# The synth\_runner Package: Utilities to Automate Synthetic Control Estimation Using synth

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Abstract. The Synthetic Control Methodology (Abadie and Gardeazabal, 2003; Abadie et al., 2010) allows for a data-driven approach to small-sample comparative studies. synth\_runner automates the process of running multiple synthetic control estimations using synth. It conducts placebo estimates in-space (estimations for the same treatment period but on all the control units). Inference (p-values) is provided by comparing the estimated main effect to the distribution of placebo effects. It allows several units to receive treatment, possibly at different time periods. Automatically generating the outcome predictors and diagnostics by splitting the pre-treatment into training and validation portions is allowed. Additionally, it provides diagnostics to assess fit and generates visualizations of results.

**Keywords:** st0001, synth\_runner, Synthetic Control Methodology, Randomization Inference

# 1 Introduction<sup>1</sup>

The Synthetic Control Methodology (SCM) (Abadie and Gardeazabal, 2003; Abadie et al., 2010) is a data-driven approach to small-sample comparative case-studies for estimating treatment effects. Similar to a difference-in-differences design, SCM exploits the differences in treated and untreated units across the event of interest. However, in contrast to a difference-in-differences design, SCM does not give all untreated units the same weight in the comparison. Instead, it generates a weighted average of the untreated units that closely matches the treated unit over the pre-treatment period. Outcomes for this synthetic control are then projected into the post-treatment period using the weights identified from the pre-treatment comparison. This projection is used as the counterfactual for the treated unit. Inference is conducted using placebo tests.

Along with their paper, Abadie et al. (2010) released the synth Stata command for single estimations. The synth\_runner package builds on top of that command to help conduct multiple estimations, inference, diagnostics, and generate visualizations of results. synth\_runner is designed to accompany synth but not supersede it. For more details about single estimations (variable weights, observation weights, covariate balance, and

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synthetic control outcomes when there are multiple time periods) use synth directly.

# 2 Synthetic Control Methodology

Abadie et al. (2010) posit the following data-generating process. Let  $D_{jt}$  be an indicator for treatment for unit j at time t. Next, let the observed outcome variable  $Y_{jt}$  be the sum of a time-varying treatment effect  $\alpha_{jt}D_{jt}$  and the no-treatment counterfactual  $Y_{jt}^N$ , which is specified using a factor model

$$Y_{jt} = \alpha_{jt} D_{jt} + Y_{jt}^{N}$$

$$= \alpha_{jt} D_{jt} + (\delta_t + \theta_t \mathbf{Z}_j + \lambda_t \mu_j + \varepsilon_{jt})$$
(1)

where  $\delta_t$  is an unknown time factor,  $\mathbf{Z}_j$  is a  $(r \times 1)$  vector of observed covariates unaffected by treatment,  $\theta_t$  is a  $(1 \times r)$  vector of unknown parameters,  $\lambda_t$  is a  $(1 \times F)$  vector of unknown factors,  $\mu_j$  is a  $(F \times 1)$  vector of unknown factor loadings, and the error  $\varepsilon_{jt}$  is independent across units and time with zero mean. Letting the first unit be the treated unit, the treatment effect is estimated by approximating the unknown  $Y_{1t}^N$  with a weighted average of untreated units

$$\hat{\alpha}_{1t} = Y_{1t} - \sum_{j>2} w_j Y_{jt}$$

Equation 1 simplifies to the traditional fixed effects equation if  $\lambda_t \mu_j = \phi_j$ . The fixed effects model allows for unobserved heterogeneity that is only time-invariant. The factor model employed by SCM generalizes this to allow for the existence of non-parallel trends between treated and untreated units after controlling for observables.

