

# Financial Econometrics II – Cross Section and Panel Data

#### Difference-in-differences

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SFI Léman PhD program – 2024, Lecture 3

#### **Overview**



- Motivation
- Difference-in-differences the basics
- Difference-in-differences implementation
- Extensions: continuous treatment & staggered diff-in-diff

## The setting



Recap – if we have a model like

$$y = \beta_0 + \beta_1 x + u$$

and  $cov(x, u) \neq 0$ , then one of the key assumptions for causal inference is violated.

- Another way to think about this is that the distribution of x
  (also after controlling for other covariates) is not random.
- For instance: firms with low leverage may have higher profits because low leverage is more likely for firms with some omitted variable contained in u that is also associated with high profits.

## **Quasi-natural experiments**



- Ideally, the researcher could simply run experiments to achieve the necessary randomness.
  - In medicine, researchers randomly give a new drug to patients to determine the effect of this new drug.
- In corporate finance, we generally cannot do this random assignment.
  - We cannot randomly assign a firm's leverage to determine its effect on profits.
- Therefore, researchers in corporate finance rely on so-called "quasi-natural experiments."

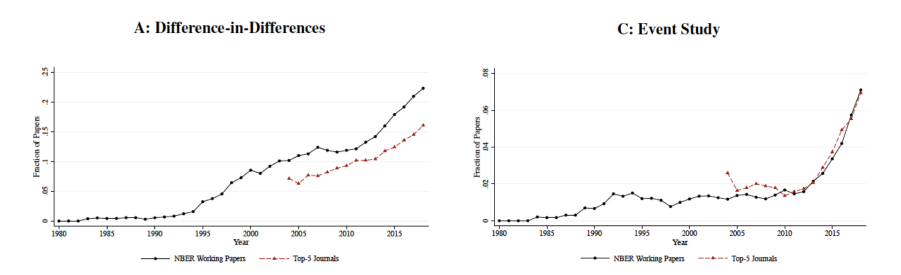
## **Quasi-natural experiments**



- Quasi-natural experiments are events that cause random assignment or a change in a variable of interest x, typically in a panel setting.
  - Some regulatory change (often at state or country level) affects the leverage of a subset of firms.
  - A macro-level financial crisis differentially affects different firms depending on their debt maturity structure pre-crisis.
- Using this type of differential quasi-random treatment to draw causal inferences in a panel setting is very popular, and usually referred to as a difference-in-differences approach.



- Difference-in-differences (or DiD) is an extremely popular approach – probably the main "identification technology" used in recent years
  - also in finance see Goldsmith-Pinkham charts in lecture 1



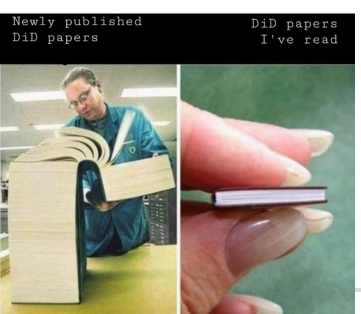
Source: Currie et al. (2020). They note: "Over time, event studies have become almost synonymous
with difference-in-differences: It is now rare to use difference-in-differences without showing an
event study graph, and conversely it is rare to show event studies without a control group."

### Recent developments in DiD



- The popularity of DiD is likely due to the fact that it is very intuitive. And for a while, applied econ/finance researchers could follow a "standard playbook".
- But: over the last 5 years, many new methodological developments in this area. Will mention some of them, although cannot possibly cover them all.





## Recent developments in DiD



- A list of recent surveys (aside from those we will see later):
  - de Chaisemartin, Clément and Xavier D'Haultfoeuille. 2023. <u>Two-way fixed</u> <u>effects and differences-in-differences with heterogeneous treatment effects:</u> <u>A survey.</u> *Econometrics Journal*.
    - These authors are also working on a textbook on DiD draft available at <a href="https://ssrn.com/abstract=4487202">https://ssrn.com/abstract=4487202</a>
  - Miller, Douglas L. 2023. <u>An introductory guide to event study</u> models. In *Journal of Economic Perspectives*.
  - Roth, Jonathan, Pedro H. C. Sant'Anna, Alyssa Bilinski, and John Poe.
     2023. What's trending in difference-in-differences? A synthesis of the recent econometrics literature. Journal of Econometrics.
- For a great 2-hour overview, see the 2023 NBER Methods Lecture by Jesse Shapiro and Liyang Sun:

https://www.nber.org/conferences/si-2023-methods-lectures-linear-panel-event-studies

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## Notation (following Roth et al. 2023)



