# **Causal Inference II**

MIXTAPE SESSION



# Roadmap

Continuous DiD

Dose causal parameter
Identification
Selection bias
Interpreting TWFE

Fuzzy DiD

Concluding remarks

#### Continuous DiD

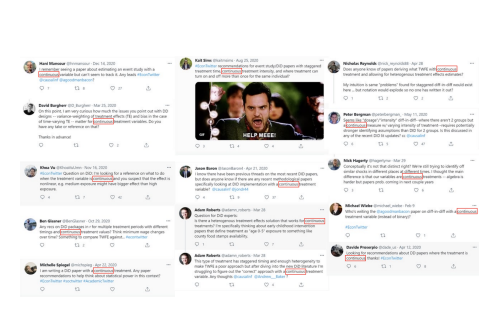
- A very common panel model will use a treatment variable which is continuous, not binary
- Examples include minimum wage papers, my JHR on abortion clinic closures causing increased travel distance, vaccinations, price elasticity of demand etc.
- Variation is in "treatment intensity" and researchers typically use TWFE for estimation, or perhaps count models like Poisson

Quotes

"The two-period regression estimator can be easily modified to allow for continuous, or at least non-binary, treatments." (Wooldridge 2005)

"A second advantage of regression DiD is that it facilitates the study of policies other than those that can be described by a dummy." (Angrist and Pischke 2008)

### Continuous DiD



#### Overview

- 1. What of what we have learned carries forward to the continuous case?
- 2. Some of the problems with continuous (maybe most) don't even have to do with differential timing, so I'm not going to cover it

# Recommended steps of causal projects

- 1. Define the parameter we want ("ATT"),
- 2. Ask what what beliefs do you need ("identification"), and
- 3. Build cranks that produce the correct numbers ("estimator")

People often skip 1 and 2 and go straight to 3 and run regressions then go back and assume exogeneity (step 2), and hope that the estimates are weighted averages of individual treatment effects (1), but that is not guaranteed

# Dangers of skipping 1 and 2

- TWFE was arguably a case where people skipped 1 and 2 and went straight to 3
- We now know that the "constant treatment effect" static specification does not recover the ATT under parallel trends, but the VWATT and requires no dynamics
- We can see this too in simulations even with matching and regression – defining the parameter ahead of time then clearly indicates what assumptions to make, pushing into specifications
- Let's look at this now

# **Data Generating Process**

- Covariate imbalance
- Heterogenous treatment effects with respect to covariates
- Linear data generating process
- Question: How do we estimate the ATE vs the ATT?

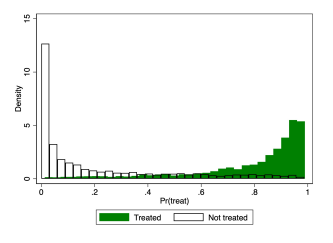
# **Data Generating Process**

- Age is generated from a normal distribution:
  - → Treatment group: mean 25, standard deviation 2.5
  - → Control group: mean 30, standard deviation 3
- GPA is generated from a normal distribution:
  - → Treatment group: mean 1.76, standard deviation 0.5
  - → Control group: mean 2.3, standard deviation 0.75
- Age and GPA are centered around their respective means
- Squared terms and interaction terms are generated:
  - → Age squared (age sq), GPA squared (gpa sq)
  - → Interaction between age and GPA (interaction)

# **Data Generating Process**

- Outcome variables are generated:
  - $\rightarrow$  No treatment (y0):  $15000+10.25\cdot age-10.5\cdot age sq+1000\cdot gpa-10.5\cdot gpa sq+500\cdot interaction <math>+\epsilon$
  - → Treatment (y1):  $y0 + 2500 + 100 \cdot age + 1000 \cdot gpa$
  - $\rightarrow$  Treatment effect (delta): y1 y0
- Average treatment effect (ATE) is estimated at 2500
- Average treatment effect on the treated (ATT) is estimated at 1971

# Covariate imbalance (expressed as propensity score)



# Regression specifications 1 and 2

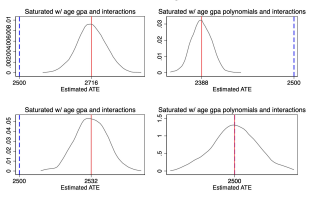
We will look at several different specifications. These first two are standard ways of incorporating covariates. You enter them in linearly as controls.

earnings = 
$$\beta_0 + \beta_1 \text{treat} + \beta_2 \text{age} + \beta_3 \text{gpa} + \epsilon$$
 (1)  
earnings =  $\beta_0 + \beta_1 \text{treat} + \beta_2 \text{age} + \beta_3 \text{age\_sq} + \beta_4 \text{gpa} + \beta_5 \text{gpa\_sq} + \beta_6 (\text{gpa} \times \text{age}) + \epsilon$  (2)

Interpretation focuses on  $\widehat{\beta}_1$ . But what does it mean? Look at top two. Remember ATE is 2500.

