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ABSTRACT

Inference with Difference-in-Differences Revisited*

A growing literature on inference in difference-in-differences (DiD) designs with grouped errors has been pessimistic about obtaining hypothesis tests of the correct size, particularly with few groups. We provide Monte Carlo evidence for three points: (i) it is possible to obtain tests of the correct size even with few groups, and in many settings very straightforward methods will achieve this; (ii) the main problem in DiD designs with grouped errors is instead low power to detect real effects; and (iii) feasible GLS estimation combined with robust inference can increase power considerably whilst maintaining correct test size – again, even with few groups.

JEL Classification: C12, C13, C21

Keywords: difference in differences, hypothesis test, power, cluster robust, feasible GLS

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1 Introduction

Difference-in-differences (DiD) designs are extremely common as a way of estimating the effects of policies or programs (henceforth ‘treatment effects’). A recent literature has highlighted that failure to appropriately quantify the uncertainty surrounding DiD estimates can lead to dramatically misleading inference (e.g. Bertrand et al, 2004; Cameron and Miller, 2013). In particular, researchers will tend to reject true null hypotheses with a probability that is far higher than the nominal size of the hypothesis test. The literature has suggested that obtaining tests that are close to the correct size requires non-standard techniques, and that it may not be possible with a small number of groups (Angrist and Pischke, 2009; Bertrand et al, 2004; Cameron et al, 2008).

In this paper we report evidence from Monte Carlo simulations that emphasises a different conclusion. We make three main points. First, in many typical DiD settings tests of the correct size can be obtained with very straightforward methods that are trivial to implement with standard statistical software (in fact, STATA’s cluster-robust inference implements these methods by default); and in settings where this works less well, a bootstrap-based approach highlighted by other authors (e.g. Cameron et al, 2008; Webb, 2013) provides a reliable alternative. All this is true even with few groups. Second, these techniques have very low power to detect real treatment effects. Thus the real challenge for inference with DiD designs is power rather than size. Third, we show that substantial gains in power can be achieved using feasible GLS. Moreover, if feasible GLS is combined with robust inference, test size can still be controlled if the parametric assumptions about the error process implicit in FGLS estimation are violated, even with few groups. We therefore recommend that applied researchers using DiD designs pay careful attention not just to consistency and test size, but also to the efficiency of their estimators.

DiD designs often use micro-data but estimate the effects of a treatment which varies only at a group level at any point in time (e.g. variation in policy across US states). A consequence is that within-group correlation of errors can substantially increase the true level of uncertainty surrounding the treatment effect (e.g. Angrist and Pischke, 2009; Cameron and Miller, 2013; Donald and Lang, 2007; Moulton, 1990; Wooldridge, 2003). Furthermore, treatment status is typically highly serially correlated. In fact, in the most common case which we focus on in this paper, treatment is an ‘absorbing state’: once a group is treated, it remains treated in all subsequent periods. This means that serially correlated error terms are likely to have a large impact on the true level of precision with which treatment effects are estimated. In a well-cited Monte Carlo study using US earnings data, Bertrand et al (2004) show that accounting only for grouped errors at the state-time level whilst ignoring serial correlation led to a 44% probability of rejecting a true null hypothesis using a nominal 5% level test. So, for example, when evaluating a labor market policy implemented in certain regions from a particular point in time onwards, a researcher should worry both that people

in the same region at the same time are affected by common labor market shocks (unrelated to the policy) and that these regional shocks are serially correlated.

A simple approach to deal with both cross-sectional and serial correlation in within-group errors would be to use the formula for a cluster-robust variance matrix due to Liang and Zeger (1986). This is consistent and Wald statistics which use it are asymptotically normal, but the asymptotics apply as the number of clusters tends to infinity. By clustering at the group level rather than the group-time level to account for serial correlation, one is often left with few clusters. The finite sample (i.e. few-clusters) performance of this approach - an empirical question - then becomes crucial, and the literature to date has come to pessimistic conclusions about it. Bertrand et al (2004) and Cameron et al (2008) use US earnings data and generate placebo state-level treatments before estimating their ‘effects’. Forming t-statistics using cluster-robust standard errors (CRSEs), they obtain 9% and 11% rejection rates using nominal 5% level tests with samples from 10 and 6 US states respectively.¹ This is a considerable improvement over using OLS standard errors, when rejection rates are more than 40%. But it is still approximately double the nominal test size.

The crucial finding of Bertrand et al and others - that inference can go badly wrong in DiD unless one is very careful - is confirmed once again here. But we also show that a simple modification to the standard cluster-robust inference procedure described above can dramatically improve test size with few clusters. One can apply a scaling factor to the OLS residuals that are plugged into the CRSE formula, and use critical values from a t distribution with degrees of freedom equal to the number of groups minus one, rather than a standard normal. When this is done, our simulations show that true test size is within about one percentage point of nominal test size with 50, 20, 10 or 6 groups. We further show that this holds under a wide range of error processes. The key situation in which the method is unreliable is when there is a large imbalance between the numbers of treatment and control groups.

Various alternative techniques for achieving correct test size have been proposed and/or tested (Bertrand et al, 2004; Cameron et al, 2008; Donald and Lang, 2007; Bester et al, 2011). Of these, only a wild cluster bootstrap-t procedure has been shown to produce tests of approximately the right size in the typical DiD setup considered in this paper (see Section 2) when the number of groups is as small as six (Cameron et al, 2008). Like using CRSEs, this is theoretically robust to heteroscedasticity and arbitrary patterns of error correlation within clusters, and to variation in error processes across clusters. It has also

¹Both papers first account for cross-sectional within-group error correlation by aggregating to the group-time level, taking mean residuals within each group-time cell from a regression of earnings on individual-level characteristics. This is a straightforward way to deal with this problem and is appropriate in typical DiD settings where the number of observations per group-time cell is large. (It will also be the approach taken in this paper.) The remaining issues for inference are dealing with a finite number of groups and any serial correlation in group-time shocks.

been shown to be quite robust to large imbalances between the numbers of treatment and control groups (Mackinnon and Webb, 2013), and hence provides an important alternative to the simpler method described above in such situations. But it is less trivial to implement and computationally more intensive.

Our second point is that, while it is generally not difficult to obtain the correct size, power to detect real effects is a serious concern. When we use the methods above to implement correctly sized hypothesis tests, we find that DiD designs have very low power. This problem is very severe with few groups. For example, with a large 30-year panel of US earnings data from 6 states, a policy implemented by half of the states that raised earnings by 5% would be detected with only 17% probability (using a test of size 0.05). The policy would have to increase earnings by 16% if the null of no policy effect is to be rejected with 80% probability.

Finally, we show that substantial gains in power can be achieved using feasible GLS. In particular, with a moderate time series dimension of at least about 10 time periods, one will often be able to increase power by modeling the serial correlation of unobservables inherent in typical DiD designs. Test size can still be controlled in a way that is robust to having small numbers of groups, and to violations of the parametric assumptions about the error process implicit in FGLS estimation, using the straightforward cluster-robust inference technique described above. We therefore recommend the use of the combination of FGLS and cluster-robust techniques in DiD applications. We also confirm that, in the absence of robust inference, test size can be significantly improved using a bias correction for the OLS estimates of the parameters of an AR process derived in Hansen (2007).

The paper proceeds as follows. Section 2 describes the standard econometric setup in DiD designs that we consider, and discusses possible solutions to the inference problems that can arise in this setting. Section 3 details the Monte Carlo design we use to test different inference methods. Section 4 presents and discusses the results of our Monte Carlo simulations. Section 5 summarizes and concludes.

2 Approaches to inference in a difference-in-differences design

We consider the standard linear DiD model

$$y_{igt} = \alpha_g + \delta_t + \beta T_{gt} + \gamma w_{igt} + v_{igt}, \quad (1)$$

where α_g and δ_t capture group (state) and time (year) fixed effects, β is the treatment effect of interest for a treatment which varies at the group-time level only, w_{igt} are individual-level control variables, and v_{igt} is the unobserved individual-level earnings shock.

Our interest lies in the performance of different methods for performing inference about

β , both in terms of type 1 and type 2 error (i.e. test size and power to detect real effects). Hence we assume that the OLS DiD estimator based on equation 1 is unbiased, i.e. $E(v_{igt}|\alpha_g, \delta_t, T_{gt}, w_{igt}) = 0$ so that $E(\hat{\beta}^{OLS}) = \beta$. (This is ensured in our Monte Carlo simulations because we generate placebo treatments randomly.)

The problem we seek to address is that the v_{igt} may not be iid within groups. Some of the variation in v_{igt} may occur at the group-time level, i.e. $v_{igt} = \varepsilon_{gt} + \xi_{igt}$. The DiD estimator is therefore effectively attempting to distinguish between the effects of a group-time level treatment and between-group differences in the evolution of group-time shocks. In addition, the group-time shocks may be serially correlated. The net result is both cross-sectional and serial correlation in within-group shocks. This is highly likely in many DiD applications, including the primary example used in this paper (and much of the previous literature) where groups are US states and the outcome of interest is earnings. The challenge, then, is to quantify accurately the additional uncertainty about β that this causes.

Given the setup described, the computation of $\hat{\beta}^{OLS}$ from a micro-data regression using equation 1 is equivalent to a two-step procedure. First, run a regression using the micro-data of y_{igt} on w_{igt} , and take the mean residual within each group-time cell. Denote these estimated covariate-adjusted group-time means as \hat{Y}_{gt} .² Then, since

$$\hat{Y}_{gt} = \alpha_g + \delta_t + \beta T_{gt} + \varepsilon_{gt} + (\hat{Y}_{gt} - Y_{gt}), \quad (2)$$

$\hat{\beta}^{OLS}$ can be obtained from a second-stage regression of \hat{Y}_{gt} on group effects, time effects and the (group-time level) treatment indicator. If state-time cell sizes are large, then estimation error in \hat{Y}_{gt} can essentially be ignored: the composite error term in equation 2 is approximately equal to ε_{gt} , the group-time shock. Equation 2 highlights that, in that case, the true precision of $\hat{\beta}^{OLS}$ depends almost entirely on the number of group-time cells rather than the number of individual-level observations.³

As we explain fully in the next section, we first aggregate the data to the state-time level in this way and ignore any estimation error (i.e. we proceed as though $\hat{Y}_{gt} = Y_{gt}$). We then estimate equation 2 and perform inference about β . As the first-stage aggregation accounts for cross-sectional error correlation within states, the key remaining issues for inference are the fact that the state-time shocks may be serially correlated and that there are a finite number of states. A number of methods have been proposed to account for these two issues. We describe them below, as well as some modest proposals of our own.

