# Summary of the course

Tutorial 7

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### Goal for today's tutorial

- Discuss the full course
  - Lecture 1: Binary choice models, censoring, truncation, and selection models (3-9)
  - Lecture 2: IV (10-13)
  - Lecture 3: Panel data models (14-20)
  - Lecture 4: Potential outcomes model (21-24)
  - Lecture 5: LATE and power analysis (25-27)
  - Lecture 6: DiD (28-30)
  - Lecture 7: RDD and RKD (31-34)

### Lecture 1: Binary choice models

ullet  $Y_i$  can only take the values 0 or 1

$$Y_i = egin{cases} 1 & ext{with probability } p_i \ 0 & ext{with probability } (1-p_i) \end{cases}$$

• For a binary model, the cumulative distribution function (cdf) is

$$p_i = P(Y_i = 1|X_i) = F(X_i'eta)$$

• For a binary model, the probability density function (density) is

$$f(Y_i|X_i) = p_i^{Y_i} (1-p_i)^{1-Y_i}$$

• To find  $\beta$ , use maximum likelihood function

$$egin{aligned} L(eta) &= \sum_{i=1}^{N} \left[ Y_i \ln p_i + (1-Y_i) \ln (1-p_i) 
ight] \ &= \sum_{i=1}^{N} \left[ Y_i \ln F(X_i'eta) + (1-Y_i) \ln (1-F(X_i'eta)) 
ight] \end{aligned}$$

ullet Notice we did not specify a particular form of the cdf  $F(X_i'eta)$ 

### Lecture 1: Binary choice models

- Linear probability model
  - $\circ \ p_i = F(X_i'eta) = X_i'eta$
  - $\circ$  Marginal effect:  $rac{\partial p_i}{\partial X_{ik}}=eta_k$
  - There is heteroskedasticity, so use robust s.e.
  - Estimated probabilities can be outside of the bounds
- Logit
  - $\circ~p_i = F(X_i'eta) = rac{exp(X_i'eta)}{1 + exp(X_i'eta)}$  the cdf of logistic distribution
  - $\circ$  Marginal effect:  $rac{\partial p_i}{\partial X_{ik}} = rac{exp(X_ieta)}{(1+exp(X_ieta)^2}eta_k$
  - $\circ$  MLE is not consistent if  $F(\cdot)$  is incorrectly specified
- Probit
  - $\circ~p_i = F(X_i'eta)$  =  $\Phi(X_i'eta)$  the cdf of standard normal distribution
  - $\circ$  Marginal effect:  $rac{\partial p_i}{\partial X_{ik}} = \phi(X_ieta)eta_k$
  - $\circ$  MLE is not consistent if  $F(\cdot)$  is incorrectly specified

#### Lecture 1: Latent structure

• Binary choice models are often written in terms of a latent structure with some latent (unobserved) variable

$$Y_i^* = X_i' eta + U_i$$

The observed outcome variable is

$$Y_i = egin{cases} 1 & ext{if} \ Y_i^* > 0 \ 0 & ext{if} \ Y_i^* \leq 0 \end{cases}$$

with

$$egin{aligned} P(Y_i = 1 | X_i) &= P(Y_i^* > 0 | X_i) \ &= P(X_i' eta + U_i > 0 | X_i) \ &= P(-U_i < X_i' eta | X_i) \ &= F(X_i' eta) \end{aligned}$$

where the cdf  $F(\cdot)$  is symmetric

## Lecture 1: Censoring and truncation

• The latent (unobserved) variable is

$$Y_i^* = X_i' eta + U_i$$

• The observe outcome variable is

$$Y_i = egin{cases} Y_i^* & ext{if} \ Y_i^* > c_i \ c_i & ext{if} \ Y_i^* \leq c_i \end{cases}$$

- Censored observations are in the sample
  - $\circ$  for them  $Y_i = c_i$  if  $Y_i^* \leq c_i$
- Truncated observations are not in the sample
  - $\circ$  for them  $Y_i$  is missing if  $Y_i^* \leq c_i$
- Ignoring censoring and truncation leads to a biased and inconsistent estimator