## 2.1 Estimation

To begin with, let there be a single unit that receives treatment. Let  $T_0$  be the number of pre-treatment periods of the T total periods. Index units  $\{1,...,J+1\}$  such that the first unit is the treated unit and the others are "donors". Let  $\mathbf{Y}_j$  be  $(T \times 1)$  the vector of outcomes for unit j and  $\mathbf{Y}_0$  be the  $(T \times J)$  matrix of outcomes for all donors. Let  $\mathbf{W}$  be a  $(J \times 1)$  observation-weight matrix  $(w_2, w_3, ..., w_{J+1})'$  where  $\sum_{j=2}^{J+1} w_j = 1$  and  $w_j \geq 0 \ \forall j \in \{2,...,J+1\}$ . A weighted average of donors over the outcome is constructed as  $\mathbf{Y}_0\mathbf{W}$ . Partition the outcome into pre-treatment and post-treatment vectors  $\mathbf{Y}_j = (\widetilde{\mathbf{Y}}_j \backslash \widetilde{\mathbf{Y}}_j)$ . Let  $\mathbf{X}$  represent a set of k pre-treatment characteristics ("predictors"). This includes  $\mathbf{Z}$  (the observed covariates above) and M linear combinations of  $\widetilde{\mathbf{Y}}$  so that k = r + M. Analogously, let  $\mathbf{X}_0$  be the  $(k \times J)$  matrix of donor predictors. Let  $\mathbf{V}$  be a  $(k \times k)$  variable-weight matrix indicating the relative significance of the predictor variables. Given  $\mathbf{Y}$  and  $\mathbf{X}$ , estimation of SCM consists of finding the optimal weighting matrices  $\mathbf{W}$  and  $\mathbf{V}$ . For a given  $\mathbf{V}$ ,  $\mathbf{W}$  is picked to minimize the RMSPE of the predictor variables,  $\|\mathbf{X}_1 - \mathbf{X}_0 \mathbf{W}\|_{\mathbf{V}}$ . In this way, the treated unit and its synthetic control look similar along

dimensions that matter for predicting pre-treatment outcomes. The inferential procedure is valid for any  $\mathbf{V}$ , but Abadie et al. (2010) suggest that  $\mathbf{V}$  be picked to minimize the prediction error of the pre-treatment outcome between the treated unit the synthetic control. Define distance measures  $\|\mathbf{A}\|_{\mathbf{B}} = \sqrt{\mathbf{A}'\mathbf{B}\mathbf{A}}$  and  $\|\mathbf{A}\| = \sqrt{\mathbf{A}'\cos(\mathbf{A})^{-1}\mathbf{A}}$ .  $\|\mathbf{\tilde{Y}}_1 - \mathbf{\tilde{Y}}_0\mathbf{W}\|$  is then the pre-treatment root mean squared prediction error (RMSPE) with a given weighted average of the control units. Define this as  $\mathbf{\tilde{s}}_1$  and define  $\mathbf{\tilde{s}}_1$  as the post-treatment RMSPE.  $\mathbf{V}$  is then picked to minimize  $\mathbf{\tilde{s}}_1$  (note that  $\mathbf{W}$  is a function of  $\mathbf{V}$ ).

If weights can be found such that the synthetic control matches the treated unit in the pre-treatment period:

$$\left\| \widetilde{\mathbf{Y}}_1 - \widetilde{\mathbf{Y}}_0 \mathbf{W} \right\| = 0 = \left\| \mathbf{Z}_1 - \mathbf{Z}_0 \mathbf{W} \right\|$$
 (2)

and  $\sum_{t=1}^{T_0} \lambda_t' \lambda_t$  is non-singular, then  $\hat{\alpha}_1$  will have a bias that goes to zero as the number of pre-intervention periods grows large relative to the scale of the  $\varepsilon_{jt}$ .

#### 2.2 Inference

After estimating the effect, statistical significance is determined by running placebo tests. Estimate the same model on each untreated unit, assuming it was treated at the same time, to get a distribution of "in-place" placebo effects. Disallow the actual treated unit from being considered for the synthetic controls of these other units. If the distribution of placebo effects yields many effects as large as the main estimate, then it is likely that the estimated effect was observed by chance. This non-parametric, exact test has the advantage of not imposing any distribution on the errors.

Suppose that the estimated effect for a particular post-treatment period is  $\hat{\alpha}_{1t}$  and that the distribution of corresponding in-place placebos is  $\hat{\alpha}_{1t}^{PL} = \{\hat{\alpha}_{jt} : j \neq 1\}$ . The two-sided p-value is then

$$p\text{-value} = \Pr\left(|\hat{\alpha}_{1t}^{PL}| \ge |\hat{\alpha}_{1t}|\right)$$
$$= \frac{\sum_{j \ne 1} 1(|\hat{\alpha}_{jt}| \ge |\hat{\alpha}_{1t}|)}{J}$$

and the one-sided p-values (for positive effects) are

$$p$$
-value =  $\Pr\left(\hat{\alpha}_{1t}^{PL} \ge \hat{\alpha}_{1t}\right)$ 

When treatment is randomized this becomes classical randomization inference<sup>2</sup>. If treatment is not randomly assigned, the p-value still has the interpretation of being the proportion of control units that have an estimated effect at least as large as that of the treated unit. Confidence intervals can be constructed by inverting the p-values for  $\hat{\alpha}_{1t}$ .

