Frontier Topics in Empirical Economics: Week 9 Causal Inference with Panel Data II

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- Up until now, we consider a plain vanilla DID setting
- Some policy is implemented at time t_0 in one set of units (treated group), but not the other set of units (untreated group)
- Let's go to more general case of TWFE ⇒ Staggered DID
- Policy can be rolled out in different places at different time

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- \blacksquare Assume we have policy implemented at group g level, rolling out in different periods
- \blacksquare Let's run the same TWFE regression for individual i in group g at time t

$$Y_{igt} = \gamma_g + \lambda_t + \delta D_{gt} + \epsilon_{igt} \tag{1}_{igt}$$

- $D_{gt} = 1$ if group g is treated at time t
- \blacksquare In homogeneous treatment effect case: δ is TE
- In heterogeneous treatment effect case: δ is a weighted sum of ATE in each group and period
- But, is this in general a good estimator? NO!

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- TWFE identifies a weighted average TE
- But some of the weights can be negative
- Thus, the weighted average may be negative even if signs of all cell TEs are positive
- Let's see why (Please read CD(2020), this is important!

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- \blacksquare Three-level of data: individual i, group g, period t
- \blacksquare $N_{g,t}$ is the number of observations in cell (g,t), N is the total number of samplesses
- \blacksquare Assume $D_{i,g,t}$ is a binary treatment, $Y_{i,g,t}(0)$ and $Y_{i,g,t}(1)$ are potential outcomes
- We have (g,t) cell-level average variables as:

$$D_{g,t} = \frac{1}{N_{g,t}} \sum_{i \in (g,t)} D_{i,g,t}, \quad Y_{g,t}(0) = \frac{1}{N_{g,t}} \sum_{i \in (g,t)} Y_{i,g,t}(0)$$
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- Assumption 1: For all (g,t), $N_{g,t} > 0$ (Full Support)
- Assumption 2: For all (g, t), $D_{i,g,t} = D_{g,t}$ (Sharp Design) Treatment is identical for everyone in the same group-time ce
- Assumption 3: Vectors $(Y_{g,t}(0), Y_{g,t}(1), D_{g,t})_{1 \le t \le T}$ are mutually independent No correlation across groups (correlations within group across time is allowed) Weaker version of iid

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Assumption 4:

For all
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, $E(Y_{g,t}(0)-Y_{g,t-1}(0)|D_{g,1},...,D_{g,T})=E(Y_{g,t}(0)-Y_{g,t-1}(0))$

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- In a more general case than staggered DID, exit is also allowed
- \blacksquare Some groups can cancel the policy after some periods, then we have $D_{g,t}=1, D_{g,t+1}=0$
- Let's run the TWFE regression:

$$Y_{igt} = \gamma_g + \lambda_t + \beta^{fe} D_{gt} + \epsilon_{igt}$$

- Assumption 5: For $t \ge 2$, $E(Y_{g,t}(0) Y_{g,t-1}(0))$ does not vary across g Common trends
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■ ATT:
$$\delta^{TR} = E[Y_{i,g,t}(1) - Y_{i,g,t}(0) | D_{g,t} = 1]$$

■ Cell average TE:
$$\Delta_{g,t} = \frac{1}{N_{g,t}} \sum_{i \in (g,t)} [Y_{i,g,t}(1) - Y_{i,g,t}(0)]$$

■ Thus, we have ATT to be an expected weighted average of cell averages

$$\delta^{TR} = E\left[\sum_{(g,t):D_{g,t}=1} \frac{N_{g,t}}{N_1} \Delta_{g,t}\right]$$
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• Let $\hat{\beta}^{fe}$ be the TWFE estimator and $\beta^{fe} = E(\hat{\beta}^{fe})$

Let $\epsilon_{g,t}$ denote the residual of the regression of $D_{g,t}$ on group and period FE

$$D_{g,t} = \alpha + \gamma_g + \lambda_t + \epsilon_{g,t} \tag{3}$$

Theorem 1 in CD(2020)

If we have Assumption 1-5, ther

$$\beta^{fe} = E\left[\sum_{(g,t):D_{g,t}=1} \frac{N_{g,t}}{N_1} w_{g,t} \Delta_{g,t}\right]$$

