1. Potential outcomes and treatment effects

LPO 8852: Regression II

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What you learned in Regression I

The mechanics and properties of linear regression models:

$$Y_{i} = \beta_{0} + \beta_{1} X_{i1} + \beta_{2} X_{i2} + \dots + \beta_{k} X_{ik} + u_{i}$$

- Model specification and interpretation
- Estimation (e.g., OLS, WLS)
- Inference: What is the standard error of your estimator? What is the estimator's sampling distribution in finite samples? In large samples? Knowledge of the sampling distribution is needed to construct valid confidence intervals and conduct hypothesis tests.

Model interpretation and statistical inference rely heavily on assumptions.

What you learned in Regression I

When I first learned econometrics. I often felt dissatisfied:

- Assumptions feel implausible
- How do we know the model is "correct"?
- There are always "omitted variables"!
- Causal interpretation feels like a pipe dream.

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Regression II

Research designs for causal inference

- When can a regression be interpreted as causal?
- What does it mean for an estimator to have a causal interpretation?
- What research designs—which may or may not use regression—make a strong case for causal interpretation?

We will consider:

- Matching and weighting estimators
- Panel data models (e.g., fixed effects)
- Difference-in-differences
- Synthetic control methods
- Instrumental variables
- Regression discontinuity

What is a causal effect?

A **causal effect** is a change in some outcome (Y) that is the result of a change in some other (manipulable) factor (X).

For simplicity, assume the factor X is a binary (0-1) variable. Example: the causal effect of taking an aspirin on headache pain, or the effect of getting a vaccine on the likelihood of contracting COVID-19.

Causal effects involve a **counterfactual** comparison between two different states of the world: e.g., Y whenever X = 1 versus Y whenever X = 0 (where nothing else is manipulated). For convenience we often refer to X as a **treatment**, even when not an "administered" treatment.

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Potential outcomes

The **potential outcomes framework** is useful for thinking about counterfactual comparisons and treatment effects. This approach is attributed to Neyman (1923) and Rubin, who later generalized the framework. It is often referred to as the **Neyman-Rubin causal model**.

Potential outcomes

Let D_i be a dichotomous indicator of a "treatment" where $D_i=1$ means unit i is "treated" and $D_i=0$ means i is "not treated." For every i there are two **potential outcomes**:

- $Y_i(1)$ or $Y_{i1} = \text{outcome when } D_i = 1$
- $Y_i(0)$ or $Y_{i0} = \text{outcome when } D_i = 0$

Note these are "all else equal" outcomes in the sense that nothing else is manipulated.

We call these potential outcomes since units are not observed in more than one state at the same time. This is the *fundamental problem of causal inference* (Holland, 1986).

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SUTVA

A common assumption invoked here is **SUTVA** (stable unit treatment variable assignment). What this says is that unit *i*'s potential outcomes do not depend on the treatment assignment of other units. Cases in which this could be violated:

- Spillovers from treated to untreated units (e.g., treatments for infectious disease, classroom peer effects, knowledge spillovers)
- "General equilibrium effects"

Violations of SUTVA create problems for what comes next. We'll ignore this possibility for now, but researchers should pay more attention to this assumption.

Potential outcomes

The observed Y_i is either $Y_i(0)$ or $Y_i(1)$:

$$Y_i = D_i Y_i(1) + (1 - D_i) Y_i(0)$$

Call the above the switching equation.

A **counterfactual** is the outcome for the unit in the other (hypothetical, unobserved) state. E.g., the counterfactual for $\underline{\text{treated}}$ i would be $Y_i(0)$.

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Example 1: job training program

Person	Di	$Y_i(0)$	$Y_i(1)$	Yi
1	1	10	14	14
2	1	8	12	12
3	1	12	16	16
4	1	8	12	12
5	1	6	10	10
6	1	4	8	8
7	0	4	8	4
8	0	6	10	6
9	0	8	12	8
10	0	4	8	4
11	0	10	14	10
12	0	8	12	8
13	0	2	6	2
14	0	1	5	1
Mean	0.429	6.5	10.5	8.2

Source: Jennifer Hill (2011) lecture notes. Assume Y_i is earnings and D_i indicates participation in job training program.