<sup>&</sup>lt;sup>2</sup>One may want to include  $\hat{\alpha}_{1t}$  in the comparison distribution as is common in randomization inference. This adds a one to the numerator and denominator of the *p*-value fraction. Abadie et al. (2015) and Cavallo et al. (2013), however, do not take this approach. With multiple treatments, there would be several approaches to adding the effects on the treated to the comparison distribution, so they are not dealt with here.

Care should be taken with these, however. As noted by Abadie et al. (2015), they do not have the standard interpretation when treatment is not considered randomly assigned.

To gauge the joint effect across all post-treatment periods Abadie et al. (2010) suggest using post-treatment RMSPE  $\vec{s}_1$ . In this case  $\vec{s}_1$  would be compared to the corresponding  $\vec{s}_1^{PL}$ .

The placebo effects may be quite large if those units where not matched well in the pre-treatment period. This would cause p-values to be too conservative. To control for this, one may want to adjust  $\hat{\alpha}_{jt}$  and  $\vec{s}_{j}$  for the quality of the pre-treatment matches. Adjustment can be achieved by two mechanisms:

- Restricting the comparison set of control effects to only include those that match well. This is done by setting a multiple m and removing all placebos j with  $\overline{s}_j > m\overline{s}_1$ .
- Dividing all effects by the corresponding pre-treatment match quality  $\vec{s}$  to get standardized (studentized) measures:  $\hat{\alpha}_{jt}/\vec{s}_{j}$  and  $\vec{s}_{j}/\vec{s}_{j}$ .

Inference can then be conducted over four quantities  $(\hat{\alpha}_{jt}, \vec{s}_j, \hat{\alpha}_{jt}/\vec{s}_j, \vec{s}_j/\vec{s}_j)$  and the comparison set can also be limited by the choice of m.

#### 2.3 Multiple Events

The extension by Cavallo et al. (2013) allows for more than one unit to experience treatment and at possibly different times. Index treated units  $g \in \{1...G\}$ . Let J be those units that never undergo treatment. For a particular treatment g, one can estimate an effect, say the first post-treatment period effect  $\hat{\alpha}_g$  (one could use any of the four types discussed above). We omit the t subscript as treatment dates may differ across events. Over all the treatments, the average effect is  $\bar{\alpha} = G^{-1} \sum_{g=1}^{G} \hat{\alpha}_g$ .

For each treatment g one generates a corresponding set of placebo effects  $\hat{\alpha}_g^{PL}$  where each untreated unit is thought of as entering treatment at the same time as unit g. If two treated units have the same treatment period, then their placebo sets will the be the same.

Averaging over the treatments to obtain  $\bar{\alpha}$  smooths out noise in the estimate. The same should be done in constructing  $\bar{\alpha}^{PL}$  the set of placebos against which the average treatment estimate is compared for inference. It should be constructed from all possible averages where a single placebo is taken from each  $\hat{\alpha}_g^{PL}$ . There are  $N_{\overline{PL}} = \prod_{g=1}^G J_g$  such possible averages<sup>3</sup>. Let i index a selection where a single placebo effect is chosen from each treatment placebo set. Let  $\bar{\alpha}^{PL(i)}$  represents the average of that placebo selection. Inference is now

$$\begin{split} p\text{-value} &= \Pr\left(|\bar{\alpha}^{PL}| \geq |\bar{\alpha}|\right) \\ &= \frac{\sum_{i=1}^{N_{\overline{PL}}} 1(|\bar{\alpha}^{PL(i)}| \geq |\bar{\alpha}|)}{N_{\overline{PL}}} \end{split}$$

<sup>&</sup>lt;sup>3</sup>The pool may be restricted by match quality. If  $J_g^m$  is the number of controls that match as well as treated unit g for the same time period, then  $N_{\overline{PL}}^m = \prod_{g=1}^G J_g^m$ .