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- How to interpret this weight $w_{g,t}$?
- You assign more weights to cells (g,t) deviating from the average treatment level and all cells in group g
- If everyone in this group, or every one in this year is not treated, but you are treated, then you will be assigned a large weight
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- Consider a simple case with 2 groups and 3 periods with equal group size
- Group 1 gets treated at periods 3; group 2 gets treated at periods 2 and 3

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	t=1	t=2	t=3
G1	×	×	√
G2	×	$\sqrt{}$	$\sqrt{}$

- Assume that $N_{g,t}/N_{g,t-1}$ does not vary across g (balanced panel)
- Consider a simple case with 2 groups and 3 periods with equal group size
- Group 1 gets treated at periods 3; group 2 gets treated at periods 2 and 3

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$$\text{Let } D_{g,..} = \sum_{t} \frac{N_{g,t}}{N_g} D_{g,t}, D_{..,t} = \sum_{g} \frac{N_{g,t}}{N_t} D_{g,t}, D_{...} = \sum_{(g,t)} \frac{N_{g,t}}{N} D_{g,t};$$

$$\epsilon_{g,t} = D_{g,t} - D_{g,..} - D_{.,t} + D_{...}$$

$$(4)$$

Thus, we have

$$\epsilon_{1,3} = 1 - \frac{1}{3} - 1 + \frac{1}{2} = \frac{1}{6}$$

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 $\epsilon_{2,3} < 0!!!$



In this special case, we have

$$\beta^{fe} = \frac{1}{2}E[\Delta_{1,3}] + E[\Delta_{2,2}] - \frac{1}{2}E[\Delta_{2,3}]$$

- We assign negative weight to $\Delta_{2,3}$
- Negative weight can make results weird
- If $E[\Delta_{1,3}] = E[\Delta_{2,2}] = 1$, $E[\Delta_{2,3}] = 4$, we have

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CD(2020): Negative Weights

- Let's see in more details why there is negative weight
- In this case, we have two switches: Group 1 at period 3, and group 2 at period 2
- Thus, we have two DID comparisons
- \blacksquare It can be proved that β^{re} is the average of these two

$$DID_{1} = E(Y_{2,2}) - E(Y_{2,1}) - [E(Y_{1,2}) - E(Y_{1,1})]$$

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• We can show that $DID_2 = E[\Delta_{1,3}] - (E[\Delta_{2,3}] - E[\Delta_{2,2}])$, NOT $DID_2 = E[\Delta_{1,3}]$

Proof

$$DID_{2} = E(Y_{1,3}) - E(Y_{1,2}) - [E(Y_{2,3}) - E(Y_{2,2})]$$

$$= E(Y_{1,3}(1)) - E(Y_{1,2}(0)) - [E(Y_{2,3}(1)) - E(Y_{2,2}(1))]$$

$$= E(Y_{1,3}(1)) - E(Y_{1,3}(0)) + E(Y_{1,3}(0)) - E(Y_{1,2}(0)) - [E(Y_{2,3}(1)) - E(Y_{2,2}(1))]$$
You cannot cancel this in a forbidden comparison!
$$= E[\Delta_{1,3}] + E(Y_{1,3}(0)) - E(Y_{1,2}(0))$$

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$$DID_2 = E[\Delta_{1,3}] - \underbrace{(E[\Delta_{2,3}] - E[\Delta_{2,2}])}_{\text{bias term}}$$

- Then, $E[\Delta_{2,3}]$ enters into β^{re} with a negative weight
- What does this equation mean?
- lacksquare It means that this DID comparison, is ATE in group 1 period 3, minus a bias term
- Bias term: changes in group 2's ATE between period 2 and 3 and 3
- \blacksquare You are using treated cells as the "control" group!! $\Leftarrow [E(Y_{2,3}) E(Y_{2,2})]$
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- In general, which kind of cells are more likely to have negative weights's
- Let's go back to the function of weight

$$D_{g,t} = \alpha + \gamma_g + \lambda_t + \epsilon_{g,t} \tag{5}$$

- When will $\epsilon_{g,t}$ become negative?
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- a (1) Compute the weights, see whether some of them are negativees.
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- Step 2: If you have many negative weights, or the threshold value of the flippec sign is small, using a new estimator
- \blacksquare CD(2020) constructs a new estimator for TWFE regression, called DID_M
- We define a new average treatment effect

$$\delta^{s} = E\left[\frac{1}{N_{s}} \sum_{(i,g,t): t \geq 2, D_{g,t} \neq D_{g,t-1}} [Y_{i,g,t}(1) - Y_{i,g,t}(0)]\right]$$

