = 0]$$

IGNORABILITY — **EXCHANGEABILITY** means that the groups (treatment/control) are exchangeable in the sense that if they were swapped, the new treatment group would observe the same outcomes as the old treatment group, and the new control group would observe the same outcomes as the old control group.

IGNORABILITY — **EXCHANGEABILITY**

$$(Y(1), Y(0)) \perp \!\!\! \perp X$$



i	X	Y	<i>Y</i> (1)	Y(0)	Y(1) - Y(0)
1	0	0	?	0	?
2	1	1	1	?	?
3	1	0	0	?	?
4	0	0	?	0	?
5	0	1	?	1	?
_				_	_

Table 2.1

$$\mathbb{E}[Y(1)|X=1] = \mathbb{E}[Y(1)|X=0]$$

$$\mathbb{E}[Y(0)|X=0] = \mathbb{E}[Y(0)|X=1]$$
which brings to

MEAN IGNORABILITY — EXCHANGEABILITY

$$\mathbb{E}[Y(1)|X = x] = \mathbb{E}[Y(1)]$$

$$\mathbb{E}[Y(0)|X = x] = \mathbb{E}[Y(0)]$$

IGNORABILITY – **EXCHANGEABILITY** is fundamental because it allows us to reduce the ATE to the associational difference:

$$\tau \triangleq \mathbb{E}[Y(1)] - \mathbb{E}[Y(0)]$$

$$= \mathbb{E}[Y(1)|X=1] - \mathbb{E}[Y(0)|X=0]$$

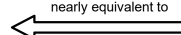
$$= \mathbb{E}[Y|X=1] - \mathbb{E}[Y|X=0]$$

MEAN (IGNORABILITY – EXCHANGEABILITY) is a weaker assumption than **FULL (IGNORABILITY – EXCHANGEABILITY)** (box below), because it only constrains the first moment of the distribution.

In general, **MEAN (IGNORABILITY – EXCHANGEABILITY)** is sufficient for ATE, but it is common to assume complete independence, as formally represented in the box below.

IGNORABILITY — **EXCHANGEABILITY**

$$(Y(1), Y(0)) \perp \!\!\! \perp X$$



 $i \quad X \quad Y \quad Y(1) \quad Y(0) \quad Y(1) - Y(0)$ $1 \quad 0 \quad 0 \quad ? \quad 0 \quad ?$ $2 \quad 1 \quad 1 \quad 1 \quad ? \quad ?$ $3 \quad 1 \quad 0 \quad 0 \quad ? \quad ?$ $4 \quad 0 \quad 0 \quad ? \quad 0 \quad ?$ $5 \quad 0 \quad 1 \quad ? \quad 1 \quad ?$

Table 2.1

$$\mathbb{E}[Y(1)|X=1] = \mathbb{E}[Y(1)|X=0]$$

$$\mathbb{E}[Y(0)|X=0] = \mathbb{E}[Y(0)|X=1]$$
 which brings to

MEAN IGNORABILITY — EXCHANGEABILITY

$$\mathbb{E}[Y(1)|X=x] = \mathbb{E}[Y(1)]$$

 $\forall x$

$$\mathbb{E}[Y(0)|X=x] = \mathbb{E}[Y(0)]$$

IGNORABILITY – **EXCHANGEABILITY** is fundamental because it allows us to reduce the ATE to the associational difference:

$$\tau \triangleq \mathbb{E}[Y(1)] - \mathbb{E}[Y(0)]$$

$$= \mathbb{E}[Y(1)|X=1] - \mathbb{E}[Y(0)|X=0]$$

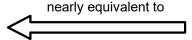
$$= \mathbb{E}[Y|X=1] - \mathbb{E}[Y|X=0]$$

An important intuition to have about **IGNORABILITY** – **EXCHANGEABILITY** is that it guarantees that the groups are comparable.

In other words, the **TREATMENT GROUP** (X = 1) and the **CONTROL GROUP** (X = 0) are the same in all relevant aspects other than the treatment X.

IGNORABILITY — EXCHANGEABILITY

$$(Y(1), Y(0)) \perp \!\!\! \perp X$$



i	X	Y	<i>Y</i> (1)	Y(0)	Y(1)-Y(0)
1	0	0	?	0	?
2	1	1	1	?	?
3	1	0	0	?	?
4	0	0	?	0	?
5	0	1	?	1	?

Table 2.1

$$\mathbb{E}[Y(1)|X=1] = \mathbb{E}[Y(1)|X=0]$$

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MEAN IGNORABILITY — EXCHANGEABILITY

$$\mathbb{E}[Y(1)|X=x] = \mathbb{E}[Y(1)]$$

 $\forall x$

$$\mathbb{E}[Y(0)|X=x] = \mathbb{E}[Y(0)]$$

The assumption of IGNORABILITY - EXCHANGEABILITY allows us to IDENTIFY CAUSAL EFFECTS.

