# FLS 6441 - Methods III: Explanation and Causation

Week 3 - Field Experiments

Jonathan Phillips

April 2019

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    - ► How much can we learn with better research design?
  - Model-Based Solutions: Not so much.

	Independence of Treatment Assignment?	Researcher Controls Treatment Assignment?
Controlled Experiments	√	$\checkmark$
Natural Ex- periments	✓	
Observational Studies		

		Independence of Treatment Assignment	Researcher Controls Treatment Assignment?
Controlled Experiments	Field Experiments	✓	✓
	Survey and Lab Experiments	✓	√
Natural Experiments	Randomized Natural Experiments	√	
	Instrumental Variables	✓	
	Discontinuities	√	
Observational Studies	Difference-in-Differences		
	Controlling for Confounding		
	Matching		
	Comparative Cases and Process Tracing		

# Section 1 Independence

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- Treatment Assignment Mechanisms that ARE independent of potential outcomes

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  - ▶ We want to estimate:

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 Potential outcomes in the treatment and control groups are now **unbiased** and representative of all the units

(4)(5)

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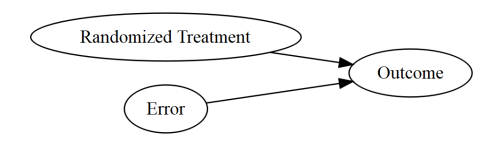
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    - ► No reverse causation is possible

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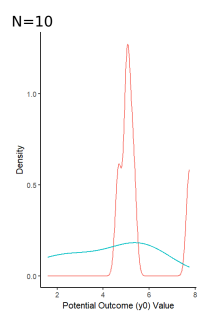
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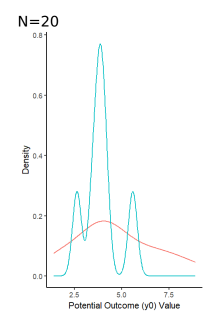
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  - ► We have no way of *verifying* if potential outcomes are biased

 Balance on potential outcomes is unlikely in small samples

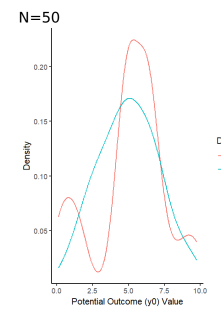
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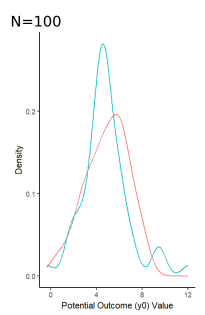
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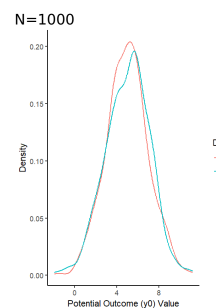
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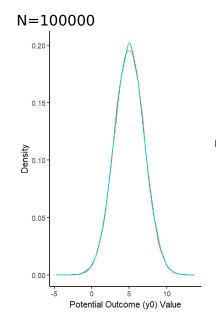
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## Section 2

# **Analysis**

$$A\hat{T}E = E(Y_1|D=1) - E(Y_0|D=0) = E(Y_1) - E(Y_0)$$

▶ If treatment is random we know that:

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▶ What is  $E(Y_1|D=1)$ ?

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  - ► And a simple T-test for statistical significance
  - NO modelling assumptions ("non-parametric")

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Implementation

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So:

$$\hat{\beta} = E(Y_{1i} - Y_{0i})$$

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► Regression Results  $(Y_i = \alpha + \beta D_i + \epsilon_i)$ :

	term	estimate	std.error	statistic	p.value
1	(Intercept)	0.03459	0.07110	0.48647	0.62664
2	treatment	0.27065	0.10044	2.69472	0.00706

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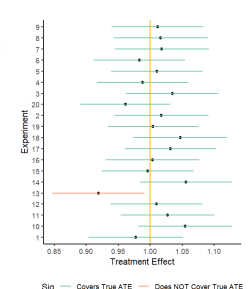
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  - ▶ But there is usually a cost trade-off

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  - ▶ To improve precision, i.e. reduce the standard errors on  $\beta$ 
    - ► The more variation in Y we can explain with covariates, the more certain we can be on the effect of D

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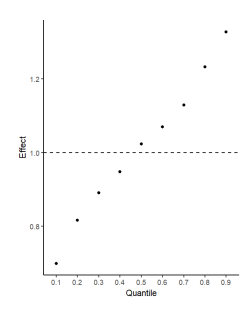
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- ► But we can also estimate Quantile treatment effects, eg. the effect of treatment on the bottom 10% of the distribution

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### Heterogeneous Effects

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- ► Only the health centre was randomly assigned, not neighbourhood income!

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- Analysis: More on how to respond to non-compliance next week

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- ► Check: Normally a difference in means T-test of covariates between treatment and control groups

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- ▶ But we can test observable pre-treatment covariates
- If covariates are the same in the treatment and control groups, this variable cannot explain any differences in outcomes
- ► If lots of variables are balanced, it's likely potential outcomes are too
- ► Check: Normally a difference in means T-test of covariates between treatment and control groups
- Check: Or a Kolmogorov-Smirnov (KS) Test of identical distributions

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- ▶ Why are spillovers a problem?
  - ► **Design:** Limit risk of spillovers, eg. leave 20 miles between each unit in sampling
  - ► Check: Qualitative fieldwork
  - Analysis: Try to measure spillovers

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- ▶ ...Or do we want to measure these additional effects?

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#### **Parallel Treatments**

- Eg. Measurement bias: Researchers give treated units 'the benefit of the doubt' and record higher outcomes for them
- Or Hawthorne Effects: Participants respond to being studied, not treatment (more next week)

► **Design:** Careful specification of treatment and control

# Downstream Consequence of Treatment



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## Parallel Treatment



## Section 4

## **Implementation**

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  - ► Political pressure
  - ▶ We don't want to be guinea pigs!

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- ▶ What's the difference between these three options?
- ▶ What % treated? 50:50 is usually most efficient

- ▶ Blocking
- Randomization is inefficient and risky

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  - ► Blocking means randomizing within fixed groups
  - ► Eg. We have 10 states and a total sample size of 5000 so we fix 250 treated and 250 control in each state
- ▶ "Block what you can; randomize what you cannot"

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- ► Both work in the same way randomization avoids selection (into the data/treatment)

# Section 5

# Critiquing

# Critiquing Field Experiments

► Field experiments are easy to evaluate. What can go wrong??

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- ► We want to learn about generalizable political processes.

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  - The places that agree to field experiments are not representative

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  - 3. **General Equilibrium Effects:** Average test scores went from 70% to 90%, so the exam board readjusted the test and made it harder.

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- ► Treatment could not be scaled (Every village cannot get visits from Columbia professors twice a year)
- And politics was ignored (No implementation unless you give locals responsibility, but then lose control)

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- Selection bias in research findings