- Will start with the DiD estimator in the simplest, canonical case binary treatment, two periods (pre/post)
  - example: the famous Card-Krueger (1994) minimum wage study comparing fast-food employment in NJ vs. PA before vs. after NJ increased the minimum wage
- Notation:  $D_i = 0/1$ : treated no/yes; t = 1 before treatm., t = 2 after
- Last lecture, briefly introduced "potential outcomes" useful here:
  - Denote  $Y_{i,t}(0)$  unit is potential outcome if untreated;  $Y_{i,t}(1)$  if treated. Would ideally like Average Treatment Effect:  $E[Y_{i,2}(1) Y_{i,2}(0)]$
  - Observed outcome is  $Y_{i,t} = D_i Y_{i,t}(1) + (1 D_i) Y_{i,t}(0)$
  - DiD can uncover Average Treatment Effect on the Treated (ATT):

$$\tau_2 = E[Y_{i,2}(1) - Y_{i,2}(0) | D_i = 1]$$

## Two key assumptions



- The challenge in estimating  $\tau_2$  is that the untreated potential outcome  $Y_{i,2}(0)$  for the treatment group is never observed
- The central idea behind DiD is that we can use the untreated group to construct this counterfactual outcome
- This requires the "parallel trends" (PT) assumption:

$$E[Y_{i,2}(0) - Y_{i,1}(0) | D_i = 1] = E[Y_{i,2}(0) - Y_{i,1}(0) | D_i = 0]$$

- In words: without treatment, the average outcome for the treated and untreated groups would have evolved in parallel
  - Note: this is about the change across the two periods, not the level
- DGP where this holds:  $Y_{i,t}(0) = \alpha_i + \phi_t + \varepsilon_{it}$  with  $E(\varepsilon_{it}|D_i) = 0$ 
  - Treatment can be related to  $\alpha_i$ , but not to the trend  $\varepsilon_{it}$

## Two key assumptions



Second key assumption is "no anticipation":

$$Y_{i,1}(0) = Y_{i,1}(1)$$
 for all *i* with  $D_i = 1$ 

- In words: in the pre-treatment period, getting subsequently treated does not affect outcome yet.
- What does this get us? Rearrange PT assumption from last slide:

$$E[Y_{i,2}(0)|D_i = 1] = E[Y_{i,1}(0)|D_i = 1] + E[Y_{i,2}(0) - Y_{i,1}(0)|D_i = 0]$$
(by no anticip.) 
$$= E[Y_{i,1}(1)|D_i = 1] + E[Y_{i,2}(0) - Y_{i,1}(0)|D_i = 0]$$

$$\equiv E[Y_{i,1}|D_i = 1] + E[Y_{i,2} - Y_{i,1}|D_i = 0]$$

So then we can identify

$$au_2 = \mathrm{E}[Y_{i,2}(1) - Y_{i,2}(0) \, | D_i = 1]$$

$$= \mathrm{E}[Y_{i,2} - Y_{i,1} \, | D_i = 1] - \mathrm{E}[Y_{i,2} - Y_{i,1} \, | D_i = 0]$$
change for treated change for untreated

#### The difference-in-differences estimator



To estimate

$$\tau_2 = E[Y_{i,2} - Y_{i,1} | D_i = 1] - E[Y_{i,2} - Y_{i,1} | D_i = 0]$$

we use the sample analogue:

$$\widehat{\tau}_2 = (\overline{Y}_{t=2,D=1} - \overline{Y}_{t=1,D=1}) - (\overline{Y}_{t=2,D=0} - \overline{Y}_{t=1,D=0})$$

 Example from Card and Krueger (1994), where NJ was "treated" with an increase in minimum wage:

Variable	Stores by state		
	PA (i)	NJ (ii)	Difference, NJ – PA (iii)
<ol> <li>FTE employment before,</li></ol>	23.33	20.44	-2.89
all available observations	(1.35)	(0.51)	(1.44)
<ol><li>FTE employment after,</li></ol>	21.17	21.03	-0.14 (1.07)
all available observations	(0.94)	(0.52)	
<ol> <li>Change in mean FTE</li></ol>	-2.16	0.59	2.76
employment	(1.25)	(0.54)	(1.36)

## Intuition – single differences



 Consider observing the treated group only, and attempting to estimate the treatment effect as

$$\bar{Y}_{t=2,D=1} - \bar{Y}_{t=1,D=1}$$

- This would "work" only if without the treatment, the expected outcome would have remained unchanged between the two periods.
- Conversely, consider observing only the post-period, and attempting to estimate the treatment effect as

$$\overline{Y}_{t=2,D=1} - \overline{Y}_{t=2,D=0}$$

- This would "work" only if without the treatment, the expected outcome would have been identical across the two groups – implausible unless treatment was fully randomly assigned.
- (See Roberts-Whited for more formal discussion of single-diff cases)