- To IDENTIFY A CAUSAL EFFECT is to reduce a causal expression to a purely statistical expression.
 - To reduce an expression from one that uses potential outcome notation to one that uses only statistical notation such as X, Z, Y, expectations, and conditioning.
 - We can calculate the **CAUSAL EFFECT** from just the **OBSERVATIONAL DISTRIBUTION** P(X, Z, Y)

IDENTIFIABILITY

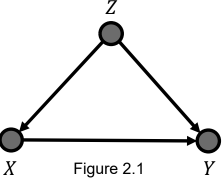
A causal quantity (e.g. $\mathbb{E}[Y(x)]$) is identifiable if we can compute it from a purely statistical quantity (e.g. $\mathbb{E}[Y|X=x]$).

IGNORABILITY — **EXCHANGEABILITY** is extremely important, but how realistic of an assumption is it?

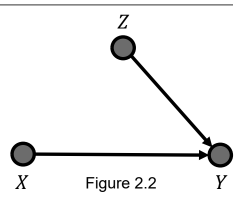
IGNORABILITY — EXCHANGEABILITY

$$(Y(1), Y(0)) \perp \!\!\! \perp X$$

COMPLETELY UNREALISTIC, confounding is likely to happen in most data we observe.



We can make the **IGNORABILITY** — **EXCHANGEABILITY** assumption realistic by performing **RANDOMIZED EXPERIMENTS**, which force the treatment X to not be caused by anything but a coin toss, so then we have the causal structure shown in Figure 2.2.



IDENTIFIABILITY

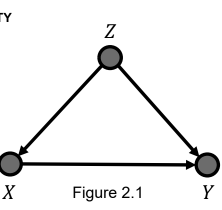
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IGNORABILITY — EXCHANGEABILITY

$$(Y(1), Y(0)) \perp \!\!\! \perp X$$

COMPLETELY UNREALISTIC, confounding is likely to happen in most data we observe.



In **OBSERVATIONAL DATA**, it is unrealistic to assume that the groups (treatment and control) are exchangeable. In other words, there is no reason to expect that the groups (treatment and control) are the same in all relevant variables $Z \in \mathbf{Z}$ other than the treatment X.

However, if we control for relevant variables by conditioning, then maybe the groups will be exchangeable.

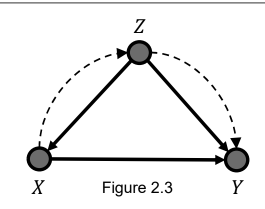
- What do we mean by "relevant variables"?
- For now, let's just say they are all of the covariate variables Z.

CONDITIONAL EXCHANGEABILITY - UNCONFOUNDEDNESS

$$(Y(1), Y(0)) \perp \!\!\! \perp X \mid \mathbf{Z}$$
 where **Z** are the covariate variables.

Although the treatment X and potential outcomes Y(1) and Y(0) may be unconditionally associated (due to confounding), within levels of Z, they are not associated.

No confounding within levels of Z because **CONTROLLING FOR** Z makes the treatment group (X = 1) and the control group (X = 0) comparable.



We do not have **IGNORABILITY** — **EXCHANGEABILITY** in the data because Z is a common cause of X and Y.

NON-CAUSAL ASSOCIATION between X and Y, flows along the following path:

$$X \leftarrow Z \rightarrow Y$$

However, **CONDITIONAL EXCHANGEABILITY** — **UNCONFOUNDEDNESS** holds in the data.

Indeed, when conditioning on Z, non-causal association between X and Y no longer exists.

Non-causal association is "**BLOCKED**" at Z by conditioning on Z.

CONDITIONAL EXCHANGEABILITY — **UNCONFOUNDEDNESS** is the main assumption necessary for causal inference.

CONDITIONAL EXCHANGEABILITY — UNCONFOUNDEDNESS

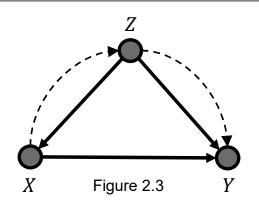
 $(Y(1), Y(0)) \perp \!\!\! \perp X \mid \mathbf{Z}$ where **Z** are the covariate variables.