# ATE estimates across different specifications

#### OLS Estimates of ATE with heterogenous treatment effects



Four kernel density plots of estimated coefficients from 1000 simulations

# ATT Calculation: Regression 3

Saturated regressions: interact treatment dummy with all covariates.

Regression Specification:

earnings = 
$$\beta_0 + \beta_{1t}$$
treat +  $\beta_{2a}$ age +  $\beta_{3g}$ gpa +  $\beta_{4ta}$ (treat × age) +  $\beta_{5tg}$ (treat × gpa) +  $\beta_{6age}$ (age × gpa) +  $\beta_{6tag}$ (treat × age × gpa) +  $\epsilon$ 

Estimated ATE is the coefficient,  $\widehat{\beta_{1t}}$ , but how do we get the estimated ATT?

Calculating ATT:

$$\mathsf{ATT}_3 = \beta_{1t} + \beta_{4ta} \cdot \mathsf{a\bar{g}e} + \beta_{5tq} \cdot \mathsf{g\bar{p}a} + \beta_{6taq} \cdot \mathsf{a\bar{g}e} \cdot \mathsf{g\bar{p}a}$$

using the means of all covariates

# ATT Calculation: Regression 4

Saturated regressions: interact it with covariates, higher order polynomials, and all interactions

Regression Specification:

earnings 
$$= \beta_0 + \beta_{1t} \operatorname{treat} + \beta_{2a} \operatorname{age} + \beta_{3a_Sq} \operatorname{age} \operatorname{sq} + \beta_{4g} \operatorname{gpa} + \beta_{5g_Sq} \operatorname{gpa} \operatorname{sq} + \\ \beta_{6ta} (\operatorname{treat} \times \operatorname{age}) + \beta_{7ta_Sq} (\operatorname{treat} \times \operatorname{age} \operatorname{sq}) + \beta_{8tg} (\operatorname{treat} \times \operatorname{gpa}) + \\ \beta_{9tg_Sq} (\operatorname{treat} \times \operatorname{gpa} \operatorname{sq}) + \beta_{10ag} (\operatorname{age} \times \operatorname{gpa}) + \beta_{11a_Sqg} (\operatorname{age} \operatorname{sq} \times \operatorname{gpa}) + \\ \beta_{12ag_Sq} (\operatorname{age} \times \operatorname{gpa} \operatorname{sq}) + \beta_{13a_Sqg_Sq} (\operatorname{age} \operatorname{sq} \times \operatorname{gpa} \operatorname{sq}) + \epsilon$$

Estimated ATE is the coefficient,  $\widehat{\beta_{1t}}$ , but how do we get the estimated ATT?

Calculating ATT:

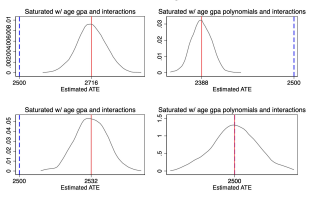
$$\begin{array}{ll} \text{earnings} & = & \beta_0 + \beta_{1t} \text{treat} + \beta_{2a} \text{age} + \beta_{3a\_sq} \text{age\_sq} + \beta_{4g} \text{gpa} + \beta_{5g\_sq} \text{gpa\_sq} + \beta_{6ta} (\text{treat} \times \text{age}) \\ & + \beta_{7ta\_sq} (\text{treat} \times \text{age\_sq}) + \beta_{8tg} (\text{treat} \times \text{gpa}) + \beta_{9tg\_sq} (\text{treat} \times \text{gpa\_sq}) + \beta_{10ag} (\text{age} \times \text{gpa}) \\ & + \beta_{11a\_sqg} (\text{age\_sq} \times \text{gpa}) + \beta_{12ag\_sq} (\text{age} \times \text{gpa\_sq}) + \beta_{13a\_sqg\_sq} (\text{age\_sq} \times \text{gpa\_sq}) + \epsilon \end{array}$$

# Interpretations

- ATE is 2500
- ATT is 1980
- Key point here: the same regression contains both parameters, but only when done correctly, and only when interpreted correctly

# ATE estimates across different specifications

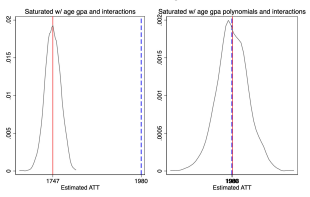
#### OLS Estimates of ATE with heterogenous treatment effects



