Causality and Directed Acyclic Graphs

Alexander Quispe

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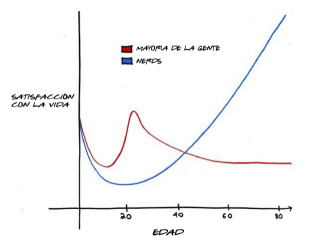
Correlation vs causality

Figure: Fuente link



Correlation vs causality

Figure: Fuente: https://twitter.com/llopatin/status/1540800782198382593



IDENTIFICATION

This lecture is based on notes from "The Effect Book" by Nick Huntington-Klein.

- Where's Your Variation?
- Identification

IDENTIFICATION

- Using theory, paint the most accurate picture possible of what the data-generating process looks like
- Use that data generating process to figure out the reasons our data might look the way it does that don't answer our research question
- Find ways to block out those alternate reasons and so dig out the variation we need

Causality and Directed Acyclic Graphs

- "Directed" because the edges imply a direction: because the arrow from L to A is into A, L may cause A, but not the other way around.
- "Acyclic" because there are no cycles: a variable can not cause itself, either directly or through another variable
- Suppose in our study individuals are randomly assigned to heart transplant A with a probability that depends on the severity of their disease L. Then L is a common cause of A and Y

Directed Acyclic Graphs

- Figure 1 represents an observational study in which we are willing to assume that the assignment of heart transplant A has as parent disease severity L and no other causes of Y. This also represents a conditionally randomized experiment.
- Figure 2 represents a marginally randomized experiment.

- Lets analyse figure 3, which is an observational study.
- A is carrying a lighter, L cigarette smoking and Y is lung cancer.
- We know that carrying a lighter A has no causal effect on lung cancer Y.
- MAIN QUESTION = carrying a lighter A is associated with lung cancer Y?
- how do you attack this research question?

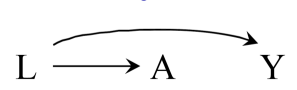
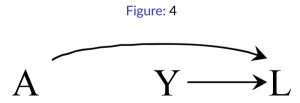


Figure: 3

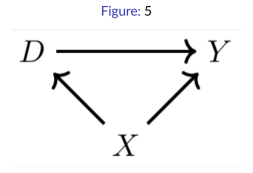
- Having information about the treatment *A* improves our ability to predict the outcome *Y*, even though *A* does not have a causal effect on *Y*.
- The investigator will make a mistake if he concludes that A has a causal effect on Y just because A and Y are associated.

- Suppose you know that certain genetic haplotype A has no causal effect on anyone's risk of becoming a cigarette smoker Y
- Both the haplotype *A* and cigarette smoking *Y* have a causal effect on the risk of heart disease *L*.
- The common effect L is referred to as a **collider** on the path $A \rightarrow L \leftarrow Y$
- Question : A and Y are associated?



- A and Y are independent.
- The knowledge that both A and Y cause heart disease L is irrelevant when considering the association between A and Y.
- Colliders block the flow of association along the path on which they lie. Thus A and Y are independent because the only path between them, $A \to L \leftarrow Y$, is blocked by the collider L.

Cofounder

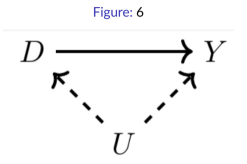


- **Direct Path** = $D \rightarrow Y$, this is causal
- Backdor Path = $D \leftarrow X \rightarrow Y$, this is not causal
- Backdoor Path creates spurious correlations between ${\it D}$ and ${\it Y}$

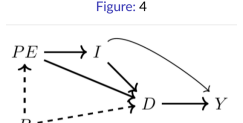
Backdoor Path

- The most important things we can learn from the DAG
- Similar to the notion of omitted variable bias in that it represents a variable that determines the outcome and the treatment variable
- Just as not controlling for a variable like that in a regression creates omitted variable bias, **leaving a backdoor open creates bias**.

Confounder is unobservable



Human capital model - DAG



- *D* be the treatment (e.g., college education)
- Y be the outcome of interest (e.g., earnings).
- *PE* be parental education
- I be family income
- *B* be unobserved background factors, such as genetics, family environment, and mental ability.

Human capital model - DAG

Figure: 8

- 1. D o Y (the causal effect of education on earnings)
- 2. $D \leftarrow I \rightarrow Y$ (backdoor path 1)
- 3. $D \leftarrow PE
 ightarrow I
 ightarrow Y$ (backdoor path 2)
- 4. $D \leftarrow B
 ightarrow PE
 ightarrow I
 ightarrow Y$ (backdoor path 3)

The problem with open backdoor paths is that they create systematic and independent correlations between D and Y

Backdoor criterion

You, as researcher, have the goal of closing these backdoor paths. And if we can close all of the otherwise open backdoor paths, then we can isolate the causal effect of D on Y. **SOLUTIONS**

- you can close that path by **conditioning** on the confounder. **"Controlling for"** the variable in a regression.
- By **not conditioning on a collider**, you will have closed that backdoor path and that takes you closer to your larger ambition to isolate some causal effect.

When all backdoor paths have been closed, we say that you have come up with a research design that satisfies the backdoor criterion.

Backdoor criterion

Figure: 9

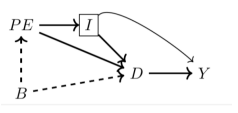


Figure: 10

$$Y_i = lpha + \delta D_i + eta I_i + arepsilon_i$$