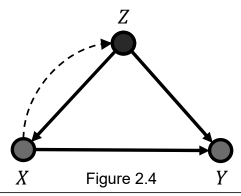
We can now identify the causal effect within levels of **Z**, just like we did with (unconditional) ignorability – exchangeability:

$$\mathbb{E}[Y(1) - Y(0) \mid \mathbf{Z}] = \mathbb{E}[Y(1)|\mathbf{Z}] - \mathbb{E}[Y(0)|\mathbf{Z}]$$

$$= \mathbb{E}[Y(1)|X = 1, \mathbf{Z}] - \mathbb{E}[Y(0)|X = 0, \mathbf{Z}]$$

$$= \mathbb{E}[Y|X = 1, \mathbf{Z}] - \mathbb{E}[Y|X = 0, \mathbf{Z}]$$





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CONDITIONAL EXCHANGEABILITY — UNCONFOUNDEDNESS

 $(Y(1), Y(0)) \perp \!\!\! \perp X \mid \mathbf{Z}$ where **Z** are the covariate variables.

We can now identify the causal effect within levels of \mathbf{Z} , just like we did with (unconditional) ignorability – exchangeability:

$$\mathbb{E}[Y(1) - Y(0) \mid \mathbf{Z}] = \mathbb{E}[Y(1)|\mathbf{Z}] - \mathbb{E}[Y(0)|\mathbf{Z}]$$
$$= \mathbb{E}[Y(1)|X = 1, \mathbf{Z}] - \mathbb{E}[Y(0)|X = 0, \mathbf{Z}]$$
$$= \mathbb{E}[Y|X = 1, \mathbf{Z}] - \mathbb{E}[Y|X = 0, \mathbf{Z}]$$

Conditional exchangeability is a core assumption for causal inference and goes by many names:

- unconfoundedness,
- conditional ignorability,
- no unobserved confounding,
- selection on observables,
- no omitted variable bias
- etc...

I will use the term

UNCONFOUNDEDNESS

If we want the marginal effect that we had before when assuming (unconditional) ignorability – exchangeability, we can get that by simply marginalizing out **Z** as follows

$$\mathbb{E}[Y(1) - Y(0)] = \mathbb{E}_{\mathbf{Z}} \mathbb{E}[Y(1) - Y(0) | \mathbf{Z}]$$
$$= \mathbb{E}_{\mathbf{Z}} [\mathbb{E}[Y | X = 1, \mathbf{Z}] - \mathbb{E}[Y | X = 0, \mathbf{Z}]]$$

We can now introduce an important result for causal inference, that we will formally prove in next lectures.

ADJUSTMENT FORMULA

Given the assumptions of unconfoundedness, positivity, consistency, and no interference, we can identify the ATE:

$$\mathbb{E}[Y(1) - Y(0)] = \mathbb{E}_{\mathbf{Z}}[\mathbb{E}[Y|X=1,\mathbf{Z}] - \mathbb{E}[Y|X=0,\mathbf{Z}]]$$

We moved from the assumption of ignorability – exchangeability to that of unconfoundedness because it seems more realistic.

However, we often cannot know for certain if unconfoundedness holds!!!

Conditional exchangeability is a core assumption for causal inference and goes by many names:

- unconfoundedness,
- conditional ignorability,
- no unobserved confounding,
- selection on observables,
- no omitted variable bias
- etc...

I will use the term

UNCONFOUNDEDNESS

There may be some **UNOBSERVED CONFOUNDERS** (W in Figure 2.3B) that are not part of $\mathbf{Z} = \{M\}$, meaning **UNCONFOUNDEDNESS IS VIOLATED**.

Fortunately, that is not a problem in **RANDOMIZED EXPERIMENTS**.

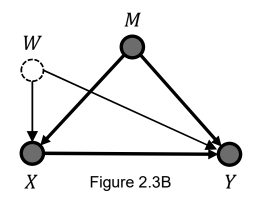
Unfortunately, it is something that we must always be conscious of in **OBSERVATIONAL DATA**.

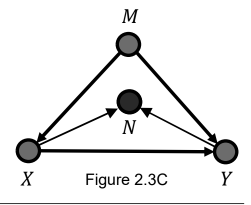
Intuitively, the best thing we can do is to observe and fit (adjust for) as many covariates into **Z** as possible to try to ensure unconfoundedness.

However, we will see in next lectures, that it is not necessarily true that conditioning on more covariates **Z** always helps our causal estimates to be less biased.

Indeed, it can be the case we obtain more biased estimates when adjusting for the "wrong covariates" in $\mathbf{Z} = \{M, N\}$.

In Figure 2.3C, adjusting for M provides unbiased estimates, while adjusting for N results in biased estimates.





Conditioning on many covariates is attractive for achieving unconfoundedness, but it can be detrimental for another reason that has to do with **POSITIVITY**.

Positivity is the condition that all subgroups of the data with different value z for covariates Z have some probability of receiving any value of treatment X.

POSITIVITY — OVERLAP — COMMON SUPPORT

For all values z of covariates z present in the population of interest (i.e., z such that P(z = z) > 0)

$$0 < P(X = 1 | \mathbf{Z} = \mathbf{z}) < 1$$

If we have a **POSITIVITY VIOLATION**, then we will be conditioning on a zero probability event.