Four kernel density plots of estimated coefficients from 1000 simulations

### ATT estimates across different specifications

#### OLS Estimates of ATT with heterogenous treatment effects

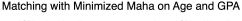


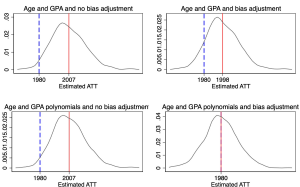
Two kernel density plots of estimated coefficients from two regressions and 1000 simulations

# Matching by minimizing Maha distance metric

- Next I just estimated the ATT using a simpler model nonparametric matching using Abadie and Imbens (2006;2011)
- It's biased when the matches aren't exact, but you can use regression adjustment to estimate the selection bias
- Similar to what augmented synth does
- Much less difficult syntax

# ATT estimates across different matching specifications





Four kernel density plots of estimated ATT from 1000 simulations

# Recommended steps of causal projects

- 1. Define the parameter we want ("ATT"),
- 2. Ask what what beliefs do you need ("identification"), and
- 3. Build cranks that produce the correct numbers ("estimator")

See how when we skep 1 and 2 and go straight to 3, heterogenous treatment effects makes major problems for interpretation? It isn't that regressions can't recover parameters, but you have to saturate when you're attempting to recover ATE or ATT, and even then it's challenging to interpret – and programming, you often don't have code that will do it for you.

# Introducing a new causal parameter

- ATT: Extensive margin causal parameter. Do this versus don't do this.
- **Dose**: Intensive margin causal parameter. Do this much versus this much.

The dose causal parameter will be based on Angrist and Imbens (1995)

#### **Parameters**

### Average treated on the treated

$$ATT(d|d) = E[Y_{it}^d - Y_{it}^0|D_{it} = d]$$

while the treatment, D, can be any amount, d, that amount is technically a particular dose. We raised the minimum wage, but we raised it to a particular wage.

#### **Parameters**

#### Average treated on the treated

$$ATT(d|d) = E[Y_{it}^d - Y_{it}^0|D_{it} = d]$$

This is "the ATT of d for the groups that chose d dosage" which uses as its comparison no dose.

# Average causal response function



Two-stage least squares estimation of average causal effects in models with variable treatment intensity

Authors Joshua D Angrist, Guido W Imbens

Publication date 1995/6/1

Journal Journal of the American statistical Association

Volume 90

Issue 430

Pages 431-442

Publisher Taylor & Francis Group

Description Two-stage least squares (TSLS) is widely used in econometrics to estimate parameters in systems of linear simultaneous equations and to solve problems of omitted-variables bias in single-equation estimation. We show here that TSLS can also be used to estimate the average causal effect of variable treatments such as drug dosage, hours of exam preparation, cigarette smoking, and years of schooling. The average causal effect in which we are interested is a conditional expectation of the difference between the outcomes of the treated and what these outcomes would have been in the absence of treatment. Given mild regularity assumptions, the probability limit of TSLS is a weighted average of per-unit average causal effects along the length of an appropriately defined causal response function. The weighting function is illustrated in an empirical example based on the relationship between schooling and earnings.

Total citations Cited by 1372



Two-stage least squares estimation of average causal effects in models with variable treatment intensity

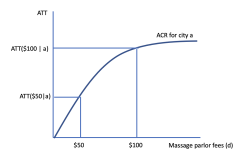
JD Angrist, GW Imbens - Journal of the American statistical Association, 1995 Cited by 1358 Related articles All 14 versions

Average causal response with variable treatment intensity \* J Angrist, G Imbens - 1995 Cited by 16 Related articles All 10 versions

# Angrist and Imbens 1995

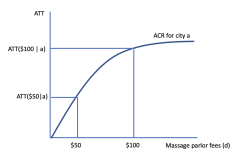
"We refer to the parameter  $\beta$  as the **average causal response (ACR)**. This parameter captures a weighed average causal responses to a unit change in treatment, for those whose treatment status is affected by the instrument. . . . "

# ATT for a given dose



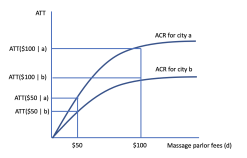
What is the effect of setting fees to \$100 versus nothing at all? It's ATT(\$100|a) for this city.

# ATT for a given dose



Assume city a did choose d=\$100. Then ATT(\$50|a) just means that that is its ATT had it chosen the lower level. The curve, in other words, is tracing out all average causal response for this city.

# ATT for a given dose



What if everyone has different responses? In other words, city a has the higher curve than city b. Then there are several comparisons possible. What is the effect of \$50 on outcomes for cities that actually chose \$50 versus those than actually chose \$100?