### Lecture 1: Censoring and truncation

• To find heta, use maximum likelihood function. Assume  $f^*(Y_i|X_i)$  is a density function of  $Y_i^*$ , then the cdf function of  $Y_i^*$  is

$$F^*(c_i|X_i) = P(Y_i^* < c_i|X_i) = \int_{-\infty}^{c_i} f^*(Y_i|X_i) dY_i$$

- Censoring
  - $\circ$  density function:  $f(Y_i|X_i)=f^*(Y_i|X_i)^{d_i}F^*(c_i|X_i)^{1-d_i}$  with  $d_i=1$  for uncensored observations
  - log-likelihood function

$$L( heta) = \sum_{i=1}^{N} \left[ d_i \ln f^*(Y_i|X_i, heta) + (1-d_i) \ln F^*(c_i|X_i, heta) 
ight]$$

- Truncation
  - $\circ$  density function:  $f(Y_i|X_i)=rac{f^*(Y_i|X_i)}{P(Y_i^*>c_i)}=rac{f^*(Y_i|X_i)}{1-F^*(c_i|X_i)}$
  - log-likelihood function

$$L( heta) = \sum_{i=1}^{N} \left[ \ln f^*(Y_i|X_i, heta) - \ln(1 - F^*(c_i|X_i, heta)) 
ight]$$

### Lecture 1: Sample selection model

- The outcome variable is observed only for a selected sample
- The sample selection model has two stages
  - Selection equation

$$I_i^* = Z_i' \gamma + V_i$$

ullet The indicator function, based on  $I_i^*$  , takes two values

$$I_i = egin{cases} 1 & ext{ if } I_i^* > 0 \ 0 & ext{ if } I_i^* \leq 0 \end{cases}$$

Regression equation

$$Y_i^* = X_i' eta + U_i$$

 $\circ$  However, we observe only  $Y_i$ 

$$Y_i = egin{cases} Y_i^* & ext{if } I_i = 1 \ ext{missing} & ext{if } I_i = 0 \end{cases}$$

### Lecture 1: Sample selection model

• To estimate the sample selection model, we make an assumption that disturbances terms are bivariate normal

$$\left[egin{array}{c} U_i \ V_i \end{array}
ight] \sim \mathcal{N}\left(0, \left[egin{array}{ccc} \sigma^2 & 
ho\sigma \ 
ho\sigma & 1 \end{array}
ight]
ight)$$

ullet Let us find expected value  $Y_i$  conditional on  $I_i=1$ , i.e. observed  $Y_i$ 

$$egin{aligned} E[Y_i|I_i = 1, Z_i, X_i] &= E[X_i'eta + U_i|I_i = 1, Z_i, X_i] \ &= X_i'eta + E[U_i|I_i = 1, Z_i, X_i] \ &= X_i'eta + E[U_i|Z_i'\gamma + V_i > 0, Z_i, X_i] \ &= X_i'eta + E[U_i| - V_i < Z_i'\gamma, Z_i, X_i] \ &= X_i'eta + 
ho\sigmarac{\phi(Z_i'\gamma)}{\Phi(Z_i'\gamma)} \end{aligned}$$

- ullet If ho=0, i.e. if  $U_i$  and  $V_i$  are independent or when  $X_i$  and  $Z_i$  are uncorrelated, OLS estimator is consistent
- If ho 
  eq 0, OLS estimator is inconsistent, and  $rac{\phi(Z_i'\gamma)}{\Phi(Z_i'\gamma)}$  is the Inverse Mills ratio which denotes selection bias