To clearly see how a positivity violation translates to division by zero, let's rewrite the right-hand side of

$$\mathbb{E}[Y(1) - Y(0)] = \mathbb{E}_{\mathbf{Z}}[\mathbb{E}[Y|X=1,\mathbf{Z}] - \mathbb{E}[Y|X=0,\mathbf{Z}]]$$

in the case of discrete covariate variables Z, to obtain

POSITIVITY IS ESSENTIAL TO DEFINE CAUSAL EFFECT!!!

$$= \sum_{\mathbf{z}} P(\mathbf{Z} = \mathbf{z}) \left(\sum_{\mathbf{y}} y P(Y = y | X = 1, \mathbf{Z} = \mathbf{z}) - \sum_{\mathbf{y}} y P(Y = y | X = 0, \mathbf{Z} = \mathbf{z}) \right)$$

$$= \sum_{\mathbf{z}} P(\mathbf{Z} = \mathbf{z}) \left(\sum_{\mathbf{y}} y \frac{P(Y = y, X = 1, \mathbf{Z} = \mathbf{z})}{P(X = 1 | \mathbf{Z} = \mathbf{z}) P(\mathbf{Z} = \mathbf{z})} - \sum_{\mathbf{y}} y \frac{P(Y = y, X = 0, \mathbf{Z} = \mathbf{z})}{P(X = 0 | \mathbf{Z} = \mathbf{z}) P(\mathbf{Z} = \mathbf{z})} \right)$$

It wouldn't make any sense to be able to estimate the causal effect of treatment (X = 1) vs. control (X = 0) in subgroup $\mathbf{Z} = \mathbf{z}$, since we see only treatment (X = 1) or only control (X = 0), i.e., we never see the alternative in subgroup $\mathbf{Z} = \mathbf{z}$.

• $if \exists z : P(X = 1 | \mathbf{Z} = z) = 0$

All units belonging to subgroup $\mathbf{Z} = \mathbf{z}$ do not receive the treatment (X = 0).

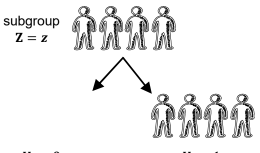
POSITIVITY — OVERLAP — COMMON SUPPORT

For all values z of covariates Z present in the population of interest (i.e., z such that P(Z = z) > 0)

$$0 < P(X = 1 | \mathbf{Z} = \mathbf{z}) < 1$$

• $if \exists z : P(X = 1 | \mathbf{Z} = z) = 1$

All units belonging to subgroup $\mathbf{Z} = \mathbf{z}$ do receive the treatment (X = 1).



X=1

Positivity is also referred to as **OVERLAP**, in the sense we want the covariate distribution of the treatment group (X = 1)

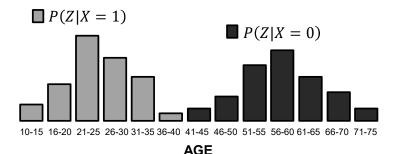
$$P(\mathbf{Z}|X=1)$$

to overlap with the covariate distribution of the control group (X = 0)

$$P(\mathbf{Z}|X=0)$$

This is why another common alias for positivity is **COMMON SUPPORT**.

NO POSITIVITY - NO OVERLAP - NO COMMON SUPPORT



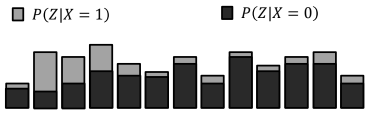
POSITIVITY — OVERLAP — COMMON SUPPORT

For all values z of covariates Z present in the population of interest (i.e., z such that P(Z = z) > 0)

$$0 < P(X = 1 | \mathbf{Z} = \mathbf{z}) < 1$$

In the case where the covariates \mathbf{Z} consist of a single variable Z (i.e., Age) we have the following graphical representation of positivity, overlap, and common support:

POSITIVITY - OVERLAP - COMMON SUPPORT



10-15 16-20 21-25 26-30 31-35 36-40 41-45 46-50 51-55 56-60 61-65 66-70 71-75

AGE

adjusting (conditioning)
on more covariates **Z**

could lead to

could lead to

higher chance of satisfying unconfoundedness

higher chance of violating positivity

POSITIVITY — OVERLAP — COMMON SUPPORT

For all values z of covariates Z present in the population of interest (i.e., z such that P(Z = z) > 0)

$$0 < P(X = 1 | \mathbf{Z} = \mathbf{z}) < 1$$

increase the "dimension" of the covariates **Z**



makes the subgroups for any level z of the covariates Z smaller



CURSE OF DIMENSIONALITY

As each subgroup $\mathbf{Z} = \mathbf{z}$ gets smaller, there is a higher and higher chance that either the whole subgroup $\mathbf{Z} = \mathbf{z}$ will have treatment (X = 1) or the whole subgroup $\mathbf{Z} = \mathbf{z}$ will have control (X = 0).