Treatment effects

The causal effect of D on Y for individual i (the **treatment effect**) is:

$$\tau_i = Y_i(1) - Y_i(0)$$

We can't know the τ_i for any individual, but we may be able to estimate an <u>average</u> of τ in some population, or some other information about the distribution of those τ s.

This information is useful for predicting what the effect might be for some other *i* (e.g., for policy and practice decisions)

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Treatment effects

We are often interested in the population average treatment effect (ATE):

$$ATE = E(\tau) = E[\underline{Y(1) - Y(0)}]$$

Or the average treatment effect on the treated (ATT):

$$ATT = E(\tau|D=1) = E[Y(1)|D=1] - \underbrace{E[Y(0)|D=1]}_{\text{not observed}}$$

Recall $E[\cdot]$ is the expectations operator (population average).

Treatment effects

Or the average treatment effect on the untreated (ATU):

$$ATU = E(\tau|D=0) = \underbrace{E[Y(1)|D=0]}_{\text{not observed}} - E[Y(0)|D=0]$$

The ATE, ATT, and ATU are **estimands**—quantities of interest in the population. Researchers are often most interested in ATT or ATE.

Note the ATE is just a weighted average of the ATT and ATU:

$$ATE = pATT + (1 - p)ATU$$

where p is the proportion treated.

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Example 1: job training program

Person	Di	$Y_i(0)$	$Y_i(1)$	Y_i
1	1	10	14	14
2	1	8	12	12
3	1	12	16	16
4	1	8	12	12
5	1	6	10	10
6	1	4	8	8
7	0	4	8	4
8	0	6	10	6
9	0	8	12	8
10	0	4	8	4
11	0	10	14	10
12	0	8	12	8
13	0	2	6	2
14	0	1	5	1
Mean	0.429	6.5	10.5	8.2

In Example 1 there are constant treatment effects:

$$ATE = ATT = ATU = 4$$

Estimating treatment effects

Suppose we calculate the simple differences in mean observed Y for two groups, D=1 and D=0 (a "naïve" estimator):

$$E[Y(1)|D=1] - E[Y(0)|D=0] = E[Y(1)|D=1] - E[Y(0)|D=0] - \underbrace{E[Y(0)|D=1] + E[Y(0)|D=1]}_{0}$$

$$E[Y(1)|D=1]-E[Y(0)|D=0] = ATT + \underbrace{E[Y(0)|D=1] - E[Y(0)|D=0]}_{\text{selection bias}}$$

Selection bias reflects differences in Y(0) between the treated and untreated group ("baseline differences" or "unobserved heterogeneity").

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Example 1: job training program

Person	Di	Educ.	Age	Y(0)	Y(1)	Y
1	1	1	26	10	14	14
2	1	1	21	8	12	12
3	1	1	30	12	16	16
4	1	1	19	8	12	12
5	1	0	25	6	10	10
6	1	0	22	4	8	8
Mean $(D=1)$	1	0.67	23.8	8	12	12
7	0	0	21	4	8	4
8	0	0	26	6	10	6
9	0	0	28	8	12	8
10	0	0	20	4	8	4
11	0	1	26	10	14	10
12	0	1	21	8	12	8
13	0	0	16	2	6	2
14	0	0	15	1	5	1
Mean $(D=0)$	0	0.25	21.6	5.4	9.4	5.4

Estimating treatment effects

In Example 1, $\tau = 4$ for everyone. But:

$$E[Y(1)|D=1] - E[Y(0)|D=0] = 12.0 - 5.4 = 6.6$$

$$E[Y(1)|D=1]-E[Y(0)|D=0] = ATT + \underbrace{E[Y(0)|D=1] - E[Y(0)|D=0]}_{\text{selection bias}}$$

$$12.0 - 5.4 = 4.0 + \underbrace{8.0 - 5.4}_{\text{selection bias}} = 6.6$$

The treated group has a higher Y(0) than the untreated group. This could be due to their higher average education and age (shown in the previous table), two things associated with higher earnings. Their Y would have been higher on average even in the absence of treatment.

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Estimating treatment effects

Think of $Y_i(0)$ as shorthand for everything about unit i other than their treatment status. Comparing mean covariates can be revealing about differences in the treated and untreated groups.