²Equivalently, one could include a full set of group-time dummies in this first regression (and omit the constant). The \hat{Y}_{gt} are the estimated coefficients on those dummies.

³If one is unsure whether this grouped error problem exists, Wooldridge (2006) points out that one could test for it. If the error term is dropped from equation 2, this imposes a set of (GT-1) restrictions on the data which can be used to compute a minimum distance estimator of β . One can then test the over-identifying restrictions. This is asymptotically valid as group-time cell sizes tend to infinity.

Standard cluster-robust inference

Our starting point is the standard OLS estimator of the standard error of $\hat{\beta}$, comparing the resulting t-statistic to standard normal critical values. This effectively assumes that the ε_{gt} in equation 2 are iid, i.e. it ignores serial correlation.

We then look at several ways of performing inference based on variants of Liang and Zeger’s (1986) cluster-robust standard error (CRSE) estimator. Their formula for a cluster-robust variance matrix is

$$\hat{V}_{CR} = (X'X)^{-1} \left(\sum_{g=1}^G X_g u_g u_g' X_g' \right) (X'X)^{-1}, \quad (3)$$

where X is the regressor matrix, X_g is the regressor matrix for group g , and u_g is the vector of regression residuals for group g . This estimator is consistent, and Wald statistics based on it are asymptotically normal, as $G \rightarrow \infty$. But it is biased, and the bias can be substantial when G is small. Intuitively, model over-fitting means that residuals will tend to be smaller in magnitude and less correlated within clusters than the true errors, meaning that CRSEs calculated using equation 3 will tend to be biased downwards. Any small- G bias is larger when the distribution of regressors is skewed: in the DiD context considered here, this is when there is an imbalance between the numbers of treatment and control groups (see Mackinnon and Webb, 2013).

Bias corrections for cluster-robust inference with few clusters

A typical way of attempting to reduce small- G bias (or, under special circumstances, to eliminate it) is effectively to scale up the residuals before plugging them into equation 3. The default in STATA is to scale by $\sqrt{\frac{G(N-1)}{(G-1)(N-k)}}$, where N is the total number of observations and k is the number of parameters.⁴ With large N , this is approximately equivalent to $\sqrt{G/(G-1)}$: the additional $\sqrt{(N-1)/(N-k)}$ is a degrees of freedom correction which makes a negligible difference in large samples (for brevity we refer to residuals scaled in this way simply as $\sqrt{G/(G-1)}$ residuals, but we use the additional $\sqrt{(N-1)/(N-k)}$ degrees of freedom adjustment so that our results can be taken as an exact test of how STATA’s default performs). This scaling of residuals leads to an unbiased CRSE estimator only under very special circumstances (see Bell and McCaffrey, 2002) and so should be viewed generally as a *bias-reducing* correction. The same applies to a second, data-dependent scaling of u_g proposed in Bell and McCaffrey (2002),⁵ and extended in Imbens and Kolesár (2012); Cameron and Miller (2013) investigate the Imbens and Kolesár (2012) adjustment in a set-up that is very similar to the one in this paper, and they show that the DOF

⁴When one uses the “vce(cluster *clustvar*)” option in a regression.

⁵This minimizes the expected sum of squared differences between the scaled residuals and the true errors in the baseline case where errors are iid.

adjustment is of minimal importance in balanced DiD designs.

For CRSEs formed using unscaled and $\sqrt{G/(G-1)}$ residuals, we show rejection rates when comparing the resulting t-statistics against critical values from both a standard normal and a t distribution with $G-1$ degrees of freedom. The former reference distribution is correct asymptotically as $G \rightarrow \infty$, so the implicit assumption when using it is that G is large enough for the asymptotics to be a reliable guide. The latter is a common small- G correction, again used by STATA for Wald tests and confidence intervals. As one expects with finite sample methods, in general it does not have an exact theoretical justification.⁶ However, recent work by Bester et al (2011) provides theoretical justification in certain small- G settings for the *combination* of CRSEs using $\sqrt{G/(G-1)}$ -scaled residuals and t_{G-1} critical values. Their asymptotics apply as group size tends to infinity, holding the number of groups fixed. Despite the familiar result that a CRSE estimator is not consistent with fixed G , they show that plugging $\sqrt{G/(G-1)}$ -scaled residuals into the CRSE formula nevertheless produces a covariance matrix which converges to a limiting random variable under certain conditions. Crucially, the resulting t-statistic turns out to have an asymptotic t_{G-1} distribution.⁷ This result relies on homogeneity requirements, including the need for regressor matrices to converge to the same limit within each group. This would be violated in the canonical DiD setup with a binary treatment indicator where some control groups are never treated.⁸ But the results we present in the following section suggest that, in practice, the Bester et al approach extends well (in terms of getting the test size right) to the standard DiD case.

Bootstraps

With few groups, an alternative to relying on asymptotic results (such as normality of the t-statistic) or on small sample corrections is to recover the distribution of the test statistic empirically via a bootstrap. Following Cameron et al (2008), we consider the wild cluster bootstrap-t procedure.⁹ Those authors found this to be the best (in terms of test size) of

⁶Donald and Lang (2007) show that a similar reference distribution - t_{G-2} - would provide tests of exactly the right size in the special case where the ε_{gt} were normal, homoscedastic and independent (i.e. serially uncorrelated).

⁷This result is also robust to violations of the assumption of no inter-cluster correlation, as long as data are weakly dependent and some regularity conditions are satisfied. In the context of spatial data where clusters are geographic regions, this implies robustness to the fact that there will be some clustering between observations which are spatially close but put into different clusters by the researcher. The intuition is that cluster size tending to infinity would mean that most observations per cluster are far from other clusters, and hence cluster averages will be approximately independent.

⁸The asymptotic variance of the score also needs to be the same across groups.

⁹We follow those authors in resampling clusters of residuals obtained from regressions which impose the null hypothesis, and scaling the resampled residuals by a constant drawn from a 2-point distribution: 1 and -1, each with probability 0.5. See Cameron et al (2008) for full details. We use 199 bootstrap replications, which is sufficient in this context as bootstrap simulation error will average out across Monte Carlo replications. We note that, as pointed out recently by Webb (2013), p-values are not point identified when the number of groups is very small. For example, with $G = 6$ there are only $2^G = 64$ potential unique bootstrap samples and $2^{G-1} = 32$ possible t-statistics (in absolute value).

a large number of inference techniques in settings with few groups. It outperformed other bootstrap-based approaches, as well as inference based upon t-statistics formed with CRSEs. But that paper did not consider the $\sqrt{G/(G-1)}$ residual correction, and it took critical values from the standard normal distribution, rather than from the t distribution; as we show in the next section, both of these can be useful small-G modifications to standard cluster-robust inference such that, when implemented in combination, they produce hypothesis tests of the correct size in most settings.

Modeling the error process using GLS

The final approach to dealing with the serial correlation in the group-time shocks is to use feasible Generalized Least Squares (GLS): this effectively exploits knowledge of this feature of the data to increase efficiency. A natural way to proceed is to assume an AR(k) process for the group-time shocks. FGLS can then be implemented by estimating equation 2 using OLS, as before; estimating the k AR parameters using the OLS regression residuals; using those estimates to apply the standard GLS linear transformations to the variables entering equation 2; and estimating the analog of equation 2 on the transformed variables via OLS.

Two issues arise. First, estimates of the AR(k) parameters obtained by regressing OLS residuals on k lags are inconsistent with T fixed, due to the presence of fixed group effects (Nickell, 1981; Solon, 1984). Hansen (2007) derives a bias correction which is consistent as $G \rightarrow \infty$, and develops the asymptotic properties of a FGLS estimator which uses it. But this correction may not work well with small G. Second, one may be worried about mis-specification of the error process.

However, neither of these issues affect the unbiasedness or consistency of the FGLS estimator. And it is likely that FGLS would still be more efficient than OLS: a weighting matrix based on an incorrect parametrization of the serial correlation process will often still be closer to the optimal GLS weighting matrix than the identity matrix used by standard OLS.

On the other hand, test size will generally be compromised, because the ordinary formula for the FGLS standard error depends upon the weighting matrix. But robust inference may offer a way to control test size. As noted more generally by Wooldridge (2006), the combination of FGLS estimation and robust inference is used relatively little in practice, but will often be a sensible way of realizing efficiency gains without compromising test size. One simply plugs the FGLS residuals, rather than OLS residuals, into the formula for a cluster-robust variance matrix.

Hansen (2007) considers this approach in the context of his FGLS procedure using bias-corrected estimates of the AR(k) parameters underlying the group-time error process, for the case where $G = 50$. The prevailing view is that the limitation of using cluster-robust inference is that its validity depends on having lots of groups. But one of the contributions

of this paper is to show that simple modifications to standard cluster-robust inference enable test size to be controlled, even with few groups. This suggests that it may be possible - and indeed straightforward - to use FGLS to improve power in DiD, whilst maintaining correctly sized tests, in a way that is robust to mis-specification (or mis-estimation) of the error process, even with a small number of groups. Our simulations confirm this.

3 Experimental design

We follow Bertrand et al (2004), Cameron et al (2008) and Hansen (2007) in using data on women aged 25 to 50 in their fourth interview month in the Merged Outgoing Rotation Group of the Current Population Survey. Our data include all 50 US states and the period 1979 to 2008 inclusive (i.e. $G = 50$ and $T = 30$). We focus primarily on $\log(\text{earnings})$ as the dependent variable. We also consider the case where a binary employment indicator is the dependent variable in a linear probability model.¹⁰ This covers the two most common outcomes of interest in DiD studies, according to a survey of the applied literature in Bertrand et al (2004). Our control variables are a quartic in age. As in the aforementioned papers, we first aggregate the data to the state-time level in the way just described and ignore any estimation error from this procedure (i.e. we proceed as though $\hat{Y}_{gt} = Y_{gt}$).¹¹ We then estimate equation 2. As the first-stage aggregation accounts for cross-sectional error correlation within states, the key remaining issues for inference are the fact that the state-time shocks may be serially correlated and that there are a finite number of states.¹²

In our first set of Monte Carlo simulations, we repeatedly resample states with replacement from the CPS data and randomly choose half of the states to be ‘treated’.¹³¹⁴ For all treated states in each Monte Carlo replication, the placebo treatment is applied in the

¹⁰This gives us samples based upon the 750,127 women with strictly positive earnings and the 1,170,522 women with non-missing employment status respectively.

¹¹Given large state-time cell sizes, aggregation should average out the individual-level shock component precisely. Mean cell sizes are 500 and 780 when the dependent variables are $\log(\text{earnings})$ and employment status respectively.

