

An Exercise in Data Analysis and Statistical Inference

Miles D. Williams

May 24, 2019

In this exercise, I (1) offer my best guess for the average treatment effect of the intervention and (2) summarize the evidence against the claim that the intervention had no effect. For both 1 and 2 I follow as closely as possible the standard operating procedure (SOP) suggested by Don Green’s lab at Columbia University. As the database specialist who compiled the data at hand is not available, and I do not have access to the pre-registered analysis plan (PAP), turning to Don Green’s SOP offers a standardized starting point for identifying appropriate estimators and methods for statistical inference.

1 Making Sense of the Data

The database specialist left few clues about this data before leaving. She left behind three datasets, one that contains baseline data on 152 individuals, one that contains information of the randomization design for the policy intervention, and one that contains endline data for two outcome variables (one continuous, Yc , and one binary, Yb). The baseline and endline data oddly have five duplicate rows at the bottom, which I removed prior to merging the data. Examination of the design dataset reveals that the randomized controlled trial (RCT) was done within blocks of varying sizes, with heterogeneous probability of treatment among blocks.

1.1 Identifying Variables

The baseline data contains 19 columns of covariates for each individual in the dataset. All are simply labeled with an X followed by a number (ranging from 1 to 19). All that is known initially is that $X1$ captures an individual’s “status,” which could refer to marital status, though it could also mean any number of things such as urban-rural status. $X2$ captures age, though $X3$ and $X4$ are nearly identical to $X2$. $X3$ seems to reflect an updated data draw on individuals’ ages, and $X4$ a final, more complete vector of ages. $X9$ and $X14$ both denote income; though, not the same income. One thought is that one of the variables is a corrected vector of income data, but another plausible guess is that these reflect dual incomes for a single household. $X8$ denotes groups, ranging from ‘a’ to ‘e’. $X11$ and $X19$ denote variables called `FISMOCheck` and `StatoFix2001` respectively. No description is given for what these variables are; though, my best guess is that `FISMOCheck` refers to a flexible single-master operation check for domain roles and naming on Windows, and values (D followed by a numerical value) probably represent domains relevant for certain aspects data transfer when the specialist was collecting/organizing the baseline dataset. It’s less clear to me what `StatoFix2001` is; however, it may have something to do with joining domains between a Windows operating system and some other operating system. The identities of the remaining covariates are unknown.

1.2 Dealing with Missing Data

Many of the covariates in the baseline dataset have missing values. Following the SOP, as less than 10 percent of the values in each variable is missing, I impute missing values for each of the numeric covariates with the average value for that variable among cases with non-missing data. I depart from the SOP, however, by using block-specific averages for imputed values. I did this because I noticed a considerable degree of clustering per block in values of many of the covariates. As I anticipate missing values for observations are likely to track with the block average, I rely on this value rather than the total sample average as a more accurate guess at the expected value of missing data.

Though I address missingness in baseline data as described above, attrition (missingness) remains an issue for each of the outcome variables in the endline data. There is a roughly 7.9 percent attrition rate for Yc , the continuous outcome variable, and a roughly 8.6 percent attrition rate for Yb , the binary outcome variable. An initial concern is whether attrition rates are significantly different than predicted by chance between treatment and control arms of the policy intervention, or among blocks. Following the analysis, I

conduct tests to assess whether attrition poses problems in drawing inferences about the average treatment effect (ATE).

2 Choice of Estimator and Adjustment Strategy

In considering choice of estimator and adjustment strategy, I refer to the SOP for guidance. The SOP relies on OLS regression as the default estimator for generating estimates for ATEs. This is a straightforward method of obtaining differences in means between treatment and control groups, so I rely on OLS as suggested.

In devising an adjustment strategy, I consider both the data at hand and the randomization design of the policy intervention. Intervention was randomized within blocks, with different treatment probabilities per block. Blocks 1, 2, and 3 only had 4 individuals each, with 2 (half) in each block randomly assigned treatment. Blocks 4, 5, and 6 similarly had half of the individuals assigned to treatment; though 8 individuals in total were in each. Blocks 7 to 9 had 12 individuals each, but only 1/3 of the subjects in each were given treatment. Finally, block 10, by far the largest, had 80 total individuals, only 5 percent of whom received treatment. The SOP advises two strategies for such a design. The first is to use OLS to regress the outcome on (1) a treatment indicator, (2) a set of block indicator variables with one block dropped to serve as a reference category, and (3) a set of treatment block interactions where the proportion of observations within each block is subtracted from its respective 0-1 block indicator. 3 is the equivalent of mean centering the block indicators.