"When observed differences proliferate, so should our suspicions about unobserved differences" (Mastering Metrics).

Estimating treatment effects

Note the "naïve" estimator also generally fails to recover the ATE:

$$E[Y(1)|D = 1] - E[Y(0)|D = 0] = ATE + \underbrace{E[Y(0)|D = 1] - E[Y(0)|D = 0]}_{\text{selection bias}} + \underbrace{(1 - p)(ATT - ATU)}_{\text{for the properties}}$$

See *Mixtape* Potential Outcomes chapter for the algebra. In Example 1, ATT=ATU (constant treatment effect), so there is no heterogeneous treatment effect bias. (There is, however, selection bias).

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Heterogeneous treatment effects

In Example 1, ATT = ATU = ATE. In practice, ATT and ATU often differ from the ATE because units endogenously sort into treatments based on gains they expect from it. We might expect ATT > ATE > ATU.



Conditional and marginal treatment effects

Other treatment effects that are often of interest are **conditional treatment effects** (ATE, ATT or ATU). That is, the average treatment effect conditional on something else being true.

$$ATE|X = E(\tau|X) = E[Y(1) - Y(0)|X]$$

In Example 1, we might be interested the ATE conditional on Education = 0. or, the ATT conditional on age > 26.

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The experimental ideal

Under what conditions will selection bias be zero? When treatment assignment is **independent** of potential outcomes ("strong ignorability"):

$$(Y_i(1), Y_i(0)) \perp \!\!\! \perp \!\!\! D$$

One case where this holds is **randomization** to treatment. Under random assignment, $E[Y_i(0)|D_i=1]=E[Y_i(0)|D_i=0]$. The D=0 and D=1 groups are random draws from the same population. The untreated D=0 can reasonably "stand in" as a counterfactual for the treated D=1.

Note under random assignment, there is no expected heterogeneous treatment effect bias (ATT=ATU). So the mean difference in outcomes between D=0 and D=1 should give us the ATE, ATT, and ATU.

Conditional independence assumption

In the absence of randomization, it may be that treatment assignment is independent of potential outcomes *conditional* on some X:

$$(Y_i(1), Y_i(0)) \perp \!\!\! \perp \!\!\! D | X$$

In other words, i's with the same X have the same distribution of Y(1) and Y(0).

This is the **conditional independence assumption** (or CIA), also known as **selection on observables**. A big assumption, but may not be unreasonable in some circumstances. We'll come back to this in Lecture 2.

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Regression and causality

What does this have to do with regression? We often hope that regression will help us estimate average treatment effects. Suppose we estimate the following simple regression with the hope of estimating the causal effect of D on Y:

$$Y_i = \beta_0 + \beta_1 D_i + u_i$$

When will this regression have a causal interpretation?

★ Answer: When it describes differences in average potential outcomes for a reference population of interest.

Let's express β_1 in terms of mean potential outcomes. In large samples, you already know β_1 will consistently estimate:

$$\beta_1 = E[Y_i|D_i = 1] - E[Y_i|D_i = 0]$$

Which is the same as:

$$\beta_1 = E[Y(1)|D=1] - E[Y(0)|D=0]$$

Is this the parameter we care about? Does it represent differences in average potential outcomes for a population of interest? Does it estimate a treatment effect for a population of interest?

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Regression and causality

Farlier we saw:

$$E[Y(1)|D=1]-E[Y(0)|D=0] = ATT + \underbrace{E[Y(0)|D=1] - E[Y(0)|D=0]}_{\text{selection biss}}$$

and:

$$E[Y(1)|D=1] - E[Y(0)|D=0] = ATE + \underbrace{E[Y(0)|D=1] - E[Y(0)|D=0]}_{\text{selection bias}} + \underbrace{(1-\rho)(ATT-ATU)}_{\text{heterogeneous treatment effect bias}$$

So, \underline{no} : β_1 will not generally give us a treatment effect we care about! A larger sample size will not help.

We also saw that:

- If D_i is randomly assigned, this difference in population means corresponds to the ATE: E[Y(1) - Y(0)] (and the ATT, ATU).
- Under randomization, the regression <u>does</u> reveal a difference in potential outcomes for a population of interest.
- Without random assignment, this is not generally true!