¹²We recommend the first-step aggregation not only to make the estimation simpler computationally. We find that, even with moderate numbers of groups, test size can not be reliably controlled if one attempts to conduct cluster-robust inference straight from the micro-data (i.e. if one tries to account for all cross-sectional and serial correlation in within-group errors in a single step). This issue was also evident in the results of Cameron et al (2008) and is noted in Hansen (2007).

¹³In treating exactly half of the states, we follow the main approach in Bertrand et al (2004) and Cameron et al (2008). This is the most favorable possible choice in terms of the resulting precision of treatment effect estimates, as it maximizes between-group variation in treatment status.

¹⁴In the particular example we use here where groups are geographical units, the assumption of no inter-group error correlation is not likely to be reasonable close to the groups’ boundaries. This is an advantage of generating placebo treatments randomly in the experiment: Barrios et al (2012) show that, as long as there is no *cross-cluster* spatial correlation in treatment status, correct test size is robust to some correlation in the error terms across clusters, as long as the data are weakly dependent so that error correlation decays with distance. Hence, we will not be confusing the impacts on test size of inadequately accounting for grouped errors with the impacts of (incorrectly) assuming that the earnings shocks of people in geographical proximity but in different states are independent.

same randomly chosen year and in all subsequent years.¹⁵ We estimate the ‘effect’ of this placebo treatment by estimating equation 2. We initially use OLS, and later feasible GLS, for estimation. Our interest lies in the performance of different methods for performing inference about β , both in terms of type 1 and type 2 error (i.e. test size and power to detect real effects). To examine the effects of having differing numbers of groups, we run variants where we resample 50, 20, 10 and 6 states.

We first report how often the null hypothesis of no treatment effect is rejected using tests of nominal size 0.05 when using different inference methods. We show that tests of correct size can be achieved, even with as few as six groups, by forming a t-statistic using a simple variant of Liang and Zeger’s (1986) cluster-robust standard error estimator and comparing it to critical values from a t_{G-1} distribution.

We check the robustness of this result in two ways. First, we repeat the same Monte Carlo experiment but use simulated state-time shocks rather than those from the CPS, allowing them to evolve according to an AR(1) process where we vary both the amount of serial correlation and the degree of non-normality in the white noise. Second, we vary the fraction of groups that are treated, to explore robustness to unbalanced designs.

We then look at power by reporting how often the null of no effect is rejected when there are real treatment effects of various sizes, when using correctly sized tests. And we compute minimum detectable effects (MDEs) as first defined in Bloom (1995): the smallest effects that would lead to a rejection of the null hypothesis (of no effect) with given probabilities. To do this, we use the same Monte Carlo procedures as described above to simulate the distribution of the t-statistic under the null hypothesis.¹⁶ For power of $x\%$, the MDE depends only on the $(100-x)$ th centile of this distribution, the critical values from the t_{G-1} distribution, and the standard error (see later). We therefore recover the entire relationship between power and MDEs. We do this for DiD designs with varying numbers of groups.

Finally, we show how power can be improved by using FGLS rather than OLS estimation. We rerun the Monte Carlo simulations, this time implementing FGLS (rather than OLS). We assume an AR(2) process for the group-time shocks. We estimate the 2 AR parameters in two ways. First, we simply regress the residuals from OLS estimation of equation 2 on two lags. With fixed T and fixed group effects, this produces inconsistent estimates of the AR parameters. Second, we apply to these estimates the bias correction derived by Hansen (2007). This correction is consistent as G goes to infinity. We label these “FGLS” and “BC-FGLS” respectively. In both cases, we explore what happens when the estimator is used with and without cluster-robust inference. We use the cluster-robust technique that we have shown to work well even when G is small: using CRSEs with $\sqrt{G/(G-1)}$ residuals and t_{G-1} critical values.

¹⁵The treatment year is chosen from a uniform distribution between 1988 and 2002.

¹⁶This is necessary because, with few clusters, the t-statistic generally has an unknown distribution.

4 Results

4.1 Rejection rates when the null is true

Table 1 contains results from our first Monte Carlo simulations, using the CPS log(earnings) data. It shows the rate with which the null of no effect is rejected when generating placebo treatments, estimating equation 2 by OLS, and using methods to perform inference about β . All hypothesis tests are of nominal size 0.05. Hence, rejection rates that deviate significantly from 0.05 indicate incorrect test size. We use 5000 replications. Simulation standard errors are shown in parentheses. The standard error for an estimated rejection rate \hat{r} is $se(\hat{r}) = \sqrt{\hat{r}(1 - \hat{r})/4999}$.

The first row of table 1 shows the rejection rates obtained assuming iid errors, i.e. by simply forming a t-statistic using the OLS standard error and comparing to standard normal critical values. Rejection rates exceed 40%, more than eight times the nominal test size. This essentially replicates the result in Bertrand et al (2004).

Forming CRSEs using unscaled OLS residuals and comparing the resulting t-statistic to standard normal critical values results in rejection rates that are too high, particularly with small G . Using t_{G-1} rather than the standard normal as the reference distribution is enough to achieve approximately the correct test size when $G \geq 20$, but not with 6 or 10 groups.

The $\sqrt{G/(G-1)}$ residual correction, combined with t_{G-1} critical values, achieves a test size that deviates by less than 1 percentage point from the nominal test size when G ranges between 6 and 50. The same residual correction combined with standard normal critical values also works well for moderate G but, as expected, these critical values result in over-rejection when G is small.

The final row of table 1 shows rejection rates obtained using the wild cluster bootstrap-t procedure. We essentially replicate previous findings: it also performs very well relative to most tested alternatives.

In summary, table 1 suggests that tests of the correct size can be obtained using very straightforward methods even with very few groups. In particular, this is achieved by computing a t-statistic with CRSEs that use residuals scaled by $\sqrt{\frac{G(N-1)}{(G-1)(N-K)}} \approx \sqrt{G/(G-1)}$, and using critical values from a t distribution with $(G-1)$ degrees of freedom. This is trivial to implement with statistical software. In fact, if one uses a cluster-robust variance matrix in STATA by specifying the “vce(cluster *clustvar*)” option, the confidence intervals and p-values returned are based upon precisely this procedure by default.¹⁷

In table 2 we present results from an analogous set of Monte Carlo simulations using employment status rather than earnings as the dependent variable.¹⁸ The performance of different inference methods in data containing varying numbers of groups is essentially the

¹⁷This is true at the time of writing (STATA version 12.1) and has been the case since at least STATA 6.

¹⁸Given that we collapse the data to the state-time level in a first stage, this means that Y_{gt} now represents state-time employment rates rather than mean state-time earnings.

same as in Table 1. In particular, CRSEs formed using the $\sqrt{G/(G-1)}$ residual correction combined with t_{G-1} critical values perform best, with rejection rates always within 1 percentage point of the nominal test size. Again, the wild cluster bootstrap-t also performs well relative to most alternatives.

Table 1: Rejection rates for tests of nominal 5% size with placebo treatments in log-earnings data

	G=50	G=20	G=10	G=6
Assume iid	0.429 (0.007)	0.424 (0.007)	0.422 (0.007)	0.413 (0.007)
CRSE, N(0,1) critical values	0.059 (0.003)	0.073 (0.004)	0.110 (0.004)	0.175 (0.005)
CRSE, t(G-1) critical values	0.053 (0.003)	0.056 (0.003)	0.066 (0.004)	0.095 (0.004)
$\sqrt{G/(G-1)}$ residuals, N(0,1) critical values	0.049 (0.003)	0.056 (0.003)	0.071 (0.004)	0.113 (0.004)
$\sqrt{G/(G-1)}$ residuals, t(G-1) critical values	0.045 (0.003)	0.041 (0.003)	0.042 (0.003)	0.052 (0.003)
Wild cluster bootstrap-t	0.044 (0.003)	0.041 (0.003)	0.048 (0.003)	0.059 (0.003)

The Table shows the proportion of the time that the null hypothesis of no treatment effect was rejected in 5000 Monte Carlo simulations. Simulation standard errors are reported in parentheses. The simulations resample groups (US states) from CPS-MORG data, having imposed the sampling restrictions described in the text. The treatment parameter has a true coefficient of zero. G is the number of sampled states. Data from 1979 to 2008 inclusive are sampled (i.e. T = 30). Regressions are run on aggregated state-year data. The different inference methods used are discussed in the text.

We have shown so far that our conclusions apply both when the outcome of interest is earnings and when it is a binary employment status indicator. But one might still worry about the generality of these findings. We now explore two variants of the experiment: unbalanced designs where the numbers of treatment and control groups are not equal; and experiments where the error terms are simulated from various data generating processes rather than from the CPS data.

Variant 1: unbalanced designs

The accuracy of robust variance matrix estimators in finite samples depends on the skewness of the distribution of regressors as well as sample size (as illustrated in Monte Carlo work

Table 2: Rejection rates for tests of nominal 5% size with placebo treatments in employment data

	G=50	G=20	G=10	G=6
Assume iid	0.376 (0.007)	0.360 (0.007)	0.378 (0.007)	0.364 (0.007)
CRSE, N(0,1) critical values	0.062 (0.003)	0.076 (0.004)	0.119 (0.005)	0.184 (0.005)
CRSE, t(G-1) critical values	0.056 (0.003)	0.059 (0.003)	0.078 (0.004)	0.097 (0.004)
$\sqrt{G/(G-1)}$ residuals, N(0,1) critical values	0.052 (0.003)	0.058 (0.003)	0.085 (0.004)	0.114 (0.004)
$\sqrt{G/(G-1)}$ residuals, t(G-1) critical values	0.046 (0.003)	0.044 (0.003)	0.056 (0.003)	0.058 (0.003)
Wild cluster bootstrap-t	0.048 (0.003)	0.039 (0.003)	0.056 (0.003)	0.064 (0.003)

Notes as for Table 1.

by Imbens and Kolesár, 2012). With cluster-robust standard errors in the DiD context considered here, a researcher should pay attention not only to the number of groups but also to whether treatment status is skewed, i.e. whether the number of treated groups is similar to the number of controls.

With unbalanced designs, inference based on cluster-robust standard errors should produce tests of correct size less reliably than when the numbers of treatment and control groups are equal (as in the Monte Carlo experiments up to now). This has been illustrated recently by Mackinnon and Webb (2013). Scaling the residuals that are plugged into the standard CRSE formula by $\sqrt{G/(G-1)}$ (and using a t_{G-1} reference distribution) may not be a sufficient small- G correction in the presence of this imbalance.