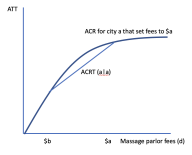
#### Parameters

#### Average treated on the treated

$$ATT(d|d) = E[Y_{it}^d - Y_{it}^0|D_{it} = d]$$

Notice that you are comparing any dose d to no treatment at all – sort of an extensive margin causal response, but that isn't the only causal concept we have. Elasticities are causal, demand curves are causal, but they aren't based on comparisons to nothing – they are intensive margin comparisons, local comparisons, adjacencies. Zero isn't the only counterfactual in other words.

# Average causal response for discrete case vs continuous



#### What is the ACRT?

- ACRT is the causal effect of dose  $D=d_j$  vs a different dose  $D=D_{j-1}$  for group d
  - $\rightarrow$  Easiest example is the demand function: at p=\$10, I buy 10 units, but at p=\$11, I buy 5 units.
  - $\rightarrow$  Causal effect of that one dollar increase is -5 units
  - → Demand curves are pairs of potential outcomes and treatments and equilibrium "selects" one of them
- Discrete/multi-valued treatment is linear difference between two ATTs for the same city
- Continuous treatment is the derivative of the function itself

# Identification for two period set up

- 1. Random sampling.
- 2. No anticipation
- 3. Parallel trends in  $Y^0$  for units of all doses

Identifying ATT(d|d)

We can estimate the ATT(d|d) using the simple DiD equation:

$$E[\Delta Y_{it}|D_i = d] - E[\Delta Y_{it}|D_i = 0]$$

No anticipation and parallel trends converts this comparison of before and after into the  $ATT(d \vert d)$ 

ATT(d|d) is using as its counterfactual the "no treatment", note. Treatment is a dosage compared to zero iow.

# Identifying ACRT

$$ATT(b|b) - ATT(a|a) = (E[\Delta Y_{it}|D_i = a] - E[\Delta Y_{it}|D_i = 0])$$
$$-(E[\Delta Y_{it}|D_i = b] - E[\Delta Y_{it}|D_i = 0])$$
$$= E[\Delta Y_{it}|D_i = a] - E[\Delta Y_{it}|D_i = b]$$

Comparing high and low dose groups.

# Identifying ACRT

$$ATT(d_{j}|d_{j}) - ATT(d_{j-1}|d_{j-1}) =$$

$$(ATT(d_{j}|d_{j}) - ATT(d_{j-1}|d_{j})) + (ATT(d_{j-1}|d_{j}) - ATT(d_{j-1}|d_{j-1})) =$$

$$(ACRT(d_{j}|d_{j})) + (ATT(d_{j-1}|d_{j}) - ATT(d_{j-1}|d_{j-1})) =$$

Part in blue is the movement along the average causal response function, the ACRT, and is causal. The part in red is selection bias.

# Identifying ACRT

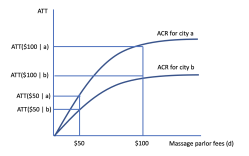
$$ATT(d_{j}|d_{j}) - ATT(d_{j-1}|d_{j-1}) =$$

$$(ATT(d_{j}|d_{j}) - ATT(d_{j-1}|d_{j})) + (ATT(d_{j-1}|d_{j}) - ATT(d_{j-1}|d_{j-1})) =$$

$$(ACRT(d_{j}|d_{j})) + (ATT(d_{j-1}|d_{j}) - ATT(d_{j-1}|d_{j-1})) =$$

Notice parallel trends allows to identify ATT terms but we need additional assumptions for this red part to vanish. We must assume that the ATT for cities that chose  $d_j$  and cities that chose  $d_{j-1}$  are the same had they both chose  $d_{j-1}$ .

## Causality and selection bias



Draw the ACRT for top curve and selection bias.

## Interpreting this

- Unrestricted heterogenous treatment effects (across dosage levels and across units with difference dose response functions) is not itself the problem
- If we randomized dosages, then  $ATT(d_{j-1}|d_j) ATT(d_{j-1}|d_{j-1}) = 0$
- Why? Because then there is no selection on gains from dosages, and average causal response functions are the same for all dosage groups
- So then when is this a problem? Sorting on gains

## Interpreting this

- When estimating treatment effects using continuous DiD, you will need to make one of two assumptions
  - 1. Strong parallel trends: Average change in  ${\cal E}[Y^0]$  for the entire sample is the same as the d group
  - 2. Parallel trends plus homogenous treatment effect functions
- Roy model like sorting on gains typically lead to violations of the second condition insofar as there is heterogenous returns to dosages across units
- So the question you have to ask yourself is do you think that cities are "optimally setting the minimum wage" around some given minimum wage?

## Stronger assumption

- I'm really not so sure I think that when it comes to state legislation that I think a Roy model is likely responsible for the equilibrium
- Solving constrained optimization problems is hard and unlikely is it the case that Florida's ATT and Georgia's ATT are terribly different from one another had both chosen the same minimum wage (but that is the bias)
- But notice where this is taking us we are getting closer and closer back to a place of assuming away the problems!