The name of the game: under what condition(s) does your regression/ estimator/research design provide a treatment effect of interest? Do those conditions hold in your case? When is your treatment effect **identified**?

• Also referred to as internal validity.

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Regression and causality

As another illustration, suppose there are constant treatment effects (δ) , so for every i, $Y_i(1) = Y_i(0) + \delta$. We don't observe potential outcomes, but rather the observed Y_i :

$$Y_i = D_i Y_i(1) + (1 - D_i) Y_i(0)$$

Plugging in for $Y_i(1)$, this can be written as a simple regression:

$$Y_i = \underbrace{E[Y(0)]}_{\beta_0} + \underbrace{\delta}_{\beta_1} D_i + \underbrace{Y_i(0) - E[Y(0)]}_{\text{residual}}$$

Note the residual for unit i is the deviation of $Y_i(0)$ from the population mean Y(0). With random assignment, D_i is uncorrelated with this residual. If there is *selection bias*—e.g., the treated tend to have higher potential outcomes than average—then there is omitted variables bias.

Now continue with constant treatment effects (δ) but suppose that potential outcomes depend linearly on X_i (and random noise u_i):

$$Y_i(0) = \alpha_0 + \alpha_1 X_i + u_i$$

$$Y_i(1) = \alpha_0 + \alpha_1 X_i + \delta + u_i$$

and that there is selection into treatment, such that D_i and X_i are related:

$$X_i = \gamma_0 + \gamma_1 D_i + w_i$$

(For example, the $D_i = 1$ group has higher X than average, $\gamma_1 > 0$).

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Regression and causality

The observed Y_i (using the switching equation) is:

$$Y_{i} = D_{i}Y_{i}(1) + (1 - D_{i})Y_{i}(0)$$

$$= D_{i}(\alpha_{0} + \alpha_{1}X_{i} + \delta + u_{i}) + (1 - D_{i})(\alpha_{0} + \alpha_{1}X_{i} + u_{i})$$

$$= \alpha_{0} + \delta D_{i} + \alpha_{1}X_{i} + u_{i}$$

If we estimate a naı̈ve simple regression of Y on D, $\alpha_1 X_i$ is in the residual:

$$Y_i = \beta_0 + \beta_1 D_i + \underbrace{v_i}_{\alpha_1 X_1 + u_i}$$

This is not a problem if X_i is uncorrelated with D_i , but in this case it is. There is omitted variables bias. X is related to potential outcomes and the D=1 and D=0 groups differ in their mean X.

If we plug in what we know about how X_i is related to D_i :

$$Y_i = \alpha_0 + \delta D_i + \alpha_1 (\gamma_0 + \gamma_1 D_i + w_i) + u_i$$

= $\alpha_0 + (\delta + \alpha_1 \gamma_1) D_i + \alpha_1 \gamma_0 + \alpha_1 w_i + u_i$

The slope coefficient is $\delta + \alpha_1 \gamma_1$. The latter term is the omitted variables bias.

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Regression and causality

Another way to see this. We know our estimator of β_1 in the simple regression will provide:

$$\begin{split} \beta_1 &= E[Y_i|D_i = 1] - E[Y_i|D_i = 0] \\ &= \alpha_0 + \alpha_1 E[X_i|D_i = 1] + \delta - \alpha_0 - \alpha_1 E[X_i|D_i = 0] \\ &= \delta + \underbrace{\alpha_1 (E[X_i|D_i = 1] - E[X_i|D_i = 0])}_{\text{selection bias}} \\ &= \delta + \underbrace{\alpha_1 (\gamma_0 + \gamma_1 - \gamma_0)}_{\text{selection bias}} \end{split}$$

selection bia

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Conditional independence assumption

This is a pretty simple case where estimating a regression that conditions on (controls for) X would eliminate the selection bias. Here, the only reason treated and untreated units differ in their potential outcomes (on average) is that they have different levels of X. This is a case of "selection" on observables"

$$Y_i = \beta_0 + \beta_1 D_i + \alpha_1 X_i + u_i$$

The conditional independence assumption holds here. Holding X constant, there is no association between treatment and potential outcomes.