Tables 3 and 4 repeat the simulations presented in Table 1 but with varying degrees of imbalance, for the cases with 10 and 50 groups respectively. G_1 denotes the number of treated groups. The first columns repeat the results for the balanced designs shown in Table 1, and subsequent columns reduce the number of treated groups. When $G = 10$, the simple method that works well under a wide a wide range of balanced designs - using CRSEs with $\sqrt{G/(G-1)}$ -scaled residuals and a t_{G-1} reference distribution - continues to achieve correct test size with $G_1 = 4$, but over-rejects a little when G_1 drops to 3, and more severely when it drops to 2. The wild cluster bootstrap-t continues to work well with $G_1 = 3$ but also performs poorly with $G_1 = 2$ (with significant *under*-rejection). When $G = 50$, both

methods continue to produce tests of the close to the right size when $G1$ drops as low as 10. With $G1 = 5$ the bootstrap is needed to conduct approximately accurate inference, while the simpler method over-rejects.

In summary these simulations confirm that, if the imbalance between treatment and control groups is large enough, using CRSEs with $\sqrt{G/(G-1)}$ -scaled residuals and a t_{G-1} reference distribution can lead to over-rejection. But as emphasised in Mackinnon and Webb (2013), the wild cluster bootstrap-t is relatively robust in this regard. Hence, although a slightly less trivial procedure is necessary to get test size right with very unbalanced designs, it can be done even with few groups.¹⁹

Table 3: Rejection rates for tests of nominal 5% size with placebo treatments in log-earnings data with 10 groups

	$G1 = 5$	$G1 = 4$	$G1 = 3$	$G1 = 2$
Assume iid	0.422 (0.007)	0.408 (0.007)	0.409 (0.007)	0.405 (0.007)
CRSE, N(0,1) critical values	0.110 (0.004)	0.125 (0.005)	0.150 (0.005)	0.241 (0.006)
CRSE, t(G-1) critical values	0.066 (0.004)	0.079 (0.004)	0.105 (0.004)	0.191 (0.006)
$\sqrt{G/(G-1)}$ residuals, N(0,1) critical values	0.071 (0.004)	0.084 (0.004)	0.113 (0.004)	0.199 (0.006)
$\sqrt{G/(G-1)}$ residuals, t(G-1) critical values	0.042 (0.003)	0.051 (0.003)	0.074 (0.004)	0.150 (0.005)
Wild cluster bootstrap-t	0.048 (0.003)	0.054 (0.003)	0.052 (0.003)	0.018 (0.002)

The Table shows the proportion of the time that the null hypothesis of no treatment effect was rejected in 5000 Monte Carlo simulations. Simulation standard errors are reported in parentheses. The simulations resample groups (US states) from CPS-MORG data, having imposed the sampling restrictions described in the text. The treatment parameter has a true coefficient of zero. $G1$ denotes the number of groups that are treated. Data from 1979 to 2008 inclusive are sampled (i.e. $T = 30$). Regressions are run on aggregated state-year data. The different inference methods used are discussed in the text.

¹⁹For the extreme case - not considered here - where there is just one treated group, see Conley and Tabler (2011). They develop a method for inference which is valid in this design when there are lots of control groups.

Table 4: Rejection rates for tests of nominal 5% size with placebo treatments in log-earnings data with 50 groups

	G1 = 25	G1 = 15	G1 = 10	G1 = 5
Assume iid	0.429 (0.007)	0.418 (0.007)	0.425 (0.007)	0.413 (0.007)
CRSE, N(0,1) critical values	0.059 (0.003)	0.067 (0.004)	0.077 (0.004)	0.136 (0.005)
CRSE, t(G-1) critical values	0.053 (0.003)	0.060 (0.003)	0.068 (0.004)	0.128 (0.005)
$\sqrt{G/(G-1)}$ residuals, N(0,1) critical values	0.049 (0.003)	0.058 (0.003)	0.065 (0.003)	0.126 (0.005)
$\sqrt{G/(G-1)}$ residuals, t(G-1) critical values	0.045 (0.003)	0.052 (0.003)	0.060 (0.003)	0.119 (0.005)
Wild cluster bootstrap-t	0.044 (0.003)	0.051 (0.003)	0.046 (0.003)	0.060 (0.003)

Notes as for Table 3.

Variant 2: alternative data generating processes

We now conduct our Monte Carlo experiments using simulated data (returning to a balanced design). The earnings generating process still conforms with equation 2, but we now simulate the state-time shocks ourselves. In doing so we vary their degree of serial correlation and non-normality. The state-time shocks for each state evolve according to the AR(1) process

$$\varepsilon_{gt} = \rho\varepsilon_{g,t-1} + \sqrt{\frac{0.004(1-0.4^2)(d-2)}{d}}\omega_{gt}, \quad t = 2, \dots, 30$$

$$\varepsilon_{g1} = \sqrt{\frac{0.004d}{d-2}}\omega_{g1},$$

where ω_{gt} is iid across groups and time and is drawn from a t distribution with d degrees of freedom. To control the degree of non-normality in the white noise, we vary d between 4 (very high non-normality) and 120 (at which point the t distribution is essentially standard normal). To control the degree of serial correlation, we vary ρ . We also examine a scenario in which the data generating process is heterogeneous, by drawing ρ separately for each state from a uniform distribution between 0 and 1. The scaling applied to ω_{gt} ensures that,

when $\rho = 0.4$, the variance of ε_{gt} is equal to 0.004 - approximately the empirical variance of the residuals in the CPS data. This means that the degree of serial correlation is allowed to affect the stationary variance of ε_{gt} , but the distribution of the white noise is not. We generate the initial condition (ε_{g1}) such that its variance matches the stationary variance of state-time shocks in other time periods.

In each Monte Carlo replication, we first resample states with replacement from the CPS data and randomly choose treated states and the year in which the placebo treatment begins, just as before. We then regress Y_{gt} on state and year fixed effects only. For each state-time combination, we simulate the outcome variable by summing the relevant (estimated) state effect, the relevant (estimated) year effect, and the random state-year shock generated as above. We then estimate the DiD model using the transformed outcome and conduct the hypothesis test on β . We use 10,000 replications.

Tables 5 to 8 report rejection rates for various combinations of ρ and d , when varying the number of groups between 50 and 6. They show that our finding is robust to a very wide range of error processes. Rejection rates remain within about a percentage point of the nominal test size under all of the tested combinations of degrees of serial correlation, non-normality in the white noise, and number of groups.

4.2 Power to detect real effects

Our findings in the previous section indicate that controlling test size need not be a major concern in DiD designs. However, we now show that power to detect real treatment effects with tests of correct size can be extremely low.

Rejection rates in table 9 indicate power to detect treatment effects on earnings of approximately 2% (precisely, 0.02 log-points), 5%, 10% and 15%. This is based upon the same Monte Carlo replications as table 1, except we transform the dependent variable: for example, to look at power to detect a 5% effect we add $0.05T_{gt}$ to Y_{gt} .

We focus on the two methods that we have shown to produce approximately correctly sized hypothesis tests even when the number of groups is small: the $\sqrt{G/(G-1)}$ residual scaling combined with t_{G-1} critical values, and the wild cluster bootstrap-t. Nevertheless, to ensure that we are comparing the power of hypothesis tests which have *exactly* the same size, we adopt the useful procedure suggested by Davidson and Mackinnon (1998). The nominal significance level used to determine whether to reject the null hypothesis is that which gives a test of true size 0.05. This nominal significance level is obtained from the 5th percentile of the empirical distribution of p-values from Monte Carlo simulations under a true null (i.e. the simulations underlying the results in table 1). As the results in Table 1 suggest, for both of these methods this is a number very close (but not generally identical) to 0.05. All results reported in Table 9 use this 'size-adjusted' measure of power.

The results indicate that power is a serious issue in these designs. A 2% effect would be

Table 5: Rejection rates for tests of nominal 5% size using $\sqrt{G/(G-1)}$ -CRSEs and t_{G-1} critical values with 50 groups (simulated data)

	$\rho = 0$	$\rho = 0.2$	$\rho = 0.4$	$\rho = 0.6$	$\rho = 0.8$	ρ varies with g
d=4	0.041 (0.002)	0.045 (0.002)	0.047 (0.002)	0.048 (0.002)	0.048 (0.002)	0.044 (0.002)
d=20	0.049 (0.002)	0.045 (0.002)	0.046 (0.002)	0.044 (0.002)	0.046 (0.002)	0.043 (0.002)
d=60	0.043 (0.002)	0.047 (0.002)	0.047 (0.002)	0.046 (0.002)	0.046 (0.002)	0.049 (0.002)
d=120	0.046 (0.002)	0.047 (0.002)	0.047 (0.002)	0.048 (0.002)	0.049 (0.002)	0.045 (0.002)

The Table shows the proportion of the time that the null hypothesis of no treatment effect was rejected in 10000 Monte Carlo simulations. Simulation standard errors are reported in parentheses. The simulations resample groups (US states) from CPS-MORG data, having imposed the sampling restrictions described in the text. The treatment parameter has a true coefficient of zero. Data from 1979 to 2008 inclusive are sampled (i.e. $T = 30$). Regressions are run on aggregated state-year data. Simulated log-earnings are generated by effectively replacing the empirical regression residuals with a simulated error term generated according to an AR(1) process. Each cell in the table represents a different AR(1) process. ρ denotes the AR(1) parameter. In the final column the AR(1) parameter is drawn separately for each group, from a uniform distribution between 0 and 1. d denotes the degrees of freedom of the scaled t distribution from which the white noise is drawn (hence it controls the degree of non-normality). See text for full details.

Table 6: Rejection rates for tests of nominal 5% size using $\sqrt{G/(G-1)}$ -CRSEs and t_{G-1} critical values with 20 groups (simulated data)

	$\rho = 0$	$\rho = 0.2$	$\rho = 0.4$	$\rho = 0.6$	$\rho = 0.8$	ρ varies with g
d=4	0.048 (0.002)	0.047 (0.002)	0.049 (0.002)	0.045 (0.002)	0.045 (0.002)	0.050 (0.002)
d=20	0.049 (0.002)	0.045 (0.002)	0.050 (0.002)	0.044 (0.002)	0.047 (0.002)	0.047 (0.002)
d=60	0.049 (0.002)	0.048 (0.002)	0.050 (0.002)	0.050 (0.002)	0.044 (0.002)	0.047 (0.002)
d=120	0.049 (0.002)	0.043 (0.002)	0.049 (0.002)	0.044 (0.002)	0.046 (0.002)	0.048 (0.002)

Notes as for Table 5.