- If  $E(U_i|X_i) 
  eq 0$ , there is endogeneity problem
- ullet In this case OLS provides a biased and inconsistent  $\hat{eta}$
- Sources of endogeneity
  - Omitted variables
  - Reverse causality
  - Measurement error
- A solution is to use an instrument that should be
  - $\circ$  Relevant:  $\operatorname{cov}(Z_i, X_i) 
    eq 0$
  - $\circ$  Valid (exogenous):  $\mathrm{cov}(Z_i,U_i)=0$
- Use two-stage least squares (IV) estimator
  - First stage

$$X_i = \gamma_0 + \gamma_1 Z_i + V_i \ \Longrightarrow \hat{X}_i = \hat{\gamma_0} + \hat{\gamma_1} Z_i$$

Second stage

$$Y_i = \beta_0 + \beta_1 \hat{X}_i + U_i^*$$

$$\implies \hat{\beta}_{1,2SLS}$$

•  $\hat{eta}_{1,2SLS}$  has the following form

$$\hat{eta}_{1, ext{2SLS}} = rac{\sum_{i=1}^{n} \left(Z_i - ar{Z}_n
ight) \left(Y_i - ar{Y}_n
ight)}{\sum_{j=1}^{n} \left(Z_j - ar{Z}_n
ight) \left(X_j - ar{X}_n
ight)}$$

•  $\hat{eta}_{1,2SLS}$  is consistent

$$ext{plim}_{n o\infty}\hat{eta}_{1,2SLS} = rac{ ext{cov}(Z_i,Y_i)}{ ext{cov}(Z_i,X_i)} = eta_1 + rac{ ext{cov}(Z_i,U_i)}{ ext{cov}(Z_i,X_i)} = eta_1$$

•  $\hat{eta}_{1.2SLS}$  is biased

$$E[\hat{eta}_{1,2SLS}] = eta_1 + \sum_{i=1}^n \mathrm{E}\left[rac{rac{1}{n}\left(Z_i - ar{Z}_n
ight)U_i}{rac{1}{n}\sum_{j=1}^n\left(Z_j - ar{Z}_n
ight)\left(X_j - ar{X}_n
ight)}
ight] 
eq eta_1$$

 Do you want to derive more consistency and unbiasedness of estimators? Take the core course Advanced Econometrics I

- To test exogeneity of  $X_i$ , use the Hausman test
  - $\circ H_0$ :  $X_i$  is exogenous, i.e. OLS and 2SLS are both consistent

$$\circ$$
 Test statistic:  $H=rac{(\hat{eta}_{1,2SLS}-\hat{eta}_{1,OLS})^2}{\mathrm{var}(\hat{eta}_{1,2SLS}-\hat{eta}_{1,OLS})}\sim \chi^2(1)$ 

- $\circ$  Reject if  $H>\chi^2_lpha(1)$
- To test validity, use the Sargan test (over-identification required)
  - $\circ$   $H_0$ : all instruments are valid
  - Find the second-stage residuals and regress them on the instruments

$$U_i = \delta_0 + \delta_1 Z_{1,i}, + \ldots + \delta_M Z_{M,i} + e_i \sim \chi^2(M-1)$$

- $\circ$  Test statistic:  $H=nR^2$
- $\circ$  Reject if  $H>\chi^2_lpha(M-1)$
- ullet IV is consistent if instrument is relevant (F-test >10), but bias can be large

$$ext{Bias IV} \sim rac{\left\{\# ext{ instruments} 
ight\} imes 
ho(U_i, V_i) imes \left(1 - R_{ ext{partial}}^2
ight)}{\left\{\# ext{ observations} 
ight\} imes R_{ ext{partial}}^2}$$

where  $R^2_{
m partial}$  is the contribution of the instruments to  $R^2$  in the first-stage

ullet IV is weak if  $\mathrm{cov}(Z_i,X_i)$  is small. Recall

$$ext{plim}_{n o\infty}\hat{eta}_{1,2SLS} = rac{ ext{cov}(Z_i,Y_i)}{ ext{cov}(Z_i,X_i)}$$