## 2.4 Diagnostics

Cavallo et al. (2013) perform two basic checks to see if the synthetic control serves as a valid counterfactual. The first is to check if a weighted average of donors is able to approximate the treated unit in the pre-treatment. This should be satisfied if the treated unit lies within the convex hull of the control units. One can visually compare the difference in pre-treatment outcomes between a unit and its synthetic control. Additionally one could look at the distribution of pre-treatment RMSPE's and see what proportion of control units have values at least as high as that of the treated unit. Cavallo et al. (2013) discard several events from study as they can not be matched appropriately.

Secondly, one can exclude some pre-treatment outcomes from the list of predictors and see if the synthetic control matches well the treated unit in these periods.<sup>4</sup> As this is still pre-treatment, the synthetic control should match well. The initial section of the pre-treatment period is often designated the "training" period with the later part being the "validation" period. Cavallo et al. (2013) set aside the first half of the pre-treatment period as the training period.

## 3 The synth\_runner Package

The synth\_runner package contains several tools to help conduct SCM estimation. It requires the synth package which can be obtained from the SSC archive. The main program is synth\_runner, which is outlined here. Additionally, there are simple graphing utilities (effect\_graphs, pval\_graphs, single\_treatment\_graphs) that show basic graphs. These are explained in the following code examples and can be modified easily.

#### 3.1 Syntax

synth\_runner depvar predictorvars [, trunit(#) trperiod(#) d(varname) trends
pre\_limit\_mult(real) training\_propr(real) keep(file) replace ci pvals1s max\_lead(#)
n\_pl\_avgs(string) parallel deterministicout pred\_prog(string) drop\_units\_prog(string)
synthsettings ]

Post-estimation graphing commands are shown in the examples below. The syntax is similar to the synth command. New options include d, trends, pre\_limit\_mult, training\_propr, ci, pvals1s, n\_pl\_avgs. Options not explicitly matched will be passed to synth as synthsettings.

## 3.2 Settings

Required Settings:

- depvar the outcome variable.
- predictorvars the list of predictor variables. See help synth help for more details.

 $<sup>^4</sup>$ Note also that unless some pre-treatment outcome variables are dropped from the set of predictors, all other covariate predictors are rendered redundant. The optimization of V will put no weight on those additional predictors in terms of predicting pre-treatment outcomes.

For specifying the unit and time period of treatment, there are two methods. Exactly one of these is required.

- trunit(#) and trperiod(#). This syntax (used by synth) can be used when there is a single unit entering treatment.
- d(varname). The d variable should be a binary variable which is 1 for treated units in treated periods, and 0 everywhere else. This allows for multiple units to undergo treatment, possibly at different times.

### Options:

- trends will force synth to match on the trends in the outcome variable. It does this by scaling each unit's outcome variable so that it is 1 in the last pre-treatment period.
- pre\_limit\_mult( $real \ge 1$ ) will not include placebo effects in the pool for inference if the match quality of that control (pre-treatment RMSPE) is greater than  $pre\ limit\ mult$  times the match quality of the treated unit.
- training\_propr( $0 \le real \le 1$ ) instructs synth\_runner to automatically generate the outcome predictors. The default (0) is to not generate any (the user then includes the desired ones in predictorvars). If set to a number greater than 0, then that initial proportion of the pre-treatment period is used as a training period with the rest being the validation period. Outcome predictors for every time in the training period will be added to the synth commands. Diagnostics of the fit for the validation period will be outputted. If the value is between 0 and 1, there will be at least one training period and at least one validation period. If it is set to 1, then all the pre-treatment period outcome variables will be used as predictors. This will make other covariate predictors redundant.
- ci outputs confidence intervals from randomization inference for raw effect estimates. These should only be used if the treatment is randomly assigned. If treatment is not randomly assigned then these confidence intervals do not have the standard interpretation.
- pvals1s outputs one-sided p-values in addition to the two-sided p-values.
- keep(filename) saves a dataset with the results. This is only allowed if there is a single period in which unit(s) enter treatment. It is easy to merge this in the initial dataset. If keep(filename) is specified, it will hold the following variables:
  - panelvar contains the respective panel unit (from the tsset panel unit variable panelvar).
  - timevar contains the respective time period (from the tsset panel time variable timevar).