- $N_s = \sum_{(g,t): t \ge 2, D_{g,t} \ne D_{g,t-1}} N_{g,t}$, number of obs changing their treatment status from t-1 to t
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- Assumption 9: Strong Exogeneity for Y(1) (corresponding to A4) For all (g,t), $E(Y_{g,t}(1)-Y_{g,t-1}(1)|D_{g,1},...D_{g,T})=E(Y_{g,t}(1)-Y_{g,t-1}(1))$
- Assumption 10: Common Trends for Y(1) (corresponding to A5) For $t \ge 2$, $E(Y_{g,t}(1) Y_{g,t-1}(1))$ does not vary across g
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- Assumption 11: Existence of "Stable" Groups (existence of control groups)
 - * (i) If there is g such that $D_{g,t-1}=0$, $D_{g,s}=1$, there exists g' such that
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 - a (ii) If there is g such that $D_{g,n-1}=1, D_{g,n}=0$, there exists g such that $D_{g,n-1}=D_{g,n}=1$
- Assumption 12: Mean Independence between a group's outcome and other groups' treatment (No spillover)
 - For all g, t, $E(Y_{g,t}(0)|D) = E(Y_{g,t}(0)|D_g), E(Y_{g,t}(1)|D) = E(Y_{g,t}(1)|D_g)$

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- Let's define the DID_M estimator
- Let $N_{d,d',t} = \sum_{g:D_g,t=d,D_g,t-1=d'} N_g,t$, that is, number of obs with treatment d at t and d' at t-1
- Let's define two parts of DID comparisons

$$DID_{+,t} = \sum_{g:D_{g,t}=1,D_{g,t-1}=0} \frac{N_{g,t}}{N_{1,0,t}} (Y_{g,t} - Y_{g,t-1}) - \sum_{g:D_{g,t}=D_{g,t-1}=0} \frac{N_{g,t}}{N_{0,0,t}} (Y_{g,t} - Y_{g,t-1})$$

$$DID_{-,t} = \sum_{g:D_{g,t}=1} \frac{N_{g,t}}{N_{1,1,t}} (Y_{g,t} - Y_{g,t-1}) - \sum_{g:D_{g,t}=1} \frac{N_{g,t}}{N_{0,1,t}} (Y_{g,t} - Y_{g,t-1})$$

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$$DID_{M} = \sum_{t=2}^{T} \left(\frac{N_{1,0,t}}{N_{s}} DID_{+,t} + \frac{N_{0,1,t}}{N_{s}} DID_{-,t} \right)$$
 (6)

Theorem 3 in CD(2020)

If we have Assumptions 1,2,4,5, and 9-12 ther

$$E[DID_M] = \delta^3$$

The DID_M estimator is a weighted average of joiners' and leavers' treatment effect. It is an unbiased estimator of δ^s , that is, the ATE of all switching cells.

 \blacksquare *DID_M* estimator is defined as

$$DID_{M} = \sum_{t=2}^{T} \left(\frac{N_{1,0,t}}{N_{s}} DID_{+,t} + \frac{N_{0,1,t}}{N_{s}} DID_{-,t} \right)$$
 (6)

Theorem 3 in CD(2020)

If we have Assumptions 1,2,4,5, and 9-12 then

$$E[DID_M] = \delta^s$$

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Homework 2: Will plain vanilla DID (like Card and Krueger (1994)) suffer from the same issue when we use TWFE estimator? Explain your answer in words. (Do not use math)