Table 7: Rejection rates for tests of nominal 5% size using $\sqrt{G/(G-1)}$ -CRSEs and t_{G-1} critical values with 10 groups (simulated data)

	$\rho = 0$	$\rho = 0.2$	$\rho = 0.4$	$\rho = 0.6$	$\rho = 0.8$	ρ varies with g
d=4	0.054 (0.002)	0.052 (0.002)	0.049 (0.002)	0.051 (0.002)	0.056 (0.002)	0.053 (0.002)
d=20	0.051 (0.002)	0.050 (0.002)	0.050 (0.002)	0.049 (0.002)	0.049 (0.002)	0.051 (0.002)
d=60	0.053 (0.002)	0.050 (0.002)	0.050 (0.002)	0.047 (0.002)	0.053 (0.002)	0.053 (0.002)
d=120	0.052 (0.002)	0.051 (0.002)	0.053 (0.002)	0.053 (0.002)	0.054 (0.002)	0.055 (0.002)

Notes as for Table 5.

Table 8: Rejection rates for tests of nominal 5% size using $\sqrt{G/(G-1)}$ -CRSEs and t_{G-1} critical values with 6 groups (simulated data)

	$\rho = 0$	$\rho = 0.2$	$\rho = 0.4$	$\rho = 0.6$	$\rho = 0.8$	ρ varies with g
d=4	0.056 (0.002)	0.064 (0.002)	0.063 (0.002)	0.057 (0.002)	0.060 (0.002)	0.061 (0.002)
d=20	0.061 (0.002)	0.062 (0.002)	0.064 (0.002)	0.060 (0.002)	0.061 (0.002)	0.060 (0.002)
d=60	0.062 (0.002)	0.063 (0.002)	0.061 (0.002)	0.064 (0.002)	0.064 (0.002)	0.064 (0.002)
d=120	0.060 (0.002)	0.067 (0.002)	0.065 (0.002)	0.063 (0.002)	0.063 (0.002)	0.060 (0.002)

Notes as for Table 5.

detected with a probability of less than 1 in 4, even with data from all 50 US states. To detect a 5% effect with a probability of about 80% - a conventional benchmark for power - one would need data on all 50 US states. Power declines much further with G . With 6 states, a researcher would have less than a 50-50 chance of detecting even a 10% effect, a 17% chance of detecting a 5% effect, and a 7% chance of detecting a 2% effect (power barely greater than the size of the test). In other words, it is unlikely that one would detect effects of a typically realistic magnitude using a correctly sized test, and highly unlikely when the number of groups is small.

A comparison of the two inference methods suggests that their power is similar for all combinations of number of groups and size of treatment effect. If anything, the simpler $\sqrt{G/(G-1)}$ residual scaling combined with t_{G-1} critical values tends to have slightly higher size-adjusted power than the wild cluster bootstrap-t.

Figures 1 and 2 document power more comprehensively by showing the minimum effects that would be detected (i.e. that would lead to a rejection of the null of no effect) with given probabilities - a way of assessing statistical power first outlined by Bloom (1995). We vary power between 1% and 99% and compute the minimum detectable effects (MDEs) in each case. We continue just with the hypothesis test that uses CRSEs with $\sqrt{G/(G-1)}$ -scaled residuals and t_{G-1} critical values.

For a given level of power, x , the MDE is

$$MDE(x) = se(\hat{\beta}) [c_u - p_{1-x}^t], \quad (4)$$

where $se(\hat{\beta})$ is the $\sqrt{G/(G-1)}$ -corrected CRSE estimate, c_u is the upper critical value (the 97.5th percentile of the t_{G-1} distribution), and p_{1-x}^t is the $(1-x)th$ percentile of the t-statistic under the null hypothesis of no treatment effect.

We proceed with the same Monte Carlo design underlying the results in Table 1. Monte Carlo replications provide us with an estimate of the distribution of the t-statistic under the null. They also provide repeated estimates of the $\sqrt{G/(G-1)}$ -corrected CRSE: we plug each of those estimates into equation 4 in turn, and take the average. Due to the low computational intensity of this approach, we are able to use 100,000 Monte Carlo replications so that simulation error is negligible. We use equation 4 to compute MDEs for power ranging from 1% to 99%.

Figure 1 plots MDEs against power when the number of groups is 50, 20, 10 and 6. With earnings data on the entire US population (50 states), one would need a treatment effect of about 3.5% to have even a 50-50 chance of detecting it. With a sample from 6 US states - by no means an extreme example in the applied DiD literature - the MDE on earnings is about 16% for 80% power and 11% for 50% power. Figure 2 shows the analogous results using a binary employment indicator as the dependent variable. This leads to similar conclusions. For 80% power, the MDE on the employment rate with data from all 50 states is about 2

Table 9: Rejection rates for tests of true 5% size with different treatment effects (β) in log-earnings data

	G=50	G=20	G=10	G=6
$\beta = 0.02$: $\sqrt{G/(G-1)}$ -CRSEs, $t(G-1)$ critical values	0.238 (0.006)	0.134 (0.005)	0.088 (0.004)	0.074 (0.004)
$\beta = 0.02$: wild cluster bootstrap-t	0.225 (0.006)	0.125 (0.005)	0.093 (0.004)	0.074 (0.004)
$\beta = 0.05$: $\sqrt{G/(G-1)}$ -CRSEs, $t(G-1)$ critical values	0.822 (0.005)	0.513 (0.007)	0.273 (0.006)	0.168 (0.005)
$\beta = 0.05$: wild cluster bootstrap-t	0.799 (0.006)	0.490 (0.007)	0.283 (0.006)	0.167 (0.005)
$\beta = 0.10$: $\sqrt{G/(G-1)}$ -CRSEs, $t(G-1)$ critical values	1.000 (0.000)	0.919 (0.004)	0.718 (0.006)	0.448 (0.007)
$\beta = 0.10$: wild cluster bootstrap-t	0.999 (0.000)	0.898 (0.004)	0.712 (0.006)	0.429 (0.007)
$\beta = 0.15$: $\sqrt{G/(G-1)}$ -CRSEs, $t(G-1)$ critical values	1.000 (.)	0.995 (0.001)	0.904 (0.004)	0.755 (0.006)
$\beta = 0.15$: wild cluster bootstrap-t	1.000 (.)	0.992 (0.001)	0.896 (0.004)	0.700 (0.006)

The Table shows the proportion of the time that the null hypothesis of no treatment effect was rejected in 5000 Monte Carlo simulations. Simulation standard errors are reported in parentheses. The simulations resample states from CPS-MORG data, having imposed the sampling restrictions described in the text. β is the true value of the treatment parameter. G is the number of sampled states. Data from 1979 to 2008 inclusive are sampled (i.e. $T = 30$). Regressions are run on aggregated state-year data. The inference methods used are discussed in the text. We adjust for test size when making power comparisons using the procedure outlined by Davidson and Mackinnon (1998). See text for details.

percentage points, rising to 6.5 percentage points with 6 states.²⁰

4.3 Increasing power with feasible GLS

The previous subsection argued that lack of power is a key problem in typical DiD designs. This suggests that there may be large gains from efforts to improve the efficiency of estimation. The serial correlation problem inherent in a typical DiD study also suggests one way to go about this: exploit knowledge of this feature of the data using feasible GLS. Here we implement the FGLS estimation procedure suggested by Hansen (2007), with and without the straightforward robust inference procedure that we have shown to produce correctly sized tests even with few groups (see Section 2 for details). We show that, in combination with our recommended robust inference technique, FGLS can improve power considerably whilst maintaining correctly sized tests, in a way that is robust to mis-specification (or mis-estimation) of the error process, even with few groups.

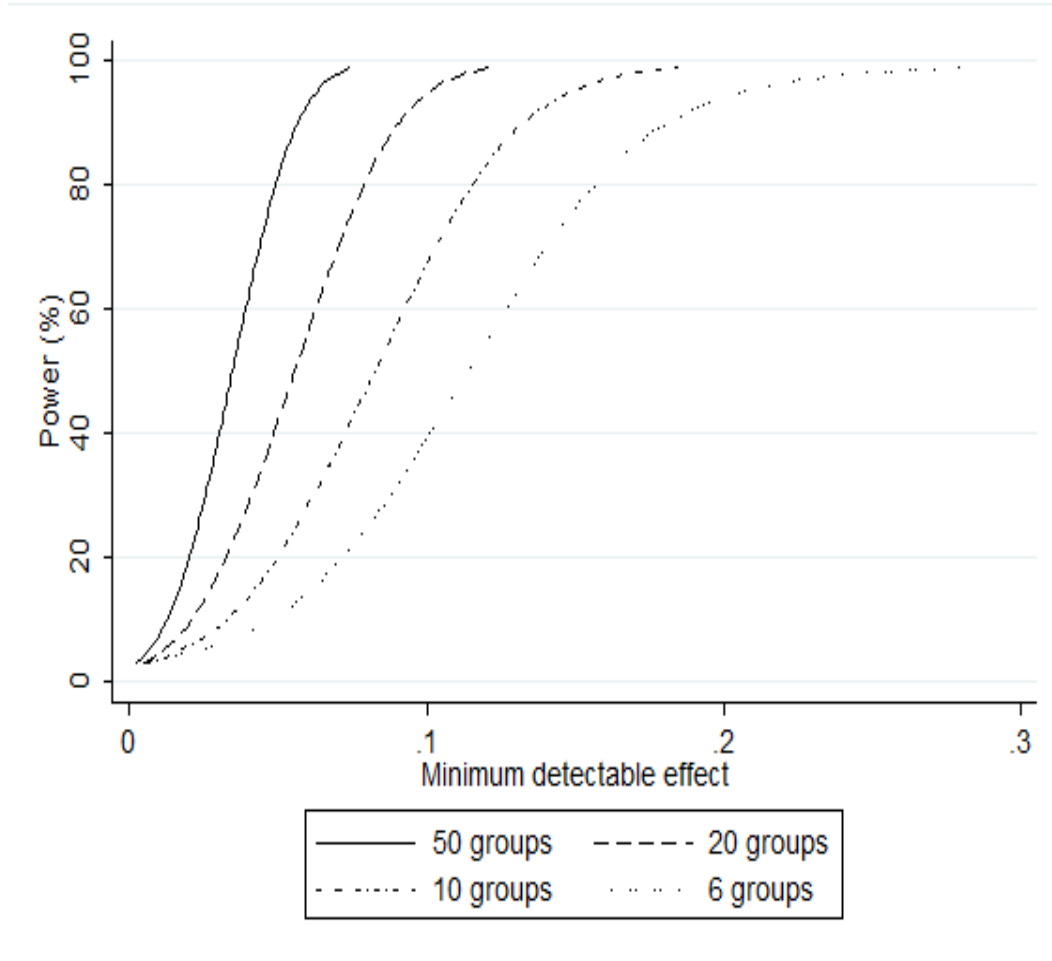
Table 10 shows rejection rates under a true null hypothesis (no treatment effect). The first row reiterates the good size properties of OLS estimation combined with CRSEs that use $\sqrt{G/(G-1)}$ -scaled residuals and t_{G-1} critical values (i.e. it repeats row four of Table 1). The second row shows that FGLS without the bias correction and without robust inference gives tests at least double the nominal size. Note, however, that even this size distortion is considerably smaller than with OLS without robust inference. Hansen’s bias correction for the estimated parameters of the AR process reduces this size distortion, though still returns rejection rates greater than the nominal test size without robust inference (fourth row), particularly when G is small. This is what we would expect, because the bias correction is consistent as $G \rightarrow \infty$. But, as with OLS, the size of the test can be controlled using robust inference, even with few groups, using the methods described earlier in this paper: when doing this, the rejection rate remains within about 1 percentage point of the nominal test size. This is true both for FGLS and BC-FGLS (third and fifth rows).