- When  $\mathrm{cov}(Z_i,X_i)$  is close to 0, i.e. instrument is irrelevant, then the sampling variation in  $\mathrm{cov}(Z_i,X_i)$  is not helpful to estimate  $\beta_{1,2SLS}$
- Weak instruments can be detected in the first-stage using a t-test or a F-test
  - $\circ$  Rule of thumb: instrument is weak if bias IV is larger than 10% of the bias of OLS

$$rac{ ext{Bias IV}}{ ext{Bias OLS}} pprox rac{ \{\# ext{ instruments } \} }{ \{\# ext{ observations } \} imes R_{ ext{partial}}^2 }$$

 Do you want to study more about weak IV? Take the field course Advanced Microeconometrics

ullet Assume N individuals observed over T periods

$$Y_{it} = lpha + X'_{it}eta + \eta_i + U_{it}$$

- $\circ$   $\eta_i$  is an individual specific effect which captures unobserved heterogeneity
- How to estimate this model?
  - Pooled OLS
  - Fixed-effects model
  - Random-effects model
- Assumptions for all three models
  - $\circ$  Strict exogeneity:  $E[U_{it}|X_{i1},\ldots,X_{iT},\eta_i]=0$  allows for only static panel models
  - $\circ$  Weak exogeneity:  $E[U_{it}|X_{it},\eta_i]=0$  allows models to be dynamic
- Do you want to study dynamic panel models? Take the field course Applied Microeconometrics

Pooled OLS

$$Y_{it} = \alpha + X'_{it}\beta + U^*_{it}$$

where  $U_{it}^* = \eta_i + U_{it}$ 

- If  $E[\eta_i|X_{i1},\ldots,X_{iT}] 
  eq 0$ , i.e. individual specific effects are correlated with regressors, the OLS estimator of eta is biased and inconsistent
- If  $E[\eta_i|X_{i1},\ldots,X_{iT}]=0$ , the OLS estimator of  $\beta$  is unbiased and consistent, but we still have that  $E[U_{it}^*U_{is}^*] \neq 0 \implies$  use clustered s.e.

Fixed-effects model

$$Y_{it} = \alpha + X'_{it}\beta + \eta_i + U_{it}$$

where  $\eta_i$  is fixed

- Within estimation
  - Estimation

$$egin{aligned} Y_{it} - ar{Y}_i &= lpha + X_{it}'eta + \eta_i + U_{it} - (lpha + ar{X_i}'eta + \eta_i + ar{U_i}) \ &= (X_{it} - ar{X}_i)'eta + (U_{it} - ar{U_i}) \end{aligned}$$

- $\circ$  Assumption:  $E[(X_{it}-ar{X}_i)'(U_{it}-ar{U}_i)]=0$
- First-difference
  - Estimation

$$Y_{it} - Y_{it-1} = lpha + X_{it}'eta + \eta_i + U_{it} - (lpha + X_{it-1}'eta + \eta_i + U_{it-1}) \ = (X_{it} - X_{it-1})'eta + (U_{it} - U_{it-1})$$

- $\circ$  Assumption:  $E[(X_{it}-X_{it-1})'(U_{it}-U_{it-1})]=0$
- Do you want to combine IV and FE? Take the core course Advanced Econometrics II

- If strict exogeneity is violated, both FE estimators are not consistent
- To test strict exogeneity, use the following specifications
  - $\circ$  For T=2, both estimators are the same so check

$$Y_{it} - Y_{it-1} = (X_{it} - X_{it-1})' eta + X_{it}' \gamma + (U_{it} - U_{it-1})$$

- $\circ \ H_0: \gamma = 0$ , use a t-test or F-test to check that
- $\circ$  For T>2, the estimators should be close

$$Y_{it} = \alpha + X'_{it}\beta + X'_{it+1}\gamma + \eta_i + U_{it}$$

 $\circ \ H_0: \gamma = 0$ , use a t-test or F-test to check that

• To check serial correlation, estimate the first difference regression

$$egin{aligned} Y_{it} - Y_{it-1} &= (X_{it} - X_{it-1})' eta + (U_{it} - U_{it-1}) \ &= (X_{it} - X_{it-1})' eta + E_{it} \ &\Longrightarrow \hat{E}_{it} = (Y_{it} - Y_{it-1}) - (X_{it} - X_{it-1})' \hat{eta} \end{aligned}$$