## Interpretation of estimate

- So we are back to interpretation what is the interpretation, then, of  $\widehat{\delta}^{TWFE}$  with continuous DiD?
- Callaway, Goodman-Bacon and Sant'anna (2022) have a decomposition which I encourage you to read

# TWFE as weighted average of ACRT parameters

$$\hat{\delta}^{TWFE} = \int_{d=L}^{d=U} w_1(l) ACRT(l), dl + w_0 \frac{ATE(d_L)}{d_L}$$

- They assume strong parallel trends and find that TWFE coefficient is weighted average of ACRT function
- All weights are non-negative and sum to one
- But weights are strange in that they are maximized at dosage equal to the mean dosage of the entire dataset

#### New estimator

- There is a new estimator in their new version, but I haven't read it yet
- Today I just wanted to emphasize the basic principles though define the parameter, figure out the assumptions, then build the estimators
- Just running regressions doesn't work once we have heterogenous treatment effects

#### Conclusion

- Very interesting area, a bit challenging, my suggestion is make clear your assumptions
- Most intuitive for me is the parallel trends plus homogenous treatment effects across units around the dosages
- But interpretations are hard because you're having to think about a new parameter, and you could have nonlinearities in the ACRT that since TWFE is a weighted average of them, could flip sign
- Not a problem but could be a challenge for you as you try to make sense of it (particularly given how the weights are)

## Roadmap

Continuous DiD

Dose causal parameter
Identification
Selection bias
Interpreting TWFE

Fuzzy DiD

Concluding remarks

# Sharp DiD

- In a "sharp" DiD, a group gets treated in period 1, a control group does not
- Parallel trends allows you to identify ATT
- We discussed several methods
- But sometimes the lines between treatment and control groups get "fuzzy"

#### **Fuzziness**

- In a "fuzzy" DiD design, there's growth in treatment occurring among units for reasons other than the treatment assignment in the control group
  - → They discuss an early 2000s Duflo paper where Indonesia pushed for more primary schooling
  - ightarrow Used earlier cohorts as controls bc they were already past the age
  - → But they saw growth in schools too
- In many applications, the "treatment rate" increase more in some groups than in others but there is no group that goes from fully untreated to fully treated
- But there is no group that also remains fully untreated

#### **Fuzzy estimators**

 Popular fuzzy estimator (10% of AERs from 2010-2012) divides DiD of the outcome by the DiD of the treatment

$$Wald_{DiD} = \frac{\left(E[Y_k|Post] - E[Y_k|Pre]\right) - \left(E[Y_U|Post] - E[Y_U|Pre]\right)}{\left(E[D_k|Post] - E[D_k|Pre]\right) - \left(E[D_U|Post] - E[D_U|Pre]\right)}$$

- It's Wald IV in that we scale the reduced form by the first stage but they call it Wald DiD
- de Chaisemartin and D'Haultfoeuille (2017) estimates the LATE for groups who go from untreated to treated

## Two proposed estimators

#### Propose two other estimators

- 1. Time corrected Wald ratio,  $Wald_{TC}$  relies on PT within subgroup of units sharing the same treatment at the first date
- 2. Changes in changes extension,  $Wald_{CiC}$  extension of Athey and Imbens (2006) "changes in changes" paper. Generalizes CiC to fuzzy. CiC is invariant to outcome scaling but puts restrictions on the full distribution of potential outcomes instead of the mean

## Personal takeaway

- Two main values of this paper that I found:
  - → Situations where the control group is getting treated with unrelated policy shocks
  - → Continuous treatments
- Code to do it is simple but in Stata

### Most basic notation

For any random variable, R, we interpret as  $R_{dgt}$  as treatment status, treatment group, time

$$R_{101} \sim R|D=1, G=0, T=1$$

Individual treatment status (D) is whether a unit is treated regardless of group; Group (G) is treatment or control *groups*; Time (T) is before or after

Sharp:  $D = G \times T$ ; Fuzzy:  $D \neq G \times T$ 

### Cases under consideration

Case 1: Share of treated units in control don't change between periods

$$E[D_{01}] = E[D_{00}]$$

Wald $_{DiD}$  identifies the LATE parameter for "switchers" (i.e., people whose treatment status changed between 0 and 1) if parallel trends hols and if the ATE of treated units at both dates is stable over time; proposes new estimators that don't depend on this

Stable ATE isn't required in a typical "sharp" DiD

### Cases under consideration

Case 2: Share of treated units changes over time in control

$$E[D_{01}] > E[D_{00}]$$

 $\mathsf{Wald}_{DiD}$  identifies the LATE of switchers under PT and stable ATE assumption and LATE of treatment and control group switchers are the same

Under certain assumptions, their alternative estimator will only be partially identified, and it depends on the size of the change of treated units in the control.