Note also that we assumed constant treatment effects here.

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Example 2: private colleges

Does attending a selective private college result in higher (log) earnings?

	No		Selection counts				
	(1)	(2)	(3)	-	4)	(5)	н
Private school	.212 (.060)	.152 (467)	.139	.0		.031 j.062	.081
Own SAT score + 100		.051 (.008)	.024			.036 (.006)	.001
Log parental income			(.026)				.139 (325
Female			-,398 (J012				29 (8)
Black			003 (031)				-31
Hispanic			(.057)				.80t (.034
Asian			.189 (220.)				.155
Other/missing race			166 (118)				185 J.117
High school top 10%			.067 [.026]				(000)
High school rank missing			.003 (/025)				806 (-023
Addese			.107 (.027)				.092 (.024)
Average SAT score of schools applied to + 100				.110 (424)		162 (22)	.877 (012)
Sest two applications				,071 (,013)	(.0	62 11)	.058 (-010)
Sent three applications				.093 (021)	,0 (0	19)	.065 (.017)
Sust four or more applications				.339 (.024)	100		10501 10501

Example 2: private colleges

Does the regression in column (1) have a causal interpretation?

Attendance at a private college is not randomly assigned; we should be concerned that the coefficient on private school does not describe differences in average potential outcomes for any population of interest. It may be that students attending selective private colleges are better qualified on a number of dimensions than students not attending such colleges. The causal effect is not identified.

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Omitted variables bias

Suppose that potential outcomes (log earnings) are governed by:

$$Y_i(0) = \alpha + \gamma A_i + u_i$$

$$Y_i(1) = \alpha + \gamma A_i + \beta + u_i$$

 A_i is a measure of "ability" and u_i is a random error term. $P_i=1$ is an indicator for private college attendance ("treatment"). The switching equation gives us:

$$Y_i = \alpha + \beta P_i + \gamma A_i + u_i$$

Call this the "long" regression, which has a causal interpretation here. Relabel some coefficients:

$$Y_i = \alpha^{\ell} + \beta^{\ell} P_i + \gamma A_i + u_i^{\ell}$$

Omitted variables bias

Suppose instead we estimated the "short" regression (as in column (1) in Example 2):

$$Y_i = \alpha^s + \beta^s P_i + u_i^s$$

We know the model with the causal interpretation is the "long" regression $(\gamma \neq 0)$, so there may be *omitted variables bias* if A_i is related to P_i .

The error term in the short regression is: $u_i^s = \gamma A_i + u_i^{\ell}$.

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Omitted variables bias formula

There is a formal (and mechanical) link between β^s and β^ℓ :

$$\beta^s = \beta^\ell + \pi_1 \gamma$$

Where:

- γ comes from the long regression: it is the relationship between Y_i
 and A_i (conditional on P_i).
- π_1 comes from an "auxiliary" regression of the omitted variable (A_i) on the included variable (P_i) .

$$A_i = \pi_0 + \pi_1 P_i + v_i$$

Example 2

Table 2.5: auxiliary regressions where A_i is SAT score (in hundreds)

	No	selection	controls		Selectio	e cons	trok	Dairesses		TABLE 2.5 cts: Omir			
	(1)	(2)	(3)	Į.	D (5)	(6)	riivate sa	moor ene	cus: Omit			_
Private school	.212 (.060)	.152	.139	.0.			,837 (,839)		Own	SAT score -		nt variable Log	g parer
Own SAT score ÷ 100		.051	(,006)		,03 4,00		.009 (-006)		(1)	(2)	(3)	(4)	- (
Log parental income		(,000)	.181				.159	Private school	1.165 (.196)	(.188)	.066 (.112)	.128 (.035)	(.0
Female			398				-3%	Female		367 (.076)			.0
Black			003 (.031)				-,037 (,035)	Black		-1.947 (.079)			1.0
Hispanic			.027				,001 (054)	Hispanic		-1.185 (.168)			2 (.0
Asian			.189				.155	Asian		014 (.116)			0
Other/missing race			166 (:118)				189 (J.117)	Other/missing race		521 (.293)			0
High school top 10%			.067				.064	High school top 10%		.948 (.107)			0 (.0
High school rank missing			.003				006	High school rank missing		.556 (.102)			(.02
Adulese			.107				(992 (024)	Arblew		318 (.147)			.03
Average SAT score of schools applied to + 100				.110	(022)		077 012)	Average SAT score of schools applied to + 100			.777 (.058)		
Sent two applications				.071	.062		9101	Sent two applications			.252 (J077)		
Sent three applications				.093	(019)		966 1171	Sent these applications			.375 (.106)		
Seat four or more applications				.139	.127	.0		Seat four or more applications			.330		