Table 11 turns attention to power, showing rejection rates when there is a 5% treatment effect on earnings. Again, the first row reiterates the earlier finding that OLS estimation combined with a correctly sized test provides low power (i.e. it repeats row four of Table ??), particularly with few groups. As Hansen (2007) showed in the case where $G = 50$, the FGLS procedures deliver substantial improvements in power. Combined with robust inference which delivers the correct test size, BC-FGLS detects the treatment effect with 96% probability, whereas OLS detects it with 80% probability. Table 11 shows that FGLS also delivers very substantial proportionate power gains relative to OLS with smaller G : with $G = 6$, power is 18% using OLS and 29% using FGLS. Using Hansen’s bias correction delivers a little more power than ‘ordinary’ FGLS.

Figure 3 illustrates the power gains more comprehensively, plotting MDEs against power

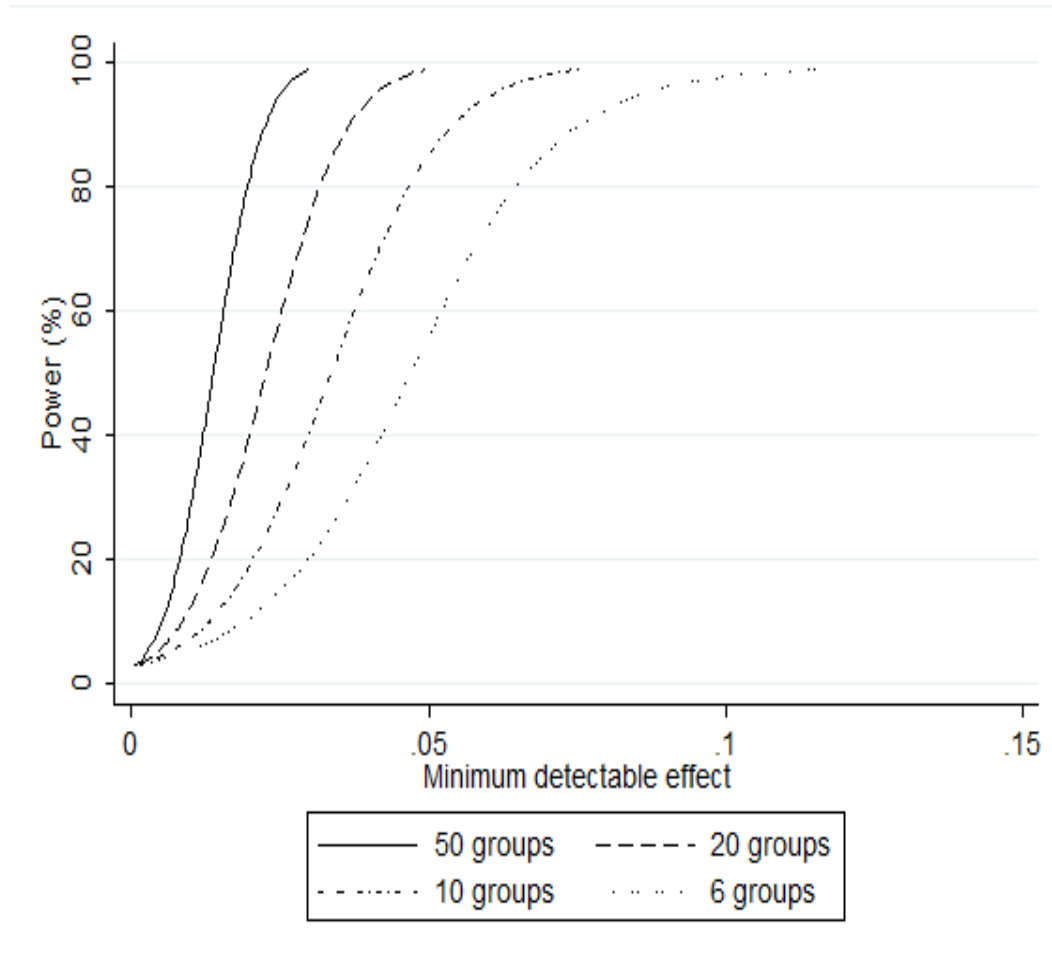
²⁰The baseline employment rate in the sample is 67%.

Figure 1: Minimum detectable effects on log-earnings using $\sqrt{G/(G-1)}$ -CRSEs and t_{G-1} critical values and tests of size 0.05



The Figure shows the proportion of the time that the null hypothesis of no treatment effect is rejected when the treatment parameter has a true coefficient ranging from 0 to 0.3. Numbers are computed using the results of 100000 Monte Carlo simulations combined with equation 4, as described in the text. The simulations resample states from CPS-MORG data, having imposed the sampling restrictions described in the text. Data from 1979 to 2008 inclusive are sampled (i.e. $T = 30$). Regressions are run on aggregated state-year data.

Figure 2: Minimum detectable effects on employment rates using $\sqrt{G/(G-1)}$ -CRSEs and t_{G-1} critical values and tests of size 0.05



The Figure shows the proportion of the time that the null hypothesis of no treatment effect is rejected when the treatment parameter has a true coefficient ranging from 0 to 0.15. Numbers are computed using the results of 100000 Monte Carlo simulations combined with equation 4, as described in the text. The simulations resample states from CPS-MORG data, having imposed the sampling restrictions described in the text. Data from 1979 to 2008 inclusive are sampled (i.e. $T = 30$). Regressions are run on aggregated state-year data.

and comparing the results for OLS and BC-FGLS estimation with varying numbers of groups (always combined with cluster-robust inference, so that test size is correct). The power gains from BC-FGLS are substantial.

Tables 12 and 13 and Figure 4 repeat this analysis for the case where the outcome of interest is a binary employment status indicator. This confirms that these conclusions all hold in that case: FGLS delivers substantial power gains over OLS, and this can be done whilst controlling test size, even with few groups.

We now further explore the robustness of these results to different data settings. Of particular interest are cases where the parametric assumptions about the serial correlation process inherent in the FGLS procedure are incorrect. In such cases, can test size still be reliably controlled (even with few groups), and what power gains (if any) can FGLS offer?

We continue to use FGLS estimation based on the assumption of an AR(2) process for the state-time shocks which is the same for all states. We explore its properties under two forms of mis-specification of the error process. First, we simulate an AR(2) process which is heterogeneous across states. Second, we simulate an MA(1) process with parameter 0.5.²¹ In each case, we effectively replace the empirical log-earnings residuals in the CPS (from a regression of state-year earnings on state and year fixed effects) with our simulated error terms, as in the robustness checks underlying tables 5 to 8.

Table 14 shows the results of these simulations under the null hypothesis of no treatment effect using tests of nominal size 0.05, for the case where $G = 10$. The first column re-iterates the rate at which the null hypothesis is rejected when using the empirical CPS error process (i.e. it repeats column 3 of table 10). The next two columns report the same statistics under the simulated error processes described above. The results show that the previous results on test size hold under these alternative processes: without robust inference, Hansen’s bias correction for the AR parameter estimates brings true test size closer to the nominal size when using FGLS estimation (although there is still some over-rejection); but test size can be controlled reliably using our suggested robust inference technique, whether estimation is carried out using OLS, FGLS or BC-FGLS.²²

Table 15 reports power to detect a treatment effect of 0.05 log-points on earnings, again for the case where $G = 10$.²³ The second column shows that, using correctly sized hypothesis

²¹For the heterogeneous AR(2) process, the coefficient on the first lag (α_1^g) is drawn from a uniform distribution between zero and one for each state. The coefficient on the second lag is set equal to $0.5 * \min(\alpha_1^g, 1 - \alpha_1^g)$, which ensures stationarity. For both the heterogeneous AR(2) and (homogeneous) MA(1) processes, the white noise in the process is normally distributed. Its variance is chosen so that the stationary variance of the simulated error term matches the empirical variance of the log-earnings residuals in the CPS (0.04).

²²We showed in Section 4.1 that this inference technique is not reliable if there is a large imbalance between the numbers of treatment and control groups; but that the wild cluster bootstrap-t procedure is relatively robust in such settings. FGLS estimation combined with bootstrap-based inference would be a sensible alternative in those situations.

²³We have also conducted this analysis with $G = 50$. Conclusions are qualitatively the same, although of course the power of all procedures is higher with more groups.

tests (i.e. those that use our suggested robust inference technique), FGLS estimation does offer substantial power gains over OLS even when based upon the incorrect assumption that the AR(2) error process is homogeneous across states.²⁴ The third column shows that, where the true error process is MA(1) rather than AR(2), there is no power gain from FGLS. Intuitively this makes sense: where the parametric assumptions about the serial correlation in the error terms are a very poor approximation to the true process, FGLS does not offer efficiency gains; where the assumptions are a better approximation, FGLS does offer efficiency gains over the OLS estimator, which does not exploit any knowledge of the nature of the error process.

Finally, we show how these results vary with the number of time periods available. This is an important dimension to explore, because we would expect the gains from modeling serial correlation to be more significant when T is large. Tables 16 and 17 repeat the simulations with $G = 10$ using CPS log-earnings as the outcome variable in cases where $T = 20$ and $T = 10$. This shows that test size can still be controlled using our recommended robust inference approach with few time periods, whether estimation is based on OLS or FGLS. It also shows, however, that the power gains from FGLS diminish with T , and with $T = 10$ the power of OLS and FGLS (when combined with inference which provides a correctly sized test) are essentially the same.

An interesting feature of Table 17 is that with OLS and robust inference, power actually declines with T .²⁵ Further inspection of the underlying simulations suggests that there is a genuine increase in the precision of OLS estimation as T falls: the estimated policy effects become more tightly distributed around their true value.