- lead contains the respective time period relative to the treatment period. Lead = 1 specifies the first period of treatment.
- effect contains the difference between the unit's outcome and its synthetic control for that time period.
- pre\_rmspe contains the pre-treatment match quality in terms of Root Mean
   Squared Predictive Error. It is constant for a unit.
- post\_rmspe contains a measure of the post-treatment effect (jointly over all
  post-treatment time periods) in terms of Root Mean Squared Predictive Error.
  It is constant for a unit.
- depvar\_scaled (if the match was done on trends) is the unit's outcome variable normalized so that its last pre-treatment period outcome is 1.
- effect\_scaled (if the match was done on trends) is the difference between
   the unit's scaled outcome and its scaled synthetic control for that time period.
- replace replaces the dataset specified in keep(filename) if it already exists.
- n\_pl\_avgs(string) controls the number of placebo averages to compute for inference. The total possible grows exponentially with the number of treated events. If omitted, the default behavior is cap the number of averages computed at 1,000,000 and if the total is more than that to sample (with replacement) the full distribution. The option n\_pl\_avgs(all) can be used to override this behavior and compute all the possible averages. The option n\_pl\_avgs(#) can be used to specify a specific number less than the total number of averages possible.
- max\_lead(int) will limit the number of post-treatment periods analyzed.
- parallel will enable parallel processing if the parallel command (at least version 1.18.1) is installed and configured.
- deterministicoutput when used with parallel, will eliminates displayed output that would vary depending on the machine (e.g. timers and number of parallel clusters) so that log files can be easily compared across runs.
- pred\_program is the name of a program that, when passed the period, will return via r(predictors) a synth-style predictor string. See Example 3.
- drop\_units\_progis the name of a program that, when passed the unit to be considered treated, will drop other units that should not be considered when forming the synthetic control. Commonly this is because they are neighboring or interfering units. See Example 3.
- synthsettings pass-through options sent to synth. See help synth for more information. The following which are disallowed: counit, figure, resultsperiod.

## 3.3 Saved Results

synth\_runner returns the following scalars and matrices:

•	_	9
S	Scalars	
	e(n_pl)	The number of placebo averages used for
		comparison.
	e(pval_joint_post)	The proportion of placebos that have a
		post-treatment RMSPE at least as large as the
		average for the treated units.
	e(pval_joint_post_t)	The proportion of placebos that have a ratio of
		post-treatment RMSPE over pre-treatment
		RMSPE at least as large as the average ratio for
		the treated units.
	e(avg_pre_rmspe_p)	The proportion of placebos that have a
		pre-treatment RMSPE at least as large as the
		average of the treated units. A measure of fit.
		Concerning if significant.
	e(avg_val_rmspe_p)	When specifying training_propr, this is the
		proportion of placebos that have a RMSPE for
		the validation period at least as large as the
		average of the treated units. A measure of fit.
		Concerning if significant.
N	Matrices	
	e(treat_control)	The average treatment outcome (centered around
		treatment) and the average of the outcome of
		those units' synthetic controls for the pre- and
		post-treatment periods.
	e(b)	A vector with the per-period effects (unit's actual
		outcome minus the outcome of its synthetic
		control) for post-treatment periods.
	e(pvals)	A vector of the proportions of placebo effects that
		are at least as large as the main effect for each
		post-treatment period.
	e(pvals_std)	A vector of the proportions of placebo
		standardized effects that are at least as large as

# 3.4 Example Usage

The following examples use a dataset from the synth package. Ensure that synth was installed with ancillary files (e.g., ssc install synth, all). This panel dataset contains

post-treatment period.

the main standardized effect for each

information for 39 US States for the years 1970-2000 (see Abadie et al. 2010 for details).

```
. sysuse smoking
(Tobacco Sales in 39 US States)
. tsset state year
    panel variable: state (strongly balanced)
    time variable: year, 1970 to 2000
    delta: 1 unit
```

#### Example 1

Reconstruct Example 1 from the synth help file (note this is not the exact estimation strategy use in Abadie et al. 2010):

Post-treatment results: Effects, p-values, p-values (psuedo t-stats)

	ı		pvals	•	
	-+				
lead1	1	-7.887098	.1315789	0	
lead2	1	-9.693599	.1842105	0	
lead3	1	-13.8027	.2105263	0	
lead4	1	-13.344	.1315789	0	
lead5	1	-17.0624	.1052632	0	
lead6	1	-20.8943	.0789474	0	
lead7	1	-19.8568	.1315789	.0263158	
lead8	1	-21.0405	.1578947	0	
lead9	1	-21.4914	.1052632	.0263158	
lead10	1	-19.1642	.1842105	.0263158	
lead11	1	-24.554	.1052632	0	
lead12	1	-24.2687	.1052632	.0263158	
. merge 1:1	st	tate vear u	sing `keepf	ile', nogene	rat

. merge 1:1 state year using `keepfile´, nogenerate
(label state already defined)

```
Result # of obs.
-----
not matched 0
matched 1,209
```

```
. gen cigsale_synth = cigsale-effect
```

The program notes progress towards estimating prediction errors and for conducting inference. Results for post-treatment periods are shown by default. In this example they are negative and significant by the standardized effect measure indicating that the law it studied likely had a negative effect on cigarette sales. We then merge in the synthetic control results and construct the effect variable.