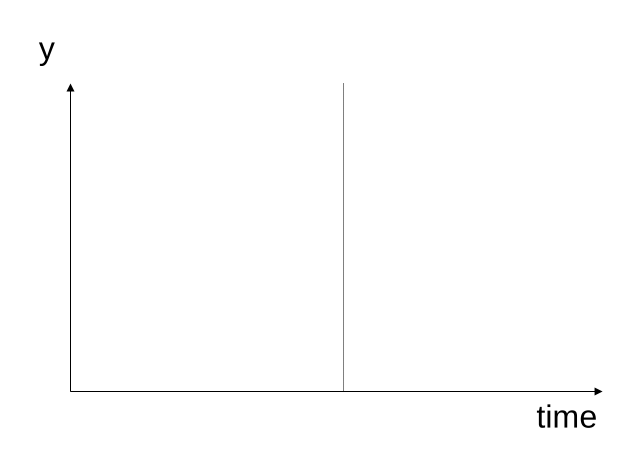
 Rather than calculating means manually, we commonly use the regression version of the difference-in-differences estimator:

$$y_{i,t} = \beta_0 + \beta_1 p_t + \beta_2 d_i + \beta_3 (d_i \times p_t) + u_{i,t}$$

- $p_t$  equals 1 if period t is after treatment, and zero otherwise
- $d_i$  equals 1 if unit i is in treated group, and zero otherwise.
  - $\beta_1$ : measures average change in y due to trends common to both treated and control units
  - $\beta_2$ : measures average difference in level of y between treated and control units in the pre-treatment period.
  - $\beta_3$ : measures the average differential change in y from the pre- to post-treatment period for the treatment group **relative** to the change in y for the control group  $\rightarrow$  can easily be shown to equal  $\tau_2$



$$y_{i,t} = \beta_0 + \beta_1 p_t + \beta_2 d_i + \beta_3 (d_i \times p_t) + u_{i,t}$$





- The difference-in-differences estimator takes care of two major identification threats:
- Any permanent, time-invariant, difference between the treatment and control groups is differenced away by inclusion of the d indicator variable.
- 2. Any common trend affecting both the treatment and control group is also differenced away by inclusion of the p indicator variable.
- Threats to the validity of the diff-in-diff estimator cannot come from permanent differences between the treatment and control groups, or shared trends.

## More on parallel trends

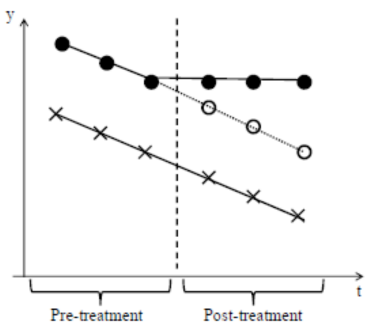


- As seen above, the crucial assumption with diff-in-diff is the "parallel trends" assumption: In the absence of treatment, the average change in the outcome variable would have been the same for both the treatment and control groups.
  - Formally in the regression version: cov(d, u) = cov(p, u) = cov(dp, u) = 0
  - Inherently untestable but can provide supportive evidence (later)
- If have multiple pre-treatment periods: requires trends in the outcomes for the treatment and control groups prior to the treatment to be the same.

### Parallel trends assumption



Figure 1: Difference-in-Differences Intuition



- Realized Avg. Treatment Outcomes
- O Counterfactual Avg. Treatment Outcomes
- Realized Avg. Control Outcomes

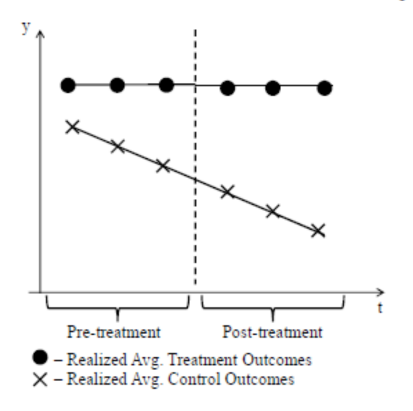
Source: Roberts and Whited (2012)

- Here, both series are trending, but that's not a problem, as long as trends in pretreatment period are parallel
  - "kink" after treatment is what identifies the effect
- Level differences in pre-period are also fine but mean that the estimation may be sensitive to functional form assumptions (e.g. log(y) vs. y – relative vs. absolute changes)
  - also, with trends as depicted, could potentially estimate the model in first differences as well

### Parallel trends assumption



Figure 2: Violation of the Parallel Trends Assumption



Source: Roberts and Whited (2012)

- Pattern depicted here is problematic for standard DiD estimator
  - Will estimate a large treatment effect just due to differential trends
- Absence of parallel pre-trends makes simple DiD estimator essentially "unusable"
  - But will return to this below
- Particular worry: treatment that happens in response to evolution before the treatment