The second approach is to estimate a least squares dummy variable (LSDV) regression—essentially an OLS model where the outcome is regressed on the treatment indicator and block indicators (minus a reference category) without interaction. The SOP recommends this approach be used under the extreme condition that, for at least one block j , the following inequality holds:

$$\frac{N_j}{\sum_j N_j} > 20 \cdot \frac{N_j P_j (1 - P_j)}{\sum_j N_j P_j (1 - P_j)}.$$

In the above, N_j denotes the number of subjects per block j and P_j the probability of treatment per block j .

To determine which of these two approaches to use, I calculate the left and right sides of the above inequality for each of the 10 blocks included in the data. After accounting for attrition, I find that in 2 blocks, this inequality is met. I therefore rely on the second approach, estimating the treatment effect without treatment-block interactions.

I further have several baseline covariates per observation. Though in theory randomization within blocks should be independent of both observed and unobserved unit characteristics, adjusting for covariates in estimating the average treatment effect is justified on the basis that including covariates that are strongly correlated with the outcome, regardless of their association with treatment assignment, help to improve statistical power. The SOP offers guidelines for how to adjust for covariates in the analysis, as well as the number of covariates to include.

The first thing to consider is the number of individuals assigned to treatment. Let M denote individuals assigned to the treatment arm. If $M \geq 20$, the SOP recommends adjusting for covariates by regressing the outcome on (1) the treatment indicator, (2) the covariates, and (3) interactions between the treatment indicator and mean-centered values of the covariates.

If $M < 20 \leq N$, where N is the total number of observations, the SOP recommends regressing the outcome on items 1 and 2 described above, and not including treatment covariate interactions.

Finally, in the extreme case where $N < 20$, the SOP recommends estimating only the difference in means between control and treatment groups.

After attrition rates are accounted for, $N = 140$ for the continuous outcome, with $M = 32$, while $N = 139$ with $M = 32$ for the binary outcome. The SOP therefore calls for the first adjustment strategy, that is, regressing the outcome on the treatment indicator, covariates, and treatment-covariate interactions (with covariates mean-centered).

Finally, the number and choice of covariates to adjust for needs to be justified. The SOP recommends including no more than $M/20$ covariates when using interactions and no more than $N/20$ when not using interactions. As I have decided to go with the interaction adjustment strategy, I restrict the number of

covariates to no more than $M/20 = 1.6$. Since I can’t include 6/10 of a covariate, I include only 1. This procedure for limiting covariates, in addition to helping to simplify the analysis, helps to steer practitioners clear of the temptation to use “kitchen sink” regressions.

However, restricting the number of covariates to 1 means I must be extra choosy in identifying the best covariate to include. The SOP recommends using covariates highly correlated with the outcome, regardless of correlation with treatment. Some basic bivariate correlations show that **X4**, subject age, is highly correlated with both the continuous and binary outcomes. Pearson’s ρ equals 0.51 for age and the continuous outcome and -0.52 for age and the binary outcome. These estimated ρ s are much larger relative to any of the other covariates in the data, suggesting age may be the best choice among baseline covariates. To help confirm this choice, I use lasso regression, a penalized regression model that “shrinks” coefficient values toward zero, to eliminate variables less able to predict the outcomes. I regress each of the outcomes on all baseline covariates (save for the duplicate age variables). As this approach provides slightly different results with each iteration, I run a lasso regression 50,000 times and calculate the proportion of times a variable is not eliminated in the shrinkage process. **X4** and **X9** are the only variables to never be eliminated in any of the 50,000 iterations for the continuous outcome. **X4** and several other covariates are not eliminated for the binary outcome. For the former, to adjudicate between **X4** and **X9** I estimate OLS models with the outcome regressed only on an intercept, and then on either **X4** or **X9**. I use a Wald test to compare the fit of these models. I find that while both significantly improved fit, **X4** improves fit far better than **X9**. For the binary outcome, I use a similar procedure. Among the several variables that survived the lasso, **X4**, again, yields the most substantial improvement relative to other covariates. I therefore use **X4** as a control variable for both the continuous and binary outcome variables.