Omitted variables bias: example

Assessing omitted variables bias in column (1) vs. (2):

- $\hat{\beta}^s = 0.212$
- $\beta^s = \beta^\ell + \pi_1 \gamma$
- What do you think the signs of π_1 and γ are?
- The estimated $\widehat{\pi_1}=1.165$ (the difference in SAT scores between private and public college students) and $\hat{\gamma}=0.051$
- So, 0.212 = β^{ℓ} + (1.165 * 0.051). Our estimator of β using β_s is likely biased upward.
- $\hat{\beta}^{\ell} = 0.152$ (compare to column (2))

Again, this is just mechanically true for any regression with an X_1 and an X_2 . We give it substance based on our application.

Omitted variables bias

Table 10.3. The omitted variable bias formula helps us think about whether failing to control for a confounder results in an over- or under-estimate of the causal effect.

	Omitted Variable Positively Correlated with Treatment $\pi > 0$	Omitted Variable Negatively Correlated with Treatment $\pi < 0$
Omitted Variable Positively Correlated with Outcome $\gamma > 0$	Positive bias $\pi \cdot \gamma > 0$	Negative bias $\pi \cdot \gamma < 0$
Omitted Variable Negatively Correlated with Outcome $\gamma < 0$	Negative bias $\pi \cdot \gamma < 0$	Positive bias $\pi \cdot \gamma > 0$

Source: Bueno de Mesquita & Fowler (2021)

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Example 2

Of course, a regression with two explanatory variables is probably not sufficient in this example: it alone is unlikely to adequately describe differences in average potential outcomes for a population of interest.

Column (3) of Table 2.3 includes additional student covariates, such as log parental income, gender, race/ethnicity, athlete, and HS top 10%. The reduction in $\hat{\beta}$ suggests the estimator used in column (2) was still biased upward.

In a setting like this, one should still be concerned about *unobserved*, possibly *unobservable* omitted variables.

The "unobservables"



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Example 2

In a further attempt to address unobserved omitted variables, columns (4) - (6) in Table 2.3 represent what might be called a "self-revelation" model. They include as covariates the number and characteristics of schools to which students *applied*. This behavior might proxy for unobserved differences that are related to both private college attendance and earnings.

This is a creative way to try to "control" for unobserved differences through a powerful proxy.

Example 2

		1)	IBLE 2.3		
Private	school	effects:	Average	SAT	score controls

	No	election	controls		Selection controls			
	(1)	(2)	(3)	P	D	5) (6)		
Private school	.212 (.060)	.152 (.057)	.139	.0.		(13 (13) (13)		
Own SAT score ÷ 100		.051	(.006)		(.0	36 .00 (86) (86		
Log parental income			(.026)			.12		
Female			398 (.012			35 (.01		
Black			003 (.031)			-,03 (,03		
Hispanic			.027 (.052)			,000 (,054		
Asian			.189 (.035)			.155 (437		
Other/missing race			166 (.118)			18 [.13		
High school top 10%			.067 (.020)			.064		
High school rank missing			.003 (.025)			-,000		
Adulese			.107 (.027)			(,024)		
Average SAT score of schools applied to + 100				.110 (.024)	(.022)			
Seer two applications				.071 (.013)	.062 (.011)	.058 (010)		
Sent three applications				(120.)	(.019)	(.017)		
Sent four or more applications				.139 (.024)	.127 (.023)	,098 (J020)		