#### **SUTVA**



- Another, "hidden", key assumption is the so-called "Stable Unit Treatment Value Assumption", or SUTVA
- Essentially, it means that a unit's potential outcome is unaffected by the treatment assignment of other units
  - aka "no interference".
  - $Y_{i,t}(D_i)$  does not depend on  $\{D_i, j \neq i\}$
- In corporate finance / banking settings, often not realistic, since firms interact with each other (directly or via market)
  - e.g. law change in one state may affect firms in other states if their products are substitutes
- Violations will lead to biased estimated treatment effects, although can often argue that sign of effect not affected

#### **SUTVA**



- Berg, Reisinger and Streitz (2021) contain thorough discussion of this issue, and advice for researchers
- So far the issue has been mostly ignored in finance settings, but this will likely change going forward
- Always worth keeping in the back of your mind when considering diff-in-diff designs – your life is much easier when you can argue that SUTVA holds
- (Related: discussion of general eqm effects see Nakamura and Steinsson, JEP 2018, for discussion in macro context.)

#### **Overview**

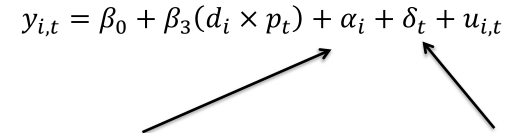


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#### **General diff-in-diff estimator**



In a panel data set, we can add firm and time fixed effects:



Firm fixed effects control for the treatment assignment

Time fixed effects control for trends and subsume  $p_t$ 

- This model may improve precision and fit.
  - Intercept is allowed to vary by firm may matter especially if have unbalanced panel
  - Allows common change in y to vary by time period (e.g. year)

## Adding controls?



- Can easily add additional covariates either to simple DiD equation from earlier, or to more general version
- Why do it? Shouldn't need to if assignment quasi-random
  - reduce error variance more precise estimate
  - "randomization check" showing that controls don't matter
  - or correcting for "conditional randomization"
- Key: controls themselves should not be affected by treatment
  - therefore, should generally only use pre-treatment controls (though in some cases can interact with time)
  - for more discussion, see Huang and Östberg (2023)
- Best practice: always report DiD estimates without controls as well, even if version with controls is your "main" specification

#### Inference in diff-in-diff



- Classic reference is Bertrand, Duflo, Mullainathan (2004).
   They discuss three approaches:
  - Block bootstrap
  - 2. Collapse into pre/post
  - 3. Clustering at the group level most common
  - With small number of clusters, even wild bootstrap may no longer work (e.g. famous Card-Krueger minimum wage study had just two groups: NJ and PA).
  - In such a case, may need to use randomization inference (see Cunningham section 4.2; Hagemann, JoE 2019, and other papers by same author)

## The event study chart



 An extension that is very commonly done, if there are multiple time periods pre- and post-treatment, is the "event study" / "dynamic diff-in-diff":

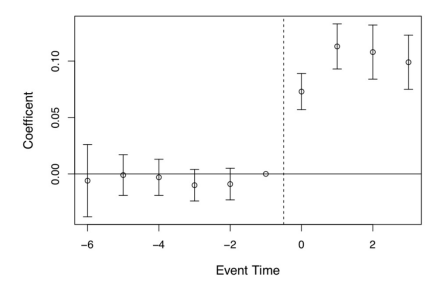
$$y_{i,t} = \alpha_i + \delta_t + \sum_{\ell=-q}^{-2} \gamma_{\ell} \mathbb{I}(t - T_{i0} = \ell) + \sum_{\ell=0}^{m} \lambda_{\ell} \mathbb{I}(t - T_{i0} = \ell) + u_{i,t}$$

- treatment starts at time  $T_{i0}$  (may differ across i)
- the  $\gamma_{\ell}$  coefficients show differential evolution prior to the treatment. Would like those to be close to zero "no pre-trends"
- the  $\lambda_{\ell}$  coefficients show differential evolution after treatment ideally "monotonic" and statistically significant (at least over some period). If effect only happens "late", sheds doubt on validity.
- some flexibility as to which period is omitted (most common: -1, but also see 0 or start of pre-period)

## **Event study chart – examples**



- Cunningham book, Section 9.4.3 discusses study by Miller et al. (QJE 2021) on how expansion of Medicaid in some states affected mortality. Time unit: years.
  - X-axis = "event time" because policy adoption staggered across states: 21 states expanded Medicaid in 2014, 3 states in 2015, 2 states in 2016, and 1 state in 2017. Cf. later



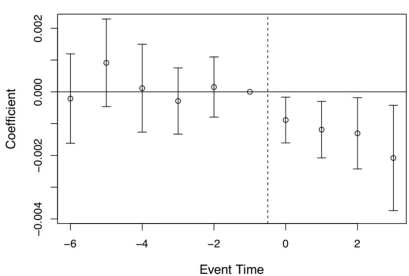


Figure 9.5: Estimates of Medicaid expansion's effects on **coverage** using leads and lags in an event study model. Reprint from Miller et al. (2019).