Given the above discussion, I generate ATE estimates using the following specifications to be estimated by OLS:

$$\ln(Y_i^c) = \beta_1 z_i + \beta_2 x_i + \beta_3 z_i \cdot (x_i - \bar{x}_j) + \mathbf{B}\alpha + \varepsilon_i, \quad (1)$$

$$Y_i^b = \gamma_1 z_i + \gamma_2 x_i + \gamma_3 z_i \cdot (x_i - \bar{x}_j) + \mathbf{B}\eta + v_i, \quad (2)$$

where β_1 and γ_1 denote the ATE for the block-randomized policy intervention z , denoted by **Zdesign** in the data, on the continuous and binary outcome variables respectively. x_i is subject age in years and \bar{x}_j is the average age of subjects per block j . I mean-center age on the block-specific average rather than the total average so that estimates reflect the ATE per block. **B** is a vector of block indicators where I drop block $j = 1$ to serve as the reference category.

Though not recommended by the SOP, I log-transform Y_i^c in equation 1. The continuous outcome variable has a highly skewed distribution. Values range from 1.24 to 67.16, though the mean is 7.28 and the median is 3.56. This skewness poses challenges for straightforwardly estimating the average treatment effect. Log-transforming values helps to overcome this issue without loss of data or the need to resort to methods such as robust regression. It further permits substantive interpretation of the ATE, which would not be easily facilitated by other approaches, such as a rank-based transformation. This choice does slightly change the interpretation of the estimand for the treatment affect, however. As this model has the functional form of a log-linear model, with some adjustment the estimated parameter on the treatment variable can be interpreted as the percent change in the outcome given treatment. More precisely, $\% \Delta Y_i^c = 100 \cdot (e^{\beta_1} - 1)$.

Turning to equation 2, the outcome variable is binary, which might make a model-based estimator such as logit an option vis-à-vis OLS. However, as numerous studies have shown, OLS estimates for a binary outcome are robust in the context of a RCT. I refer the reader to the SOP of Don Green’s lab for relevant citations.

3 Statistical Inference

To generate standard errors for the ATE, I rely on HC2 robust standard errors, or Bell-McCaffrey standard errors, without clustering. While OLS estimates are unbiased in the face of heteroskedasticity, OLS standard errors are not robust to non-constant variance in the data and may underestimate the size of coefficient standard errors when such violations of OLS assumptions arise. This leads to an increase in the probability of a false positive, or the type I error rate. The HC2 estimator for coefficient estimates imposes less restrictive assumptions on variance, and therefore HC2 estimates are more reliable. For this reason, the HC2 estimator

is the default recommendation of the SOP.

Though this study relies on block-randomization, I do not cluster standard errors by block. The SOP recommends using clustering only in panel settings, when multiple observations exist for a single subject or, for example, when treatment is given to an entire household with multiple members of that household included as subjects in the study. Neither is the case with this data. Further, the HC2 estimator will be undefined if a block-indicator equals 1 for at least a single block and 0 otherwise. As I rely on block indicators, clustering on blocks will preclude estimation of HC2 standard errors.

After estimating the ATE and its standard error, I use these values in generating evidence against the claim that the policy intervention had no effect. I do this by comparing the estimated t-statistic (the ratio of the ATE to its standard error) to its empirical distribution under random reassignment of treatment. I generate this empirical distribution using a Studentized permutation test. I first simulate random reassignment of treatment within blocks 10,000 times. For each of these simulated reassignments, I obtain an estimated ATE and HC2 standard error, which I then use to estimate a permuted t-statistic. I collect each of these 10,000 t-statistics and compare their distribution to the t-statistic I originally estimated. Using a two-sided test, I calculate the p-value, or the probability of observing the t-statistic I originally estimated under random reassignment of treatment. Consistent with standard practice, I use a p-value less than 0.05 as the threshold for rejecting the hypothesis of no treatment effect.

My preferred method for calculating the p-value from a permutation test is to estimate the proportion of times that $|t| \leq |t_p|$, where t is the originally calculated t-statistic and t_p is the p^{th} permuted t-statistic. The SOP, however, recommends calculating the one-sided p-value for the left and right sides of the originally calculated t-statistic and doubling the value of the smaller of the two. In the interest of following a standardized procedure, I rely on the SOP’s preferred method.

4 Results

Using the methods outlined in the preceding sections, I (1) generate estimates of the policy intervention’s effect on both the continuous and binary outcome variables and (2) provide evidence against the claim that the intervention had no effect. Table 1 summarizes the results. For the visually inclined reader, Figure 1 plots the estimated ATE with 95% confidence intervals, calculated based on the HC2 standard errors.

The first column in Table 1 indicates whether the estimates are for the continuous or binary outcome variable. The second column shows OLS estimates of the ATE. The third and fourth columns show the estimated HC2 standard errors and calculated t-statistics respectively. Finally, column five shows the p-values calculated based on the Studentized permutation test described in the previous section.