# Fuzzy design assumptions

### A1: Dominating growth of treated units in the treatment group

The treatment group is the one experiencing the larger increase in its treatment rate.

This rules out the case where the two groups experience the same evolution of their treatment rates. Let  $R_{gt} \sim R|G=g, T=t$ ; Assumption 1 implies the following conditions:

$$E(D_{11}) > E(D_{10})$$
  
 $E(D_{11}) - E(D_{10}) > E(D_{01}) - E(D_{00})$ 

# Fuzzy design assumptions

### A2: Stable percent of treated units in the control group

 $0 < E(D_{01}) = E(D_{00}) < 1$  means there is stable percent of treatment units in the control group.

This is a special case where number of treatment units in control group is fixed.

# Fuzzy design assumptions

### A3: Treatment participation equation

In the treatment group, no one switches from treatment to control. Formally this is

$$D=1$$
 if  $V \geq v_{qt}$  with every  $V \perp \!\!\! \perp T | G$ 

Where V is the propensity to get treatment,  $v_{gt}$  is a threshold specific to each group/time

### A little more notation

- We say a unit is treated as  $D(t) = 1\{V \ge v_{at}\}$
- Switchers are units who go from control to treatment between 0 and  $1 S = \{D(0) < D(1), G = 1\}$
- LATE is for switchers:  $\Delta = E(Y_{11}(1) Y_{11}(0))S$
- LQTE is also for switchers:  $\tau_q = f_{y_{11}(1)|S(q)}^{-1} F_{y_{11}(0)|S(q)}^{-1}$

### Switcher LATE/LQTE

### Why only switchers?

- Sometimes only ones affected are switchers; a policy occurs but only eligibility for some. Switchers end up treated
- Identifying more than the LATE places more restrictions and this already has like 8 assumptions

First estimator: Wald $_{DiD}$ 

Commonly used strategy in these fuzzy designs is to normalize the DiD on the outcome by the DiD on the treatment status itself (because remember, in the fuzzy design, units are *becoming* treated as well as *being in treatment groups* 

$$Wald_{DiD} = \frac{DiD_Y}{DiD_D}$$

## Wald-DiD

Let  $S'=\{D(0)\neq D(1), G=0\}$  be control group switchers. Then we define relevant parameters as:

$$\Delta' = E(Y_{01}(1) - Y_{01}(0)|S')$$

$$\alpha = \frac{[P(D_{11} = 1) - P(D_{10} = 1)]}{DiD_D}$$

## Assumptions

#### A4: Parallel trends

Standard assumption. Not worth repeating for the millionth time.

## Assumptions

#### A5: Stable treatment effect over time

In both groups, the average effect of going from 0 to d units of treatment among units with D(0)=d is stable over time. This is the same as assuming that among these units, the mean of Y(d) and Y(0) follow the same evolution over time

$$E\left[Y(d) - Y(0)|G, T = 1, D(0) = d\right] - E\left[Y(d) - Y(0)|G, T = 0, D(0) = d\right] = 0$$

for units in the switching population

## Assumptions

### A6: Homogenous treatment effect over time

Switchers have the same LATE in both groups. This isn't necessary in sharp DiD, just fuzzy

### Wald DiD theorems

There's a reason we just listed six assumptions. We need them for this traditional scaled DiD method for fuzzy designs called the Wald DiD. We'll go in order.

#### Theorem 1: Wald DiD

If A1, A3-A5 hold, then Wald DiD equals

$$\alpha \Delta + (1 - \alpha) \Delta'$$

but if A2 or A6, then Wald DiD equals  $\Delta$ 

# Interpretation of theorem 1: case 1

Case 1: when treatment grows in the control group, then  $\alpha>1$ . Then if we assume A1, A3-A5, a lot of things cancel out under A1, A3-A5, but the Wald DiD becomes a weighted *difference* of the LATEs of treatment and control group switchers in period 1.

Since it is a difference in LATEs, then even two positive LATEs can flip sign if the first is less than the second.

But if you assume A6, you just get the LATE.

Interpreting theorem 1: case 2

Case 2: When treatment diminishes in controls, then  $\alpha < 1$ .

Then under A1, A3-A5, Wald DiD will equal a weighted average of LATEs of treatment and control group switchers in period 1.

This quantity will not reverse signs, but won't equal the LATE without A6.

Interpreting theorem 1: case 3

Case 3: Treatment rate is stable in control, then  $\alpha=1$  and Wald DiD will equal LATE under A1, A3-A5.

This requires that the ATE among units treated at T=0 remain stable over time – necessary condition.