Figure 9.7: Miller et al. (2019) estimates of Medicaid expansion's effects on on annual mortality using leads and lags in an event study model

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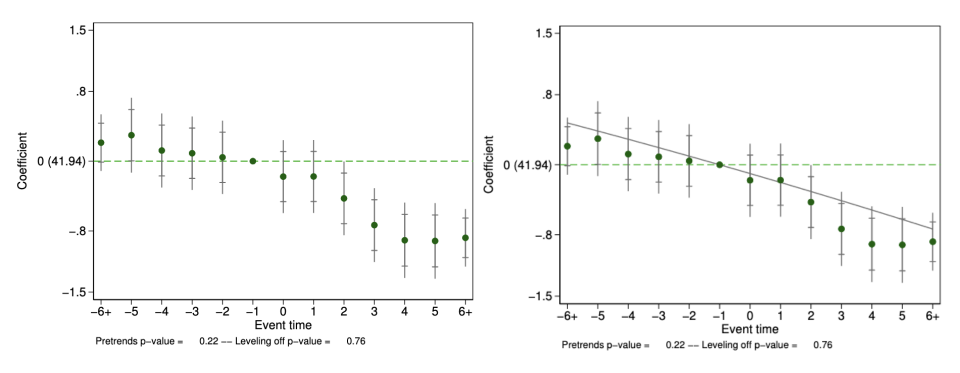
## Event study charts (and estimation) – "state of the art"

- Freyaldenhoven, Hansen, Pérez and Shapiro (2021)
   (<a href="https://scholar.harvard.edu/files/shapiro/files/eventstudy.pdf">https://scholar.harvard.edu/files/shapiro/files/eventstudy.pdf</a>)
   provide various recommendations for making event study
   plots more informative
  - incl. video series on Youtube and package ("xtevent" in Stata or "EventStudyR")
- For instance:
  - include a label for the mean of the pre-period
  - plot "uniform sup-t" confidence bands for the path of the effect, in addition to pointwise confidence intervals
    - see also <a href="https://ryanedmundkessler.github.io/software/">https://ryanedmundkessler.github.io/software/</a>
  - add p-values for "no pre-trends" and "dynamics levelling off"
  - plot path of the "least wiggly" confound whose path cannot be rejected
- Note: they focus on cumulative effects of a policy, which may or may not be what we want to show

## s:fi

# Event study charts (and estimation) – "state of the art"

Examples from Freyaldenhoven et al.:



Charts illustrate an important issue with pre-trend testing: Limited power against alternatives (often do not reject the null of no pre-trend, but would also not reject the null of *some* pre-trend)

## Notes on parallel trends / pre-trends



- Until recently, the consensus was: "no significant pre-trends" = good;
   "significant pre-trends" = you can't run your DiD
- This consensus is starting to shift see Freyaldenhoven et al. (AER 2019)
   & work by Jonathan Roth (<a href="https://jonathandroth.github.io/papers/">https://jonathandroth.github.io/papers/</a>)
- In particular "A More Credible Approach to Parallel Trends" (Rambachan & Roth, 2023) propose tools for robust inference in DiD settings where parallel trends may be violated.
  - E.g. consider restriction that the magnitude of the post-treatment violation of parallel trends can be no larger than a constant  $\overline{M}$  times the maximal pretreatment violation
  - Then, could report e.g. that the conclusion of a positive treatment effect is robust up to the value  $\overline{M} = 2$ .
  - Packages for Stata and R: <a href="https://github.com/mcaceresb/stata-honestdid#honestdid">https://github.com/mcaceresb/stata-honestdid#honestdid</a>

## **Checking validity**



- Other common validity checks that researchers often perform:
- 1. Placebo tests #1: Repeat the diff-in-diff analysis on pre-event years. That is, falsely assume that the onset of the treatment occurs one, two, three years before it actually does. The treatment effect should be statistically indistinguishable from zero.
  - event study chart essentially does that visually
- 2. Placebo tests #2: Make sure that variables that should be unaffected by the event are unaffected by the event. Replace the outcome variable of interest in the empirical model with these other variables.

## **Checking validity**



- 3. Diff-in-diff (aka triple-differences): can be seen as either another placebo (if there are subgroups that should not be affected by treatment) or as a test of mechanisms/ channels (e.g. some firms should be more affected than others)
  - as triple interactions can be hard to interpret, some authors prefer to do sample splits (and run DiD in subsamples)
- 4. Balance on observables between treatment and control groups. Helps to argue for quasi-randomization; otherwise need to add controls.
- 5. Treatment reversal: If there is a reversal of the treatment, it should cause a return to the pre-treatment behavior.