The ATE for the continuous variable is 0.519. This value denotes the average within-block difference in the logged outcome between treatment and control arms of the study, holding subject age constant at the block-specific average. This estimate equates to the following percent difference in the outcome given treatment: $100 \cdot [e^{0.519} - 1] = 68.03\%\Delta$.

The t-statistic for this estimate is roughly 3.2. Based on the permutation test, the probability of observing a statistic as extreme as that observed is quite small (0.002, or 1 in 500). This means there is little evidence against the claim that the treatment had no effect.

The ATE for the binary variable is 0.494. This value denotes the average within-block difference in the proportion of occurrences (1 values) of the binary outcome between treatment and control arms of the study, holding subject age constant at the block-specific average.

The t-statistic for this ATE is about 6.34. According to the permutation test, the probability of observing a statistic as extreme as this by random chance is practically zero. After 10,000 iterations of the permutation test, I failed to observe a t-statistic as large as the one estimated for the ATE.

Table 1: OLS estimates of treatment effects

Outcome	ATE	SE	t-statistic	p-value
Continuous (ln)	0.519	0.164	3.173	0.002
Binary	0.494	0.078	6.343	0.000

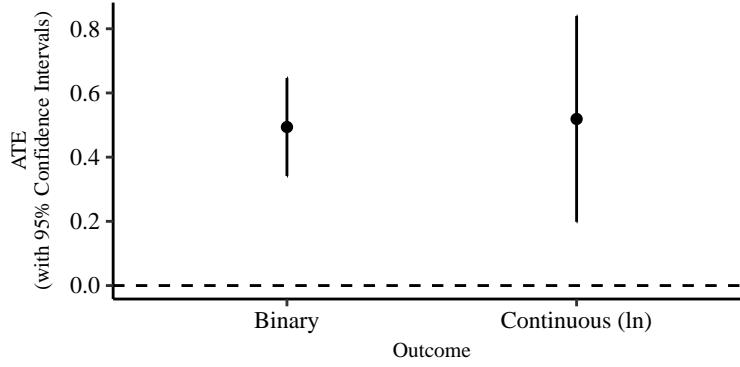


Figure 1: Coefficient plot of the average treatment effect (ATE) of the policy intervention on the log of the continuous outcome and on the binary outcome. Parametric 95% confidence intervals are shown, the values of which were estimated using calculated HC2 standard errors

As shown in Figure 1, the 95 percent confidence intervals fail to intersect with zero. This further suggests there is little evidence that the true ATE is zero as we should expect the true ATE to fall within the confidence intervals 95 percent of the time.

5 Checks for Infelicities in Design

As I mentioned in a previous section where I described missing values in the data, both outcome variables had non-zero attrition rates. Though less than 10 percent of observations had missing data on each outcome, if there are asymmetries in the distribution of missing values between arms of the trial, among blocks, or in subject age (a particularly strong predictor of both outcomes), this might lead us to worry that attrition will bias the results of the study.

As a check against the possibility that infelicities in the research design might bias estimates of the ATE, I follow the SOP's recommendation of conducting a Studentized permutation test for the F-statistic in a regression model where the outcome is a binary indicator for whether data on the outcome is missing. I specify the right-hand side of these models exactly as those used to generate ATEs for the intervention. I begin by estimating the heteroskedasticity robust Wald (F) statistic for each regression model. I then simulate 10,000 treatment reassignments within blocks and estimate new F-statistics. I then compare the original F-statistic against the distribution of F-statistics under random reassignment of treatment. If the probability of observing a F-statistic as extreme as that estimated for the observed data is less than 0.05, I consider this evidence against the null hypothesis of symmetric attrition. Using this procedure, in the case of both outcome variables, I calculate a p-value larger than this threshold, 0.75 for the continuous outcome and 0.6 for the binary outcome. This offers some assurance that asymmetric attrition should not be a major issue.

Another issue with design may be uncovered if covariate imbalances between treatment arms are greater than would be expected by chance. This might be evidence that treatment assignment was not implemented at random as intended. I therefore follow the SOP and conduct a Studentized permutation test similar to that described above, save that the outcome now is an indicator for whether an observation received treatment. I regress treatment on subject age as well as block and calculated the heteroskedasticity robust F-statistic, which I compare with the empirical distribution of the F-statistic under random reassignment of treatment. Again, if the p-value is less than 0.05, I consider this evidence against the null hypothesis of random treatment assignment. Using this procedure, I fail to reject the null with a p-value of approximately 0.2.