size of any subgroup $\mathbf{Z} = \mathbf{z}$ equal to 1



positivity is guaranteed to not hold

adjusting (conditioning)
on more covariates **Z**

could lead to

higher chance of satisfying unconfoundedness

iates **Z**

could lead to

higher chance of violating positivity



demanding too much from models and getting very bad behavior in return



fit a model to $\mathbb{E}[Y|X, \mathbf{Z}]$ using the available data (x, y, \mathbf{z})

POSITIVITY - OVERLAP - COMMON SUPPORT

For all values z of covariates Z present in the population of interest (i.e., z such that P(Z = z) > 0)

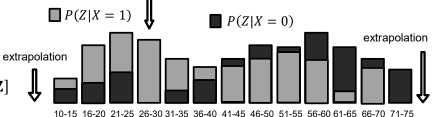
$$0 < P(X = 1 | \mathbf{Z} = \mathbf{z}) < 1$$

inputs to the model $\mathbb{E}[Y|X, \mathbf{Z}]$ are (x, \mathbf{z}) pairs, while the output is the outcome y.

$$\blacksquare \mathbb{E}[Y|X=1,\mathbf{Z}]$$

$$\blacksquare \mathbb{E}[Y|X=0,\mathbf{Z}]$$

$$P(X = 0 | \mathbf{AGE} = \mathbf{26} - \mathbf{30}) = 0$$



AGE

Another assumption is that of **NO INTERFERENCE**.

NO INTERFERENCE

The outcome Y_i of each unit "i" is unaffected by anyone else's treatment X_i , $j \neq i$.

$$Y_i(x_1, x_2, \dots, x_{i-1}, x_i, x_{i+1}, \dots, x_{n-1}, x_n) = Y_i(x_i)$$

This assumption could be violated.

TREATMENT X = "GET A DOG"



OUTCOME Y = "MY HAPPYNESS"



Violations of the no interference assumption are almost sure in network data.

Rather, the outcome Y_i of each unit "i" is only a function of treatment X_i .

We have implicitly made this assumption till now.







Communications of the second o

I hang out more

my friend

— 🚨



The last assumption is **CONSISTENCY**.

CONSISTENCY

If the treatment is X, then the observed outcome Y is the potential outcome under treatment X. Formally,

$$X = x \Longrightarrow Y = Y(x)$$

We could write this also as follows:

$$Y = Y(X)$$

It might seem like consistency is obviously true, but that is not always the case.

$$X = \begin{cases} 1 & \text{"get a dog"} \\ 0 & \text{"do not get a dog"} \end{cases}$$





Y = 1 because I needed an energetic friend





because I got an old and low energy dog

SUTVA

The **S**table **U**nit-**T**reatment **V**alue **A**ssumption (**SUTVA**) is satisfied if unit (individual) i's outcome Y_i is simply a function of unit i's treatment X_i .

SUTVA is a combination of consistency and no interference (and also deterministic potential outcomes).

"no multiple versions of treatment."



Y(1) is not well defined, since it will be 1 or 0, depending on something that is not captured by the treatment X specification.

POTENTIAL OUTCOMES 35

PART III TYING IT ALL TOGETHER

The following assumptions are all needed for solving the problem of causal inference:

CONDITIONAL EXCHANGEABILITY - UNCONFOUNDEDNESS

 $(Y(1), Y(0)) \perp \!\!\! \perp X \mid \mathbf{Z}, \mathbf{Z}$ are the covariate variables.

POSITIVITY - OVERLAP - COMMON SUPPORT

For all values z of covariates Z present in the population of interest (i.e., z such that P(Z = z) > 0)

$$0 < P(X = 1 | \mathbf{Z} = \mathbf{z}) < 1$$

SUTVA

NO INTERFERENCE

The outcome Y_i of each unit "i" is unaffected by anyone else's treatment X_j , $j \neq i$.

$$Y_i(x_1, x_2, \dots, x_{i-1}, x_i, x_{i+1}, \dots, x_{n-1}, x_n) = Y_i(x_i)$$

CONSISTENCY

If the treatment is *X*, then the observed outcome *Y* is the potential outcome under treatment *X*. Formally,

$$X = x \Longrightarrow Y = Y(x)$$

We could write this also as follows:

$$Y = Y(X)$$

ADJUSTMENT FORMULA

Given the assumptions of unconfoundedness, positivity, consistency, and no interference, we can identify the ATE:

$$\mathbb{E}[Y(1) - Y(0)] = \mathbb{E}_{\mathbf{Z}} \big[\mathbb{E}[Y|X=1,\mathbf{Z}] - \mathbb{E}[Y|X=0,\mathbf{Z}] \big]$$

NO INTERFERENCE justifies that the quantity we should be looking at for causal inference is

We now come back to give a formal proof of the **ADJUSTMENT FORMULA**.