Under A1, A3-A4, Wald DiD is equal to LATE plus a bias term involving several LATEs, and unless they cancel out exactly, Wald DiD will be different from the LATE

#### Alternative estimators

- Wald TC Time Corrected Wald DiD
- Wald CiC Changes in changes generalization to fuzzy design

Now we review alternative assumptions under which Wald TC or Wald CiC identify the LATE of switchers in the fuzzy. First let's look at Wald TC which won't depend on A4-A5.

## Alternative assumptions for the Wald TC

### A4': Conditional parallel trends

This requires Y(0) mean average follow the same trends as all the other groups.

## Wald TC estimator

Wald TC equals

$$\frac{E(Y_{11}) - E(Y_{10} + \delta_{D_{10}})}{E(D_{11} - E(D_{10})}$$

where

$$\delta_d = E[Y_{d_{01}}] - E[Y_{d_{00}}]$$

which is the change in mean outcome between periods 0 and 1 for controls and treatment status d (not groups T and C – individual units d).

#### Theorem 2

#### Theorem 2 and the Wald TC

If A1-A3 and A4', then Wald TC equals  $\Delta$ 

Note that: Wald TC equals

$$\frac{E(Y|G=1, T=1) - E(Y + (1-D)\delta_0 + D\delta_1|G=1, T=0)}{E(D|G=1, T=1) - E(D|G=1, T=0)}$$

This is almost the Wald DiD ratio except for that second term with the  $Y + (1 - D)\delta_0 + D\delta_1$  instead of just Y.

This arises because time can independently affect the outcome.

When treatment is stable for a group G, then  $\delta_0 = 0$ .

#### Comment on Theorem 2

Wald TC equals

$$\frac{E(Y|G=1,T=1) - E(Y+(1-D)\delta_0 + D\delta_1|G=1,T=0)}{E(D|G=1,T=1) - E(D|G=1,T=0)}$$

The numerator of Wald TC compares the mean outcome in the treatment group in the post period 1 to the counterfactual mean we would have had if switchers had remained untreated.

Then normalized by the change in switching, we get the LATE for switchers



Here we have continuous outcomes and an estimator for quantiles of the LATE called LQTE. New assumption is complicated but is needed for the Wald CiC

# Assumptions for changes in changes Wald ratio

### A7: Monotonicity and time invariance of unobservables

Potential outcomes are strictly increasing functions of some scalar unobserved heterogeneity term whose distribution is stationary over time. Also imposes the distribution of that unobserved heterogeneity be stationary within subgroups of units sharing the same treatment status at baseline.

#### Data restrictions

#### A8: Data restrictions

First, Y must have the same support in each of the eight  $D \times G \times T$  cells (common support). Second, the distribution of Y be continuous with positive density in each of the eight cells.

This will allow us to bound treatment effects (Athey and Imbens 2006). Now the ugliest estimator ever.

### Wald CiC estimator

Let  $Q(y)=F_{Y_{01}}^{-1}\cdot F_{Y_{00}}(Y)$  be the quantile-quantile transform of Y from period 0 to 1 in the control group. Also let:

$$F_{CiC,d(Y)} = \frac{P(D_{11} = d)F_{Y_{d11}} - P(D_{10} = d)F_{Y_{d10}}}{P(D_{11} = d) - P(D_{10} = d)}$$

And our Wald CiC estimator is:

$$W_{CiC} = \frac{E(Y_{11}) - E(Q_{D10}(Y_{10}))}{E(D_{11}) - E(D_{10})}$$

Theorem 3: Wald CIC

#### Theorem 3: Wald CiC

Under A1-A3 and A7-A8, then  $W_{CiC}$  is the LATE and equivalently we get the LQTE

$$W_{CiC} = \frac{E(Y|G=1, T=1) - E((1-D)Q_0(Y) + DQ_1(Y)|G=1, T=0)}{E(D|G=1, T=1) - E(D|G=1, T=0)}$$

Comment on theorem 3

Almost the standard Wald DiD except for that  $(1-D)Q_0(Y)+DQ_1(Y)$  instead of Y in the second term of the numerator. So again, we are simply making adjustments for the fuzziness but under different set of assumptions. This term accounts for the fact that time directly affects the outcome, but in a CiC setup.

Which to use

It's about choosing your poison. Do you want A4' or A7?

When T and C have different outcome distributions conditional on D in the first period, then scaling of the outcome may have large effect on the Wald-TC. Whereas Wald-CiC isn't sensitive to the scaling of Y.

But when the two groups have similar outcome distributions conditional on D in the first period, Wald-TC may be preferable as A4' only restricts the mean of the potential outcomes, whereas Wald-CiC restricts the entire distribution

### Extensions to non-binary, ordered treatment

#### Theorem 6

Under continuous treatments, the estimators we've been considering are equal to the average causal response parameter that Angrist and Imbens (1995) discuss. This parameter is a weighted average over all values of d of the effect of increasing treatment from d-1 to d for any switchers where treatment status goes from strictly below to strictly above d over time.