While some of the return values are matrices and can be best visualized, some are scalars and easy to examine directly.

```
. ereturn list
scalars:
              e(n_p1) = 38
         e(n_pl_used) = 38
   e(pval_joint_post) = .1315789473684211
e(pval_joint_post_s
   td)
                      = 0
   e(avg_pre_rmspe_p) = .9210526315789474
macros:
        e(properties) : "b"
matrices:
                 e(b) : 1 \times 12
         e(pvals_std) : 1 x 12
             e(pvals) : 1 x 12
     e(treat_control) : 31 x 2
. //If truly random, can modify the p-value
. di (e(pval_joint_post_std)*e(n_pl)+1)/(e(n_pl)+1)
.02564103
```

The e(pval\_joint\_post) lists the proportion of effects from control units that have post-treatment RMSPE at least as great as the treated unit. While it is not significant, it's The return e(pval\_joint\_post\_std) lists the same, but scales all values by the relevant pre-treatment RMSPE. The final measure is a diagnostic measure and it notes that the treated unit was matched better than the majority of the control units. If the treatment is considered truly at random them the true p-value is a modification that adds one to the numerator and denominator (in cases with a single treatment). This is shown for the case of the ratio of post- to pre-treatment RMSPE.

Next we create common synthetic control graphs. The single\_treatment\_graphs command creates the graphs in Figure 1 (which are easy to do when there is a single treatment). The first graphs the outcome path of all units while the second graphs the prediction differences for all units. The effect\_graphs command creates the graphs in Figure 2. One plots the outcome for the unit and it's synthetic control while the other plots the difference between the two (which for post-treatment is the "effect"). The two previous graphing commands allow the option trlinediff(real) that allows the user to offset a vertical treatment from the first treatment period. Likely options include values in the range from (first treatment period - last post-treatment period) to 0 and the default value is -1. The pval\_graphs command creates the graphs in Figure 3. These plot the p-values per-period for post-treatment years for both raw and standardized effects.

Figure 1: Graphs from single\_treatment\_graphs

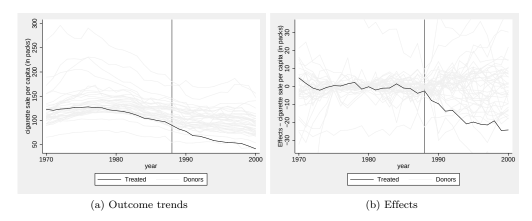


Figure 2: Graphs from effect\_graphs



. single\_treatment\_graphs, depvar(cigsale) trunit(3) trperiod(1989) ///
 trlinediff(-1) effects\_ylabels(-30(10)30) effects\_ymax(35) effects\_ymin(-35)

4

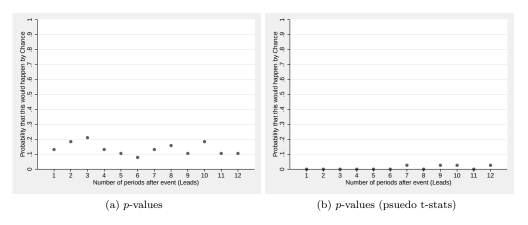
- . pval\_graphs

#### **▶** Example 2

Same treatment, but a bit more complicated setup:

- . gen byte D = (state==3 & year>=1989)
  . tempfile keepfile2
- . synth\_runner cigsale beer(1984(1)1988) lnincome(1972(1)1988) ///
   retprice age15to24, trunit(3) trperiod(1989) trends ///
   training\_propr(`=13/19´) pre\_limit\_mult(10) keep(`keepfile2´)
- . merge 1:1 state year using `keepfile2´, nogenerate
- . gen  $cigsale\_scaled\_synth = cigsale\_scaled effect\_scaled$
- . di "Proportion of control units that have a higher RMSPE than the"  $/\!/\!/$  "treated unit in the validtion period:"