The reason why this is possible in this context is as follows.²⁶ DiD regressions effectively estimate the difference in mean outcomes between post-treatment periods and pre-treatment periods (and then compare these differences across treatment and control groups). The variance of this difference is decreasing in the covariance between the error terms pre and post treatment (intuitively, if error terms pre and post treatment covaried perfectly then they would not add any noise to the difference between pre and post treatment outcomes because they would cancel out). If serial correlation between observations decays with time, then the error term from an additional time period pre (post) treatment will covary less strongly with error terms in the post (pre) treatment period than the error terms already present. Hence the ‘covariance effect’ acts to increase the variance of the DiD estimate of the treatment effect when you add another time period to the data. On the other hand, of course, the variance is also increasing in the variance of the average error terms both

²⁴We also re-ran the earlier robustness checks where state-time earnings shocks evolve according to an AR(1) process, with varying degrees of serial correlation and varying degrees of non-normality in the white noise. The same qualitative conclusions about the size and power of FGLS and OLS combined with robust inference continued to hold. These results are available from the authors on request.

²⁵The same qualitative result is reported without comment in Tables 3 to 5 of Hansen (2007).

²⁶We are extremely grateful to Joao Santos Silva for pointing out this mechanism to us.

pre and post treatment, and adding more time periods will reduce this variance. But this reduction will be small if serial correlation is high, and hence it can be dominated by the covariance effect.

An additional lesson, then, is that researchers who use OLS to perform DiD estimation should be very cautious about blindly using as many time periods as possible. With serially correlated error terms, this can result in a reduction in power. This issue does not arise with FGLS, which effectively transforms the data in a way that removes the serial correlation from the error terms.

In summary, as long as the number of time periods is not too small (approximately 10 or less), FGLS combined with robust inference is likely to offer substantial power gains; and even if it does not - because the error process is badly mis-specified - test size can still be reliably controlled, even with few groups. We therefore recommend this approach is used more routinely by applied researchers doing DiD estimation.

Table 10: Rejection rates for tests of nominal 5% size with placebo treatments in log-earnings data

	G=50	G=20	G=10	G=6
OLS, $\sqrt{G/(G-1)}$ -CRSEs, $t(G-1)$ critical values	0.045 (0.003)	0.041 (0.003)	0.042 (0.003)	0.052 (0.003)
FGLS	0.106 (0.004)	0.101 (0.004)	0.120 (0.005)	0.124 (0.005)
FGLS, $\sqrt{G/(G-1)}$ -CRSEs, $t(G-1)$ critical values	0.049 (0.003)	0.045 (0.003)	0.054 (0.003)	0.061 (0.003)
BC-FGLS	0.073 (0.004)	0.070 (0.004)	0.087 (0.004)	0.096 (0.004)
BC-FGLS, $\sqrt{G/(G-1)}$ -CRSEs, $t(G-1)$ critical values	0.049 (0.003)	0.045 (0.003)	0.058 (0.003)	0.065 (0.003)

The Table shows the proportion of the time that the null hypothesis of no treatment effect was rejected in 5000 Monte Carlo simulations. Simulation standard errors are reported in parentheses. The simulations resample groups (US states) from CPS-MORG data, having imposed the sampling restrictions described in the text. The treatment parameter has a true coefficient of zero. G is the number of sampled states. Data from 1979 to 2008 inclusive are sampled (i.e. T = 30). Regressions are run on aggregated state-year data. The different inference methods used are discussed in the text.

Table 11: Rejection rates for tests of nominal 5% size with a treatment effect of +0.05 in log-earnings data

	G=50	G=20	G=10	G=6
OLS, $\sqrt{G/(G-1)}$ -CRSEs, t(G-1) critical values	0.810 (0.006)	0.467 (0.007)	0.252 (0.006)	0.168 (0.005)
FGLS	0.985 (0.002)	0.799 (0.006)	0.573 (0.007)	0.434 (0.007)
FGLS, $\sqrt{G/(G-1)}$ -CRSEs, t(G-1) critical values	0.957 (0.003)	0.670 (0.007)	0.401 (0.007)	0.255 (0.006)
BC-FGLS	0.978 (0.002)	0.763 (0.006)	0.513 (0.007)	0.384 (0.007)
BC-FGLS, $\sqrt{G/(G-1)}$ -CRSEs, t(G-1) critical values	0.955 (0.003)	0.696 (0.007)	0.423 (0.007)	0.286 (0.006)

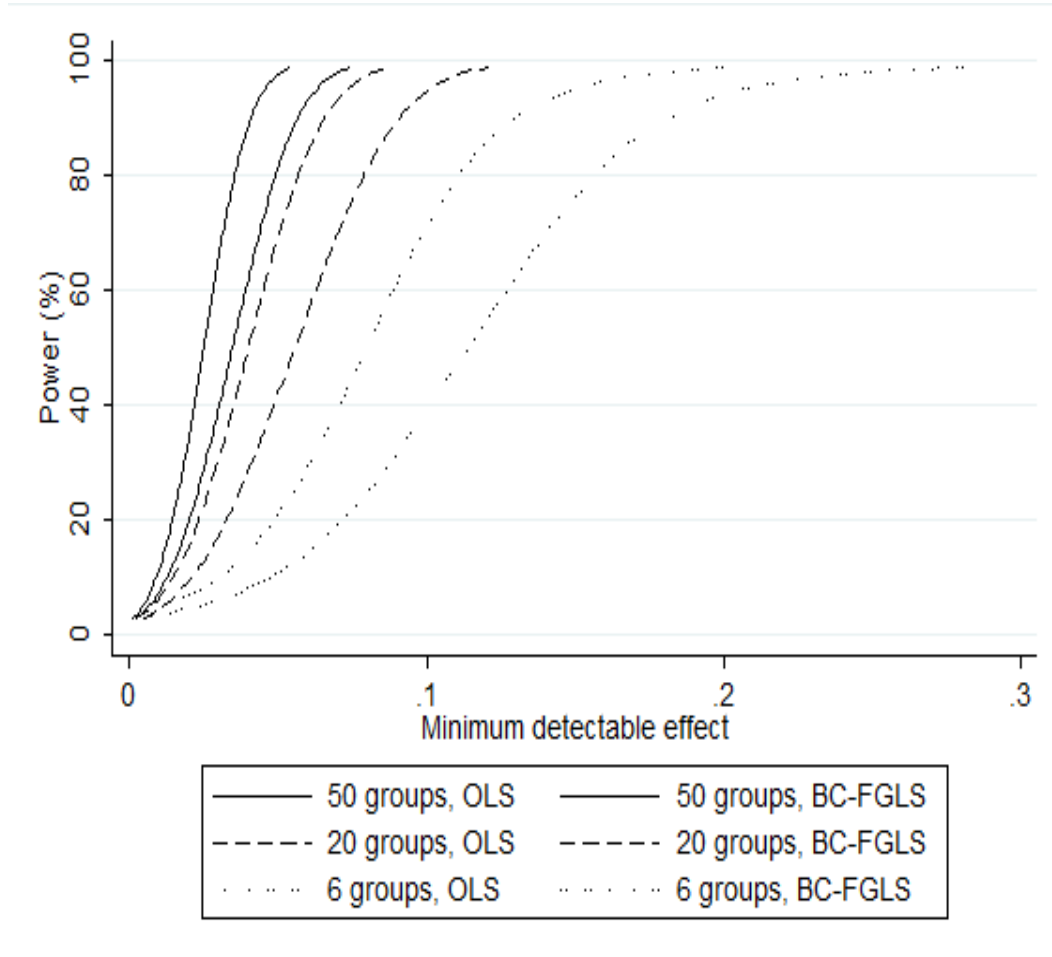
The Table shows the proportion of the time that the null hypothesis of no treatment effect was rejected in 5000 Monte Carlo simulations. Simulation standard errors are reported in parentheses. The simulations resample groups (US states) from CPS-MORG data, having imposed the sampling restrictions described in the text. The treatment parameter has a true coefficient of +0.05. G is the number of sampled states. Data from 1979 to 2008 inclusive are sampled (i.e. T = 30). Regressions are run on aggregated state-year data. The different inference methods used are discussed in the text.

Table 12: Rejection rates for tests of nominal 5% size with placebo treatments in employment data

	G=50	G=20	G=10	G=6
OLS, $\sqrt{G/(G-1)}$ -CRSEs, t(G-1) critical values	0.046 (0.003)	0.044 (0.003)	0.056 (0.003)	0.058 (0.003)
FGLS	0.182 (0.005)	0.171 (0.005)	0.202 (0.006)	0.201 (0.006)
FGLS, $\sqrt{G/(G-1)}$ -CRSEs, t(G-1) critical values	0.046 (0.003)	0.043 (0.003)	0.057 (0.003)	0.058 (0.003)
BC-FGLS	0.140 (0.005)	0.130 (0.005)	0.160 (0.005)	0.171 (0.005)
BC-FGLS, $\sqrt{G/(G-1)}$ -CRSEs, t(G-1) critical values	0.047 (0.003)	0.046 (0.003)	0.058 (0.003)	0.062 (0.003)

Notes as for Table 10.

Figure 3: Minimum detectable effects on log-earnings using $\sqrt{G/(G-1)}$ -CRSEs and t_{G-1} critical values and tests of size 0.05



The Figure shows the proportion of the time that the null hypothesis of no treatment effect is rejected when the treatment parameter has a true coefficient ranging from 0 to 0.3. Numbers are computed using the results of 100000 Monte Carlo simulations combined with equation 4, as described in the text. The simulations resample groups (US states) from CPS-MORG data, having imposed the sampling restrictions described in the text. Data from 1979 to 2008 inclusive are sampled (i.e. $T = 30$). Estimation of the treatment effect is conducted on aggregated state-year data either by OLS (reproducing part of Figure 1), or by feasible GLS assuming a AR(2) error process (homogeneous across states) and using bias-corrected AR parameter estimates as in Hansen, 2007 (denoted "BC-FGLS"). See text for full details.

Table 13: Rejection rates for tests of nominal 5% size with a treatment effect of +0.05 in employment data

	G=50	G=20	G=10	G=6
OLS, $\sqrt{G/(G-1)}$ -CRSEs, t(G-1) critical values	1.000 (.)	0.992 (0.001)	0.819 (0.005)	0.562 (0.007)
FGLS	1.000 (.)	1.000 (0.000)	0.988 (0.002)	0.934 (0.004)
FGLS, $\sqrt{G/(G-1)}$ -CRSEs, t(G-1) critical values	1.000 (.)	0.999 (0.000)	0.903 (0.004)	0.662 (0.007)
BC-FGLS	1.000 (.)	1.000 (0.000)	0.986 (0.002)	0.922 (0.004)
BC-FGLS, $\sqrt{G/(G-1)}$ -CRSEs, t(G-1) critical values	1.000 (.)	0.999 (0.000)	0.916 (0.004)	0.684 (0.007)

Notes as for Table 11.