Theorem 6 extends to a continuous treatment. Under theorem 6, each of the estimators is identifying a weighted average of the derivative of potential outcomes with respect to changing  $\boldsymbol{d}$ 

#### Stata code

Only code I know of at the moment is the fuzzydid the authors published in Stata Journal. But it allows you to specify which estimator. Here's sample code for Wald DiD:

fuzzydid lngonf g\_decr post1 inverse\_fee, did breps (1000)
cluster(county1)

where  $g_{d}ecr$  is the treatment group dummy, post1 is the post period dummy, and  $inverse_{f}ee$  is our continuous treatment variable. We specify the Wald DiD by noting did after the comma.

### Concluding remarks

- Paper is hard but worth it. It's possible your controls are getting treated for unrelated reasons, but this is testable
- The Wald DiD is a conventional approach but suffers bias without a layering in of assumptions
- Alternative estimators for when control group stabilization isn't possible or you don't want to impose treatment effect homogeneity are available
- fuzzydid can handle continuous treatments as well as dummies.

### Roadmap

Continuous DiD

Dose causal parameter
Identification
Selection bias
Interpreting TWFE

Fuzzy DiD

Concluding remarks

### Empirical micro model

- In 2014, David Card gave a speech at Michigan in which he told the history of the "empirical micro model", particularly labor
- He notes the different role that economic models have played throughout the history of modern applied (empirical) work
  - 1. Simple approximations of the theoretical model (regress earnings on education)
  - 2. Structural modeling (often highly parametrized models)
  - 3. Princeton (essentially this entire workshop) where focus is on physical (manipulated) treatment assignment
- Models are valuable even under #3, but largely to suggest question, interpret results – not to solve the identification problem

# Unrestricted heterogeneity and selection on gains

- Over time, applied economists (at least the modal one) is far less likely to believe they know enough about treatment assignment or the underlying treatment effects that they are willing to make any theoretical restrictions
- Treatment effects come from unknown production functions, and whether we have heterogeneity across treatment groups involves selection, most likely from a Roy model type of situation
- By and large, this is my conjecture, economists are either unwilling to restrict heterogeneity ex ante, and the only models many of us are by and large committed to are Roy like rational sorting models
- Without restrictions on heterogeneity ex ante, Roy models imply sorting on gains which create the very problems we find

# Consequences of being "deeply a-theoretical"

- Insofar as people sort on the gains from treatment, then given unknown and unrestricted heterogeneous treatment effects, we do not know enough to rule out situations like those in our "baker" dataset
- Baker dataset was extreme, but before this literature, it's not clear anyone would've known it was extreme
- Linear dynamics that never stabilized, large treatment effects in the ATT, heterogenous treatment profiles
- TWFE in the canonical specification performs poorly and in our case flipped sign

### Decision making under uncertainty

- If you cannot restrict heterogeneity, and Roy like models are driving selection into treatment, then there's risks
- TWFE will be efficient if model is correctly specified, which requires
  no dynamic treatment effects, and most of us when pushed cannot
  credibly claim we have enough prior knowledge to say that as we are
  often working on something for which underly production
  technology around treatment effects are not well known or external
  validity is unknown
- You will need to take this seriously, and adjust practices away from naive TWFE modeling
- You are, though, the first wave of researchers to have to, so you are unfortunately caught in a bit of limbo moment

#### Future work

- We are going to cover more differential timing modeling, but I think some of this is a bit redundant
- We start to see similar fixes to the underlying problem performed in different, though internally valid, ways
- Much of this is a bit cloudy to even the most up-to-date practitioner –
  differences between estimators are based on which control groups
  are used, whether PT holds for those control groups, and how best
  to think about the parallel trends assumption (some of which have
  stronger additive forms imposing pre-treatment parallel trends and
  others which do not)
- Cherry picking diff-in-diff could be on the horizon, and this workshop
  is meant to prepare you for this situation as worst case scenario you
  will be reading referee reports of people using methods which you
  may yourself think are inferior but which you still need to understand
- You will need to continue to invest in good judgment, as always;
   most likely going forward returns to econometric skill has shifted up

### Suggestions

- These papers have made the careers of young econometricians to be honest – the growth in citations in short period of time is very weird for econometrics papers (even the IV papers by Angrist and Imbens don't show this pattern)
- Think of a monopolistic competition model entry stops when zero profits at the margin
- Profits are shrinking but not zero at the margin
- There's a lot of activity, it will continue, and you'll need to allocate your time accordingly
- Good luck with your research and pursuit of meaning and happiness
   don't lose sight of what's important