	Dependent variable											
	Own	SAT score -	+ 100	Log parental income								
	(1)	(2)	(3)	(4)	(5)	(6)						
Private school	1.165	1.130 (.188)	.066 (.112)	.128	(.037)	,628 (465)						
Female		367 (.076)			.016 (.013)							
Black		-1.947 (.079)			359 (J019)							
Hispanic		-1.185 (.168)			259 (.050)							
Asian		014 (.116)			060 (.031)							
Other/missing race		521 (.293)			082 (.061)							
High school top 10%		.948 (.107)			066 (.011)							
High school rank missing		.556 (.102)			030 (.023)							
Addess		318 (.147)			.037							
Average SAT score of schools applied to + 100			.777 (.058)			,063 (,014						
Sent two applications			.252 (J077)			.020 (.010						
Seen these applications			.375 (.106)			.042 (J013						
Sent four or more applications			.330 (.093)			.079						

Note: This table describes the relationship between private school attendance and possonal characteristics. Dependent variables are the respondent's SAT some (diried by 100) in columns (11-11) and log particul income in columns (4)-46; Each column shows the coolinging from a eggestion of the dippodent variable on a durings for amounting a private intention and coatroll-The sample uses it 4-128. Stondard corrors are opported in parentheses.

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Example 2

In columns (4) - (6) the estimated coefficient on private school shrinks and becomes statistically insignificant.

Interestingly, the correlation between own SAT score and private school enrollment is eliminated once application behavior has been controlled for (the self-revelation model). See column (3) of Table 2.5.

Randomized controlled trials

Given their ability to eliminate selection bias, **randomized controlled trials** (**RCTs**) are considered the "gold standard" design for estimating treatment effects

- Since the treated and untreated are random draws from the same population, they should be balanced (in expectation) on observables and unobservables.
- This implies they are balanced (in <u>expectation</u>) on potential outcomes and thus on individual treatment effects.
- It is important to collect baseline data so these assumptions can be tested. While we can never test for differences in unobserved variables, differences in observed variables can be indicative of a problem.

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Randomized controlled trials

RCTs come with their own pitfalls and challenges, however. For example:

- Imperfect compliance: subjects may not comply with their treatment assignment. Estimating the ATE is not possible, but could estimate an intent-to-treat (ITT) effect, which may itself be of interest. This represents the impact of being offered treatment.
- Spillovers of treatment effect onto untreated units (SUTVA violation). Randomization at the group level may help reduce the likelihood of spillovers.
- Attrition from the study, particularly if non-random. Important to compare attrition rates for treated and untreated, and covariates for those remaining in the sample. Bounds analyses can probe the potential impact of attrition.

Randomized controlled trials

- Randomization bias: when behavior changes as a result of researcher-induced randomization (e.g., "Hawthorne effects")
- Ethical questions: is it ethical to withhold a treatment that has a likelihood of success?
- External validity: can the causal effect estimated from an RCT be generalized to other populations, places, times?

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Randomized controlled trials

Treatment effects are often estimated from RCTs using regression, although (given balance) one could estimate them from a difference in means (e.g., *t*-test). Benefits of using regression here:

- Convenient way to obtain standard errors and test for statistical significance, get confidence intervals, etc.
- Can include controls to improve precision. (These controls should not be affected by the treatment).
- Can include strata dummy variables if randomization is within strata (e.g., clustered random assignment).
- Can estimate treatment effects for subgroups using separate regressions or interaction terms.

Randomized controlled trials

Some simple RCT examples:

- Class exercise 1.4 on private school vouchers.
- RAND Health Insurance Experiment and Oregon Health Insurance Experiment in Mastering Metrics chapter 1.
- Gennetian et al., (2022) "Unconditional Cash and Family Investments in Infants: Evidence from a Large-Scale Cash Transfer Experiment in the U.S."

See Github Lecture 1 for sample RCT studies and related references.

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Causal inference with observational data

We frequently do not have the luxury of random assignment to treatment in the settings we study. The rest of the course will be devoted to designs that—when certain assumptions hold—make a strong claim to causal inference.

The RCT is an excellent benchmark, however. One of the best things you can do when designing your study with observational data is ask: what would be the ideal experiment to test my hypothesis? Is there a quasi-experimental design that approximates this for a population of interest? How close can I get to the "experimental ideal"?