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#### **Extensions of basic framework**



- Two types of extensions of the basic DiD framework are commonly considered – traditionally without much discussion of underlying assumptions, but this has changed recently:
  - 1. Continuous treatment/exposure instead of binary treatment
  - 2. Staggered treatment (rather than single pre/post period)
- In both cases, recent literature has emphasized that our standard TWFE estimators can be problematic if treatment effects are heterogeneous across units or over time (which is often plausible)

#### **Continuous treatment**



- Rather than having treated/untreated, in many settings the treatment is continuous – or different units get varying "doses" of the treatment
  - common when studying outcomes across locations with different shares of firms/households affected by some policy change
- Related to the Bartik IV design we discussed last time, but here we effectively run the "reduced form" only (and the "shifter" is pre/post rather than some aggregate variable)
- Common to just run same DiD regression with  $d_i$  continuous:

$$y_{i,t} = \alpha_i + \delta_t + \beta(d_i \times p_t) + u_{i,t}$$

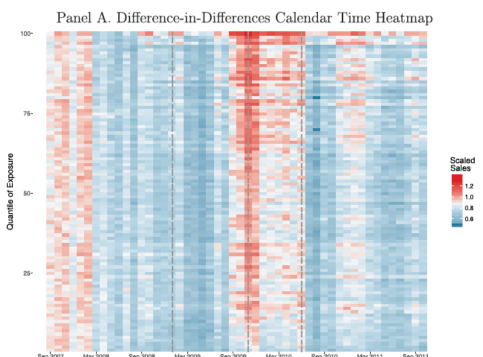
### **Continuous treatment**



- Could also turn into dummies for above/below median exposure
   that's typically done for charts
- But for estimation, appears more efficient to use all the variation
- However, also effectively assume that effect of "treatment dosage" is linear
- Useful to check that it is at least monotonic and approximately linear
  - e.g. if have sufficient data, form dummies for quintiles/deciles and estimate effects for those
- Example: Berger et al. (JF 2020) on effects of first-time homebuyer credit in 2008-10 on home sales across zip codes

## Berger et al. pre-trend & monotonicity "test" S: 1

- Exposure: "We define program exposure based on the number of potential first-time homebuyers in a ZIP code (...) [measured] as the year-2000 share of people in a ZIP code who are first-time homebuyers."
- Effects on home sales over time by centile (b/c 9k ZIP codes):



### **Continuous treatment – caveat**



- Recent work has shown that the TWFE estimator may not perform well in settings with heterogeneous treatment effects.
- In particular, Sun and Shapiro (2022) illustrate that when the treatment effects  $\beta_i$  are unit-specific, the regression

$$y_{i,t} = \alpha_i + \delta_t + \beta(d_i \times p_t) + u_{i,t}$$

may fail to recover a (weighted) average of these unit-specific effects and get a  $\beta$  estimate outside the range of all  $\beta_i$  (!)

- they discuss that having some totally untreated units (i.e.  $d_i=0$ ) can help obtain a better estimator (intuitive this helps to "anchor" the counterfactual)
- see also Callaway, Goodman-Bacon and Sant'Anna (2024) for related discussion of issues with continuous treatments

## Staggered diff-in-diff



- Often in finance, treatment doesn't happen for all units (e.g. states) at the same time – adoption over several years
- Then, "post" varies across units a "staggered" DiD
  - may have "always treated" or "never treated" units
- Very common from Baker, Larcker and Wang (2022, "BLW"):

**Table 1**Use of DiD and Staggered DiD in Finance and Accounting: 2000–2019.

	(1) DiD	(2) Staggered DiD
Journal of Finance	52	30
Journal of Financial Economics	163	85
Review of Financial Studies	138	75
Review of Finance	27	14
Journal of Financial and Quantitative Analysis	51	32
Finance	431	<b>236</b> (559)

## Staggered diff-in-diff



- While researchers approached this essentially in the same way as standard diff-in-diff – with TWFE regressions – a very active literature over the past 5 years or so has pointed out potential issues with such designs
- These occur if treatment effects are heterogeneous either across units or over time.
- Good entry point to the rapidly growing literature: BLW (2022), "How much should we trust staggered DiD?" (plus survey papers listed earlier, esp. Roth et al.)
  - https://asjadnaqvi.github.io/DiD/ provides links to various packages in Stata and R