Figure 3: Graphs from pval\_graphs



```
. di round(`e(avg_val_rmspe_p)', 0.001)
.842
. single_treatment_graphs, depvar(cigsale_scaled) ///
   effect_var(effect_scaled) trunit(3) trperiod(1989)
. effect_graphs , depvar(cigsale_scaled) trunit(3) trperiod(1989) ///
   effect_var(effect_scaled) depvar_synth(cigsale_scaled_synth)
. pval_graphs
```

Again there is a single treatment period, so output can be saved and merged back into the dataset. In this setting we (a) specify the treated units/periods with a binary variable, (b) generate the outcome predictors automatically using the initial 13 periods of the pre-treatment era (the rest is the "validation" period), (c) we match on trends, and (d) we limit during inference control units whose pre-treatment match quality more than 10 times worse than the match quality of the corresponding treatment units. Now that we have a training/validation period split there is a new diagnostic. It shows that 84% of the control units have a worse match (a higher RMSPE) during the validation period. The graphing commands are equivalent. The ones showing the range of effects and raw data are shown in Figure 4. One can see that all the lines converge on the last pre-treatment period as that is the unit that all are standardized by (and all the synthetic controls then match their real units and have zero prediction error).

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#### **▶** Example 3

Multiple treatments at different time periods:

```
cap program drop _gen_predictors
program _gen_predictors, rclass
args year
return local predictors "beer(`=`year´-4´(1)`=`year´-1´) . lnincome(`=`year´-4´(1)`=`year´-1´)"
end
cap program drop _drop_units
program _drop_units
args tunit
```

Figure 4: Graphs from single\_treatment\_graphs

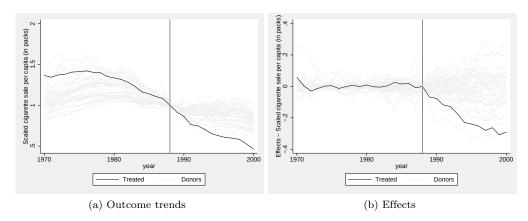
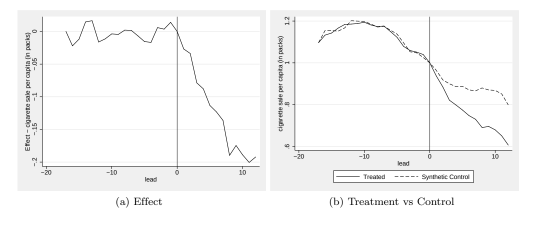


Figure 5: Graphs from effect\_graphs



```
if `tunit´==39 qui drop if inlist(state,21,38)
```

We extend Example 2 by considering a control state now to be treated (Georgia in addition to California). No treatment actually happened in Georgia in 1987. Now that we have several treatment periods we can not merge in a simple file. Some of the graphs (of single\_treatment\_graphs) can no longer be made. The option multi is now passed to effect\_graphs and those are shown in Figure 5.

4

<sup>`</sup>tunit'==3 qui drop if state==21 if

<sup>.</sup> end

gen byte D = (state==3 & year>=1989) | (state==7 & year>=1988)

<sup>.</sup> synth\_runner cigsale retprice age15to24, d(D) ///pred\_prog(\_gen\_predictors) trends training\_propr(`=13/18´) /// drop\_units\_prog(\_drop\_units)

<sup>.</sup> effect\_graphs , multi depvar(cigsale)

<sup>.</sup> pval\_graphs

## 4 Discussion

The Synthetic Control Methodology (SCM) (Abadie and Gardeazabal, 2003; Abadie et al., 2010) allows researchers to quantitatively estimate effects in many small sample settings in a manner grounded by theory. This article provides an overview of the theory of SCM and the synth\_runner package, which builds on the synth package of Abadie et al. (2010). synth\_runner provides tools to help with the common tasks of estimating a synthetic control model. It automates the process of conducting in-place placebos and calculating inference on the various possible measures. Following Cavallo et al. (2013) it (a) extends the initial estimation strategy to allow for multiple units that receive treatment (at potentially different times), (b) allows for matching on trends in the outcome variable rather than on the level, and (c) automates the process of splitting pre-treatment periods into "training" and "validation" sections. It also provides graphs of diagnostics, inference, and estimate effects.

# 5 References

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