 $= \mathbb{E}_{\mathbf{Z}} \big[\mathbb{E}[Y|X=1,\mathbf{Z}] - \mathbb{E}[Y|X=0,\mathbf{Z}] \big]$



$$\mathbb{E}[Y(1) - Y(0)] = \mathbb{E}[Y(1)] - \mathbb{E}[Y(0)]$$

$$= \mathbb{E}_{\mathbf{Z}} [\mathbb{E}[Y(1)|\mathbf{Z}] - \mathbb{E}[Y(0)|\mathbf{Z}]]$$

$$= \mathbb{E}_{\mathbf{Z}} [\mathbb{E}[Y(1)|X = 1, \mathbf{Z}] - \mathbb{E}[Y(0)|X = 0, \mathbf{Z}]]$$

AVERAGE TREATMENT EFFECT - ATE

The average treatment effect (ATE) is obtained by taking an average over the ITEs:

$$\tau \triangleq \mathbb{E}[Y_i(1) - Y_i(0)] = \mathbb{E}[Y(1) - Y(0)]$$

where we recall that the average is over the individuals "i" if $Y_i(x)$ is deterministic.

All of these assumptions tie together give us identifiability of the ATE.

(linearity of expectations)

(law of iterated expectations)

(unconfoundedness and positivity)

(consistency)

We need to introduce some terminology that will help clarify the discussion.

ESTIMAND

An estimand is a quantity that we want to estimate.

ESTIMATE

An approximation of some estimand, which we get using data.

ESTIMATOR

A function that maps a dataset to an estimate of the estimand.

$$\mathbb{E}_{\mathbf{Z}}\big[\mathbb{E}[Y|X=1,\mathbf{Z}] - \mathbb{E}[Y|X=0,\mathbf{Z}]\big]$$

estimand we care about for estimating the ATE

Given an estimand α , we let $\hat{\alpha}$ be its estimate.

ESTIMATION

The process that we use to go from data + estimand to a concrete number is known as estimation.

- CAUSAL ESTIMAND refers to any estimand that contains a potential outcome in it.
- STATISTICAL ESTIMAND refers to any estimand that does not contain a potential outcome in it.

In the following formula

$$\mathbb{E}[Y(1) - Y(0)] = \mathbb{E}_{\mathbf{Z}} \big[\mathbb{E}[Y|X=1, \mathbf{Z}] - \mathbb{E}[Y|X=0, \mathbf{Z}] \big]$$

 $\mathbb{E}[Y(1) - Y(0)]$

is the CAUSAL ESTIMAND that we are interested in

To actually estimate this causal estimand, we must translate it into a **STATISTICAL ESTIMAND**

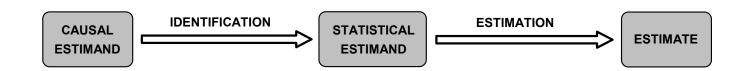
$$\mathbb{E}_{\mathbf{Z}}\big[\mathbb{E}[Y|X=1,\mathbf{Z}] - \mathbb{E}[Y|X=0,\mathbf{Z}]\big]$$

IDENTIFICATION

The process of moving from a causal estimand to an equivalent statistical estimand.

ESTIMATION

The process of moving from a statistical estimand to an estimate.



How do we do when we go to actually estimate quantities such as

$$\mathbb{E}_{\mathbf{Z}}\big[\mathbb{E}[Y|X=1,\mathbf{Z}] - \mathbb{E}[Y|X=0,\mathbf{Z}]\big]$$

We will often use a model (e.g., linear regression or some more flexible predictor from machine learning) in place of the conditional expectations

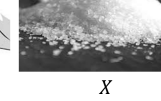
$$\mathbb{E}[Y|X=x,\mathbf{Z}=\mathbf{z}]$$

We will refer to estimators that use models like this as MODEL-ASSISTED ESTIMATORS.

We now need to discuss estimation.

We now give an example complete with estimation(*).





"systolic blood pressure"

"daily sodium intake"

Sodium intake is a continuous variable, so to easily apply

$$\mathbb{E}[Y(1) - Y(0)] = \mathbb{E}_{\mathbf{Z}}[\mathbb{E}[Y|X=1,\mathbf{Z}] - \mathbb{E}[Y|X=0,\mathbf{Z}]]$$

which is specified for binary treatment, we binarize *Y*

$$X = 1$$
 if daily sodium intake ≥ 3.5 gr.