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- Define two types of causal effects: level effect (d vs 0) and slope effect (d vs d'
- Consider a simple two-period DID case
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- We have two periods, t and t-1
- Units receive a treatment dose D_i in t, not t-1 ($D_{it-1}=0$)
- Potential outcome of individual i at time s receiving d is $Y_{is}(d)$
- Assumption 1: We have i.i.d. samples.
- Assumption 2: Full support of D,

$$D = \{0\} \cup D_+.P(D = 0) > 0, dF_D(d) > 0, \forall d \in D_+.$$
 No units are treated in $t - 1$

- Assumption 3: No anticipation effect. $Y_{it-1} = Y_{it-1}(0), Y_{it} = Y_{it}(D_i)$
- Assumption 4: Continuous treatment

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- The definition of causal effect can be much more complicated in continuous treatment case
- Since you are not only comparing d and 0, but also d and d
- 1. Level effect: $Y_t(d) Y_t(0)$ Difference between effect of some dose level d and no treatment
- 2. Slope effect: $Y'_t(d)$ The derivative of the potential outcome function. The marginal increase in the effect when dose is increased.

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$$ATT(a|b) = E[Y_t(a) - Y_t(0)|D = b], \quad ATE(d) = E[Y_t(d) - Y_t(0)]$$

- \blacksquare ATT(a|b): Average effect of dose a on units that who actually experience dose l
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- We call them Average Causal Response Function
- ACRT(d|d): Average causal response of a small change in dose d, for the group of units who actually experience dose d (similar to ATT)
- \blacksquare What is the impact for people who get dose d to get a little bit more doses
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Assumption 4: Parallel Trends.

$$\forall d, E[Y_t(0) - Y_{t-1}(0)|D = d] = E[Y_t(0) - Y_{t-1}(0)|D = 0]$$

■ It says that the path of untreated potential outcomes would have been the same for untreated group and treated group with any dose level

Under Assumptions 1 to 4, ATT
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 is identified for all $d \in D$:

$$ATT(d|d) = E[\Delta Y_t|D = d] - E[\Delta Y_t|D = 0]$$

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Under Assumptions 1 to 4, generally, ACRT(d|d) is NOT identified

$$\frac{\partial E[\Delta Y_t | D = d]}{\partial d} = ACRT(d|d) + \underbrace{\frac{\partial ATT(d|l)}{\partial l}}_{\text{Selection bias}}\Big|_{l=c}$$

- Under traditional parallel trend assumption, local comparisons of paths of outcomes mix ACRT(d|d) and a selection bias term
- The bias is the marginal change in ATT of group D = I if they get dose d
- ACRT CANNOT be identified with traditional parallel trend assumption in a DID fashion! Why?

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- For ACRT(d|d), you consider a marginal increase from d to I
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- We assume parallel trends only for group D = d and group D = l if they are not treated $(Y_t(0) Y_{t-1}(0)|D)$
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Assumption 5: Strong Parallel Trends

$$\forall d, E[Y_t(d) - Y_{t-1}(0)] = E[Y_t(d) - Y_{t-1}(0)|D = d]$$

- It says that for all doses, the average change in outcomes over time across all unitsset if they had been assigned dose d, is the same as those actually experienced dose d.
- It imposes some homogeneity on treatment effect
- Not as strong as saying "parallel trend for group D = d and any other group D = d' if they are assigned dose d'"
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■ We can non-parametrically identify ACR under strong parallel trend assumption, in a DID fashion.

Under Assumptions 1 to 3 and 5, ACR(d) is identified:

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 - where, $(i)w_1(l) \ge 0$, $w_0 > 0$, $(ii) \int_{a_0}^{a_0} w_1(l)dl + w_0 = 1$
 - The first term is the average causal effect (for the treated) of running from d_L to d_H
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Theorem 3 (b) in Callaway et al(2021

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Under strong parallel trend assumption, we eliminate the selection bias

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- Causal level effects on the treated are non-parametrically identified under common parallel trend assumption
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- Using structural method or theoretical models to help you to interpret your results
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- Linear regression is, after all, a parametric method, which imposes strong functional form assumptions
- It is a simple and elegant statistical tool
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