## **BLW** simulations – illustrating when there is a problem



— 1998 — 2007

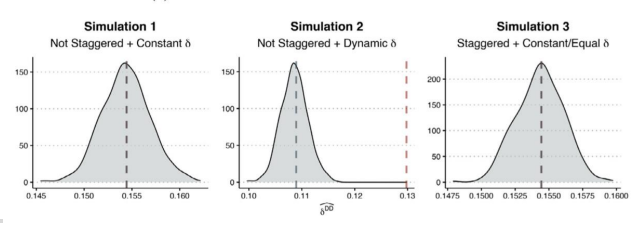
In these cases, TWFE DID does "fine":

(see paper for details about the simulations)

#### Simulation 1 Simulation 2 Simulation 3 Not Staggered + Constant δ Not Staggered + Dynamic δ Staggered + Constant/Equal δ 0.2 ROA 0.0 2000 1990 2010 1980 1990 2000 2010 1980 1990 2000 2010

(i) Trends in Outcome Path

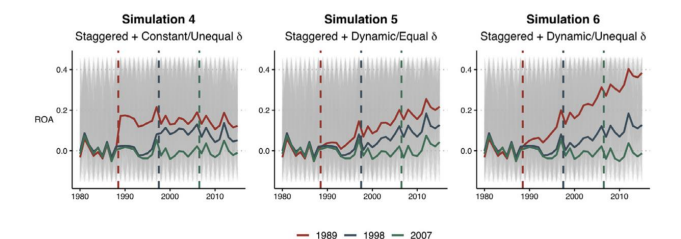
#### (ii) TWFE DiD Estimates on Simulated Data



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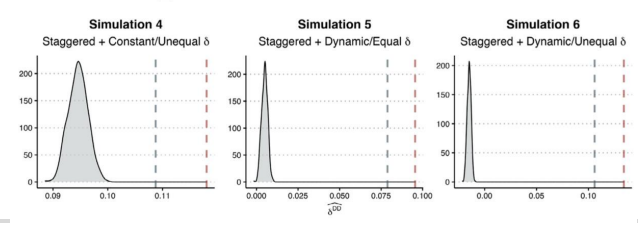
# BLW simulations – illustrating when there is a problem

In these cases, TWFE DiD does much less fine – esp. Sim. 6, where the sign flips!



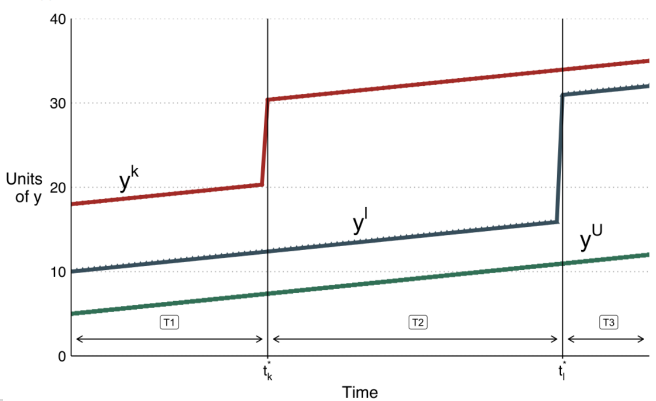
(i) Trends in Outcome Path

#### (ii) TWFE DiD Estimates on Simulated Data



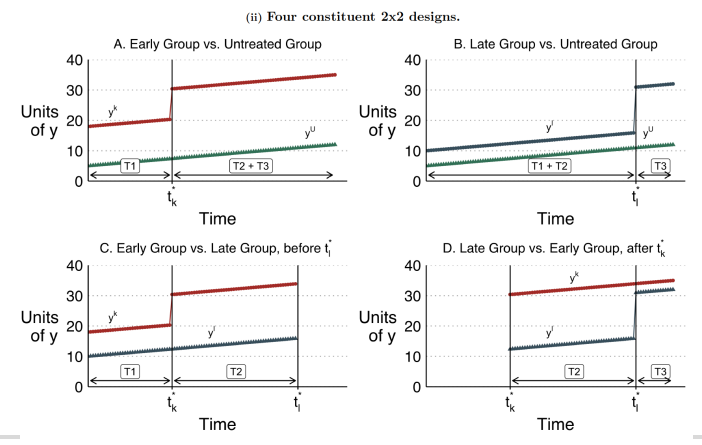
# What's the problem? Goodman-Bacon (2021) $S:\Pi$ decomposition

- Stylized setting with three groups TWFE DiD is a weighted average of all possible two-group/two-period DiD estimators
  - (i) Staggered treatment setting with three treatment groups.