X = 0 if daily sodium intake < 3.5 gr.

We estimate the causal effect of sodium intake on blood pressure.

The data also include the following covariates **Z** for each individual:

- Age
- Amount of protein in the urine

Because we are using data from a simulation, we know that the true ATE of sodium on blood pressure is 1.05.

$$\mathbb{E}[Y(1) - Y(0)] = 1.05$$

* Miguel Angel Luque-Fernandez, Michael Schomaker, Daniel Redondo-Sanchez, Maria Jose Sanchez Perez, Anand Vaidya, and Mireille E Schnitzer. 'Educational Note: Paradoxical collider effect in the analysis of non-communicable disease epidemiological data: a reproducible illustration and web application'. In: International Journal of Epidemiology 48.2 (Dec. 2018), pp. 640–653. doi: 10.1093/ije/dyy275 (cited on pages 16, 45).

How do we actually estimate the ATE?

- We assume consistency, positivity, and unconfoundedness given Z.
 This means that ATE is identified by
- 2) We then take that outer expectation over **Z** = {Age, Amount of proteine} and replace it with an empirical mean over the data, giving us the following:

$$\mathbb{E}[Y(1) - Y(0)] = \mathbb{E}_{\mathbf{Z}} \left[\mathbb{E}[Y|X = 1, \mathbf{Z}] - \mathbb{E}[Y|X = 0, \mathbf{Z}] \right]$$

$$\frac{1}{n} \sum_{i=1}^{n} \left[\mathbb{E}[Y|X = 1, \mathbf{Z} = \mathbf{z}_{i}] - \mathbb{E}[Y|X = 0, \mathbf{Z} = \mathbf{z}_{i}] \right]$$

3) To complete our estimator, we then fit a **MACHINE LEARNING MODEL** to the conditional expectation $\mathbb{E}[Y|x,z]$. We can plug in any machine learning model for $\mathbb{E}[Y|x,z]$, which gives us a **MODEL-ASSISTED ESTIMATOR**.

We use **LINEAR REGRESSION**, which works out nicely since blood pressure is generated as a linear combination of other variables (daily sodium intake, age and amount of protein in the urine).

$$\mathbb{E}[Y(1) - Y(0)] = 0.856$$

$$\%error = \frac{|0.856 - 1.05|}{1.05} \times 100\% = 18\%$$

So, if we use linear regression, which works out nicely since blood pressure is generated as a linear combination of other variables (daily sodium intake, age and amount of protein in the urine).

$$\mathbb{E}[Y(1) - Y(0)] = 0.856$$

$$\%error = \frac{|0.856 - 1.05|}{1.05} \times 100\% = 18\%$$

However, if we were to naively regress Y on only X (daily sodium intake) we would get

$$\mathbb{E}[Y(1) - Y(0)] = 5.37$$

$$\%error = \frac{|5.37 - 1.05|}{1.05} \times 100\% = 411\%$$

All of the above is obtained using the **ADJUSTMENT FORMULA** with **MODEL-ASSISTED ESTIMATION**, where:

- 1) we fit a model **M** for the conditional expectation $\mathbb{E}[Y|x, z]$
- 2) we take an empirical mean over **Z** using model **M**

ADJUSTMENT FORMULA

Given the assumptions of unconfoundedness, positivity, consistency, and no interference, we can identify the ATE:

$$\mathbb{E}[Y(1) - Y(0)] = \mathbb{E}_{\mathbf{Z}} \big[\mathbb{E}[Y|X=1, \mathbf{Z}] - \mathbb{E}[Y|X=0, \mathbf{Z}] \big]$$

Potential Outcomes

Alessio Zanga, Fabio Stella March 27, 2021

```
[1]: %%capture []pip install networkx numpy pandas statsmodels
```

```
[2]: import networkx as nx import numpy as np import pandas as pd import statsmodels.api as sm from typing import Set
```

1 Potential Outcomes

By Alessio Zanga and Fabio Stella

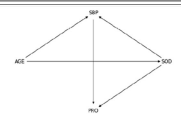
1.1 Abstract

This notebook illustrates a step-by-step example on estimating the average causal effect (ACE) of sodium on blood pressure following Luque-Fernandez et al. (2018). The main focus of this work is to highlight the differences between a naive estimate and the adjusted estimate.