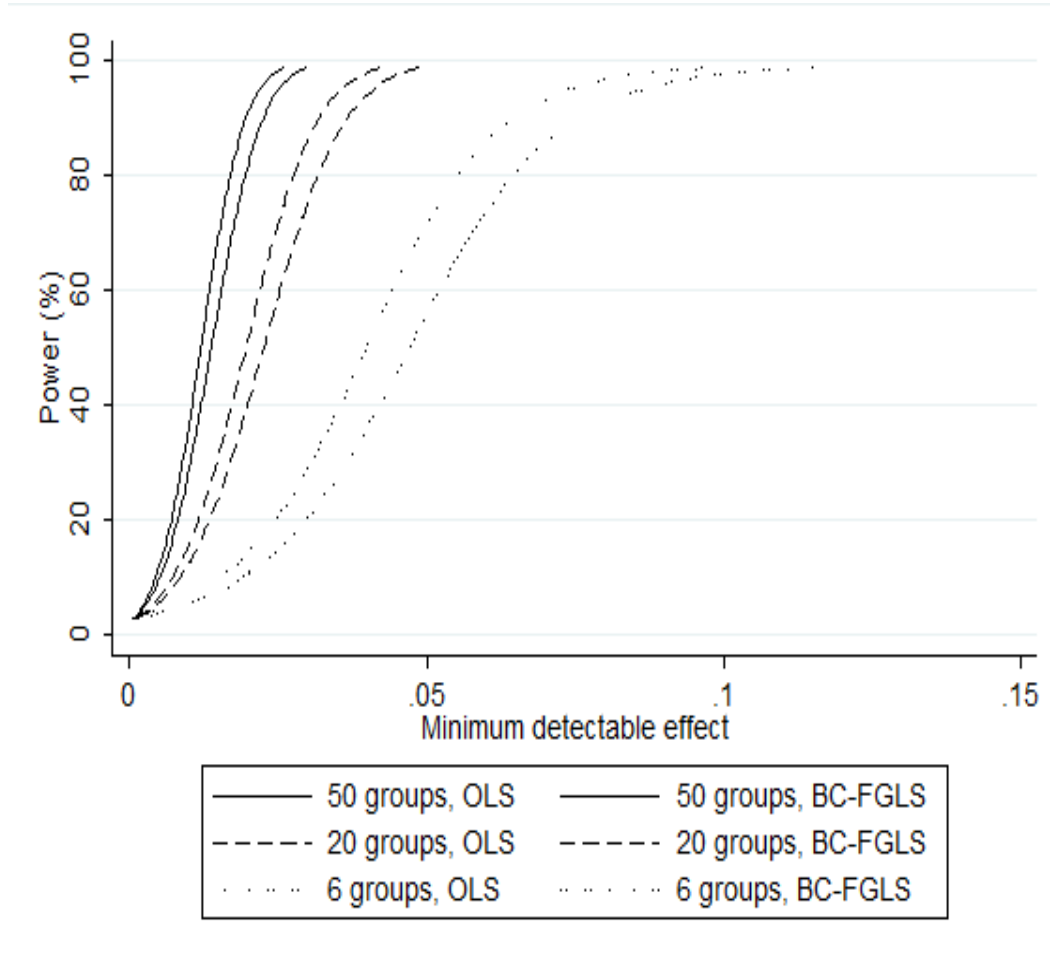
Table 14: Rejection rates for tests of nominal 5% size with placebo treatments and 10 groups in log-earnings data (simulated)

	CPS residuals	Heterogeneous AR(2)	MA(1)
OLS, $\sqrt{G/(G-1)}$ -CRSEs, t(G-1) critical values	0.049 (0.003)	0.040 (0.002)	0.052 (0.002)
FGLS	0.114 (0.004)	0.101 (0.003)	0.088 (0.003)
FGLS, $\sqrt{G/(G-1)}$ -CRSEs, t(G-1) critical values	0.054 (0.003)	0.055 (0.002)	0.051 (0.002)
BC-FGLS	0.081 (0.004)	0.072 (0.003)	0.072 (0.003)
BC-FGLS, $\sqrt{G/(G-1)}$ -CRSEs, t(G-1) critical values	0.056 (0.003)	0.059 (0.002)	0.052 (0.002)

The Table shows the proportion of the time that the null hypothesis of no treatment effect was rejected in 10000 Monte Carlo simulations. Simulation standard errors are reported in parentheses. The simulations resample groups (US states) from CPS-MORG data, having imposed the sampling restrictions described in the text. The treatment parameter has a true coefficient of zero. Data from 1979 to 2008 inclusive are sampled (i.e. $T = 30$).

Regressions are run on aggregated state-year data. The first column reproduces column 3 of table 10. Columns 2 and 3 show results after effectively replacing the empirical regression residuals with a simulated error term, generated according to an AR(2) process which varies across groups and an MA(1) process respectively. See text for full details.

Figure 4: Minimum detectable effects on employment rates using $\sqrt{G/(G-1)}$ -CRSEs and t_{G-1} critical values and tests of size 0.05



The Figure shows the proportion of the time that the null hypothesis of no treatment effect is rejected when the treatment parameter has a true coefficient ranging from 0 to 0.15. Numbers are computed using the results of 100000 Monte Carlo simulations combined with equation 4, as described in the text. The simulations resample groups (US states) from CPS-MORG data, having imposed the sampling restrictions described in the text. Data from 1979 to 2008 inclusive are sampled (i.e. $T = 30$). Estimation of the treatment effect is conducted on aggregated state-year data either by OLS (reproducing part of Figure 2), or by feasible GLS assuming a AR(2) error process (homogeneous across states) and using bias-corrected AR parameter estimates as in Hansen, 2007 (denoted "BC-FGLS"). See text for full details.

Table 15: Rejection rates for tests of nominal 5% size with a treatment effect of +0.05 and 10 groups in log-earnings data (simulated)

	CPS residuals	Heterogeneous AR(2)	MA(1)
OLS, $\sqrt{G/(G-1)}$ -CRSEs, t(G-1) critical values	0.254 (0.006)	0.510 (0.005)	0.604 (0.005)
FGLS	0.572 (0.007)	0.864 (0.003)	0.767 (0.004)
FGLS, $\sqrt{G/(G-1)}$ -CRSEs, t(G-1) critical values	0.399 (0.007)	0.723 (0.004)	0.591 (0.005)
BC-FGLS	0.518 (0.007)	0.843 (0.004)	0.736 (0.004)
BC-FGLS, $\sqrt{G/(G-1)}$ -CRSEs, t(G-1) critical values	0.423 (0.007)	0.741 (0.004)	0.593 (0.005)

The Table shows the proportion of the time that the null hypothesis of no treatment effect was rejected in 10000 Monte Carlo simulations. Simulation standard errors are reported in parentheses. The simulations resample groups (US states) from CPS-MORG data, having imposed the sampling restrictions described in the text. The treatment parameter has a true coefficient of +0.05. Data from 1979 to 2008 inclusive are sampled (i.e. $T = 30$). Regressions are run on aggregated state-year data. The first column reproduces column 3 of table 11. Columns 2 and 3 show results after effectively replacing the empirical regression residuals with a simulated error term, generated according to an AR(2) process which varies across groups and an MA(1) process respectively. See text for full details.

Table 16: Rejection rates for tests of nominal 5% size with placebo treatments and 10 groups in log-earnings data

	T=30	T=20	T=10
OLS, $\sqrt{G/(G-1)}$ -CRSEs, t(G-1) critical values	0.042 (0.003)	0.050 (0.003)	0.041 (0.003)
FGLS	0.120 (0.005)	0.131 (0.005)	0.107 (0.004)
FGLS, $\sqrt{G/(G-1)}$ -CRSEs, t(G-1) critical values	0.054 (0.003)	0.053 (0.003)	0.042 (0.003)
BC-FGLS	0.087 (0.004)	0.096 (0.004)	0.094 (0.004)
BC-FGLS, $\sqrt{G/(G-1)}$ -CRSEs, t(G-1) critical values	0.058 (0.003)	0.054 (0.003)	0.045 (0.003)

The Table shows the proportion of the time that the null hypothesis of no treatment effect was rejected in 5000 Monte Carlo simulations. Simulation standard errors are reported in parentheses. The simulations resample groups (US states) from CPS-MORG data, having imposed the sampling restrictions described in the text. The treatment parameter has a true coefficient of zero. T is the number of (consecutive) time periods (years). The first year of data is chosen from a uniform distribution between 1979 and (2009-T) in each Monte Carlo simulation. Regressions are run on aggregated state-year data. The different inference methods used are discussed in the text.

Table 17: Rejection rates for tests of nominal 5% size with a treatment effect of +0.05 and 10 groups in log-earnings data

	T=30	T=20	T=10
OLS, $\sqrt{G/(G-1)}$ -CRSEs, t(G-1) critical values	0.252 (0.006)	0.274 (0.006)	0.310 (0.007)
FGLS	0.573 (0.007)	0.545 (0.007)	0.460 (0.007)
FGLS, $\sqrt{G/(G-1)}$ -CRSEs, t(G-1) critical values	0.401 (0.007)	0.362 (0.007)	0.307 (0.007)
BC-FGLS	0.513 (0.007)	0.492 (0.007)	0.434 (0.007)
BC-FGLS, $\sqrt{G/(G-1)}$ -CRSEs, t(G-1) critical values	0.423 (0.007)	0.388 (0.007)	0.316 (0.007)

The Table shows the proportion of the time that the null hypothesis of no treatment effect was rejected in 5000 Monte Carlo simulations. Simulation standard errors are reported in parentheses. The simulations resample groups (US states) from CPS-MORG data, having imposed the sampling restrictions described in the text. The treatment parameter has a true coefficient of +0.05. T is the number of (consecutive) time periods (years). The first year of data is chosen from a uniform distribution between 1979 and (2009-T) in each Monte Carlo simulation. Regressions are run on aggregated state-year data. The different inference methods used are discussed in the text.

5 Summary and Conclusion

This paper contributes to a growing literature on inference in difference-in-differences designs with grouped errors. The literature has emphasized difficulties in obtaining correctly sized hypothesis tests, particularly with few groups, but our results suggest this is not the key challenge.

Using Monte Carlo evidence, we have made three main points. First, it is possible to obtain tests of the correct size, even with few groups, and in many settings this is possible using methods that are very straightforward to implement. Second, the main problem in difference-in-differences designs with grouped errors is instead low power to detect real effects. Third, feasible GLS estimation combined with robust inference methods can increase power considerably whilst maintaining correct test size - again, even with few groups. These findings have proven robust to a wide range of data generating processes.

We therefore recommend that applied researchers adopt GLS estimation combined with robust inference methods in practice. We also suggest that future research could usefully focus on improving power, as well as on getting test size correct, when using difference-in-difference designs.

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