# What's the problem? Goodman-Bacon (2021) $S:\Pi$ decomposition

 Stylized setting with three groups – TWFE DiD is a weighted average of all possible two-group/two-period DiD estimators



## Late vs. early

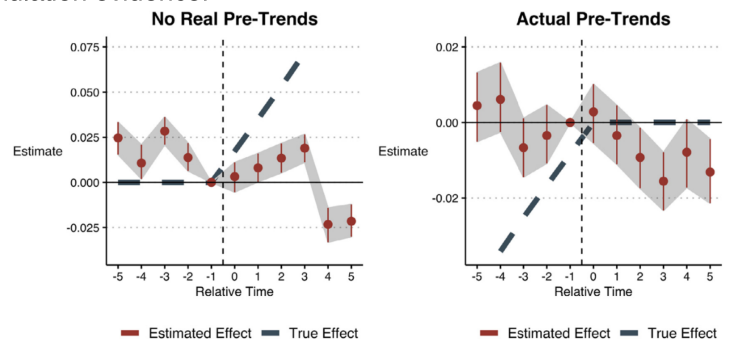


- What's the problem with Simulation 6? No untreated; "late vs. early" (constituent D) gets most weight, and "looks negative" due to weaker (but positive!) treatment effect
  - BLW: early-treated units as effective controls = "potentially problematic"
- What determines weights?
  - absolute size of the subsample
  - relative size of the treatment and effective comparison group in the subsample
  - timing of the treatment in the subsample
  - magnitude of treatment variance in the subsample
- Changing the panel length alone can change the staggered DiD estimate, even when each 2x2 DiD estimate is held constant.

## **Event study plots**



- With staggering and treatment effect heterogeneity, standard event study plots can also be dramatically "off" – BLW Section 3.2
- Simulation evidence:



Binning (e.g. <=-5, >=5) can have major effects on estimated path

## **Diagnostics & alternatives**



- Goodman-Bacon (2021) proposes a useful diagnostic, namely the weight & average DiD of each of the 4 comparison types
- Then, different new estimators have been proposed, which effectively don't use early-treated units as controls for latetreated units – BLW section 4:
  - Callaway and Sant'Anna (2021) BLW's recommendation
  - Sun and Abraham (2021)
  - Stacked Regression (used e.g. in Cengiz et al., 2019)
  - There are other alternatives see next slide.
- BLW apply these methods to two published finance papers, finding in each case that results are not robust to the alternative estimators. (Though not clear how common that is.)

## Imputation estimators



- Another type of alternative estimators are so-called "regression imputation" estimators – e.g. Borusyak et al. (2024) and Gardner et al. (2024)
- Basic idea very intuitive 2-stage approach:
  - 1. Regress outcomes on unit and time fixed effects using only the subsample of untreated observations (incl. not-yet-treated)
  - 2. Based on this regression, impute a counterfactual  $\hat{Y}_{i,t}(0)$  and estimate treatment effects relative to that
  - this yields unit-specific treatment effects that can then be aggregated to the ATT. See e.g. "did2s" package in Stata and R.
- Roth et al. (2023, sect. 3.3) compare assumptions underlying this estimator vs. Callaway-Sant'Anna and provide some guidance on which to use; see also Harmon (2024).
  - relative efficiency depends on serial correlation in errors

## **Event study plots with new estimators**



- A final thing to note is that the event study estimators from these new estimators are constructed differently from the TWFE event studies we are used to
  - illustrated for a particularly stark case in <a href="https://www.jonathandroth.com/assets/files/HetEventStudies.pdf">https://www.jonathandroth.com/assets/files/HetEventStudies.pdf</a>
- What is particularly "confusing" is that the imputation estimators
  à la Borusyak et al. (2024) do not have an omitted base period
  - and in fact the pre- and post-treatment paths should not directly be compared
- Make sure to understand the plots generated by the method(s) you use (often depend on options chosen in the relevant software packages)

## **Summary and conclusion**



- DiD is an extremely popular and intuitive methodology
- But methodological details have come under increased scrutiny in recent years, and this will certainly continue for a while – "new standards" will form.
  - see e.g. BLW Section 6 or Roth et al. (2023) for a set of recommendations, but of course not the final word.
- Life is much easier with "standard" DiD than staggered DiD
  - on the other hand, there are many solutions available to the issues pointed out in the emerging literature – may get credit for applying correctly.
  - certainly many existing staggered DiD studies will be revisited given the new methodologies

## **Summary and conclusion**



- A central issue in DiD is the parallel trends assumption untreated (or later-treated) units provide the counterfactual for treated units
  - Another very active area of research
- If you don't seem to have parallel trends between treated and control units, one option is to re-weight control units to match pre-trends and/or characteristics of treated units – this is the synthetic control approach (see Abadie, JEL 2021 and NBER methods lectures 2021 for a recent overview)
  - useful in particular if only have one treated unit
  - won't cover here but worth learning about! Example of a published paper that uses it is Zevelev (RFS 2021)