1.2 Introduction

Exceeding the recommendations for 24-h dietary sodium (SOD) intake is associated with increased levels of systolic blood pressure (SBP). Furthermore, with advancing age, the adaptive mechanism responsible for maintaining the composition and volume of the extracellular fluid is compromised. Age is a common cause of both high systolic blood pressure and impaired sodium homeostasis, acting as a confounder. However, high levels of 24-h excretion of urinary protein (PRO) are caused by sustained high SBP and increased dietary SOD, acting as a collider.

```
[3]: G = nx.DiGraph()
G.add_edges_from([("SOD", "SBP"), ("AGE", "SOD"), ("AGE", "SBP"), ("SOD", ...
...,"PRO"), ("SBP", "PRO")])
nx.draw_circular(G, node_size=1000, node_color="white", with_labels=True)
```



Assuming linear relationships between the variables and gaussian noise, a data generation process that is consistent with the represented causal graph follows directly.

```
[4]: def sample_data(size: int = int(1e6), seed: int = 31):

# Set random generator seed for results reproducibility
np.random.seed(seed)

# Sample age with mean 65 and std 5
age = np.random.normal(65, 5, size)

# Sample sodium with additive noise
sod = 0.056 * age + np.random.normal(0, 1, size)

# Binarize sodium following cutoff
sod = (sod 3.55).astype(int)

# Sample systolic blood pressure
sbp = 1.05 * sod + 2 * age + np.random.normal(0, 1, size)

# Sample uniary protein

pro = 0.4 * sod + 0.3 * sbp + np.random.normal(0, 1, size)

# Create a dataframe from sampled variables
return pd.Dataframe(from sampled variables
return pd.Dataframe(from Sampled variables
return pd.Dataframe(from Sampled variables
```

Here, the true ACE is given by the coefficient 1.05 that assign SBP a value given SOD.

```
[5]: data = sample_data()
data.describe()
```

[5]:		AGE	SOD	SBP	PRO
	count	1000000.000000	1000000.000000	1000000.000000	1000000.000000
	mean	65.001134	0.552995	130.583161	39.396720
	std	4.993518	0.497184	10.161539	3.263477
	min	40.964418	0.000000	82.816817	24.421988
	25%	61.627703	0.000000	123.720103	37.190745
	50%	65.009236	1.000000	130.597297	39.402344
	75%	68.372794	1.000000	137.445514	41.602773
	max	87.717317	1.000000	176.708462	54.573313

1.3 Methods

Assuming consistency, positivity, and unconfoundedness, the average causal effect (ACE) is defined:

$$\tau = \mathbb{E}[Y(1) - Y(0)] = \mathbb{E}_{\mathbf{Z}}[\mathbb{E}[Y(1) - Y(0)|\mathbf{Z}]] = \mathbb{E}_{\mathbf{Z}}[\mathbb{E}[Y|X = 1, \mathbf{Z}] - \mathbb{E}[Y|X = 0, \mathbf{Z}]]$$

Replacing expectations with empirical mean:

$$\hat{\tau} = \frac{1}{n} \sum_{i=1}^{n} \mathbb{E}_{Z} [\mathbb{E}[Y|X=1,Z=z_i] - \mathbb{E}[Y|X=0,Z=z_i]]$$

It is possible to choose any machine learning model for $\mathbb{E}[Y|X,Z]$, such as a linear regression model, which works out nicely since blood pressure is generated as a linear combination of other variables.

```
[6]: def ACE(data: pd.DataFrame, X: str, Y: str, Z: Set[str]):
# Define the regression model formula
formula = f"(Y) = (X)"
if len(Z) != 0: formula += "+" +" +" +",join(Z)
# Fit Ordinary Least Square regression model
estimator = sm.OLS.from_formula(formula, data).fit()
# Compute potential outcomes by fixing X
Y1 = estimator.predict(data.assign(**(X: 1)))
Y0 = estimator.predict(data.assign(**(X: 1)))
# Compute average causal effect
return np.mean(Y1 - Y0)
```

1.4 Results

The true ACE is:

The estimated ACE adjusting for AGE and PRO is:

```
[8]: t = ACE(data, X = "SOD", Y = "SBP", Z = ["AGE", "PRO"])
f"Estimated ACE: {t:.3}, Relative Error: {(np.abs((t-ace)/ace*100)):.4}%"
```

[8]: 'Estimated ACE: 0.856, Relative Error: 18.46%'

While the naive estimated ACE without adjustment is:

```
[9]: t = ACE(data, X = "SOD", Y = "SBP", Z = [])
f"Estimated ACE: {t:.3}, Relative Error: {(np.abs((t-ace)/ace*100)):.4}%"
```

[9]: 'Estimated ACE: 5.37, Relative Error: 411.5%'

1.5 Conclusions

Applying a naive regression model without adjustment leads to an estimated ACE which is four times off. A regression model with a valid adjustment set reduce the relative error to only 18%.

The associated notebook is available here

https://colab.research.google.com/github/AlessioZanga/CaMo/blob/develop/examples/potential outcomes.ipynb

COLAB EXECUTABLE