• Estimate the following model

$$\hat{E}_{it} = 
ho \hat{E}_{it-1} + e_{it}$$

ullet  $H_0:
ho=-0.5$ , use a t-test to check that. If there is no autocorrelation

$$egin{aligned} \hat{
ho} &= rac{ ext{cov}(E_{it-1}, E_{it})}{ ext{cov}(E_{it-1}^2)} = rac{ ext{cov}(U_{it-1} - U_{it-2}, U_{it} - U_{it-1})}{ ext{cov}(U_{it} - U_{it-1}, U_{it} - U_{it-1})} \ &= rac{ ext{cov}(U_{it-1}, U_{it}) - ext{cov}(U_{it-1}, U_{it-1}) - ext{cov}(U_{it-2}, U_{it}) + ext{cov}(U_{it-2}, U_{it-1})}{ ext{cov}(U_{it}, U_{it}) - ext{cov}(U_{it}, U_{it-1}) - ext{cov}(U_{it-1}, U_{it}) + ext{cov}(U_{it-1}, U_{it-1})} \ &= rac{- ext{cov}(U_{it-1}, U_{it-1})}{ ext{cov}(U_{it}, U_{it}) + ext{cov}(U_{it-1}, U_{it-1})} = rac{-\sigma_u^2}{2\sigma_u^2} = -rac{1}{2} \end{aligned}$$

If there is autocorrelation, use robust s.e.

Random-effects model

$$Y_{it} = lpha + X_{it}'eta + \eta_i + U_{it}$$

where  $\eta_i$  is random and  $E[\eta_i|X_{i1},\ldots,X_{iT}]=0$ 

- Estimation
  - $\circ$  Stack observations of all individuals  $Y_i = X_i' eta + e_T \eta_i + U_i$
- ullet If  $\sigma_\eta^2$  and  $\sigma_u^2$  known, use the GLS estimator

$$\hat{eta}_{GLS} = \sum_{i=1}^{N} (X_i' \Omega^{-1} X_i)^{-1} \sum_{i=1}^{N} (X_i' \Omega^{-1} Y_i)$$

where 
$$ext{var}(e_T\eta_i + U_i) = \sigma_u^2(I_T + rac{\sigma_\eta^2}{\sigma_u^2}e_Te_T') = \sigma_u^2\Omega$$

- ullet If  $\sigma_{\eta}^2$  and  $\sigma_{u}^2$  unknown, use the FGLS estimator
  - $\circ$  Estimate  $\sigma_u^2$  by within estimation
  - $\circ$  Estimate  $\sigma_{\eta}^2$  by between estimation
  - $\circ$  Do GLS with  $\hat{\Omega}$  instead of  $\Omega$
- ullet In general  $\sigma_{\eta}^2$  and  $\sigma_{u}^2$  are unknown, so one has to apply FGLS

- FE or RE model
  - RE can deal with time-invariant regressors
  - RE can be used to make predictions outside the sample
  - $\circ$  RE has a stronger assumption that  $E[\eta_i|X_{i1},\ldots,X_{iT}]=0$
  - FE robust against correlation of individual effects and regressors
  - FE is less efficient and parameter estimates might be noisy
- Use the Mundlak procedure, to test which model to use
  - Estimate the RE model

$$Y_{it} = X_{it}'eta + ar{X}_i'\gamma + \omega_i + U_{it}$$

where  $\eta_i = {ar X}_i' \gamma + \omega_i$  and  $\omega_i$  is a random effect that is uncorrelated with  $X_{it}$ 

- $\circ~H_0: \gamma = 0$ , i.e. the random effects model should be used
- Alternative use the Hausman test

$$H=(\hat{eta}_{FE}-\hat{eta}_{RE})'[ ext{var}(\hat{eta}_{FE})- ext{var}(\hat{eta}_{RE})]^{-1}(\hat{eta}_{FE}-\hat{eta}_{RE})\sim \chi^2(R)$$

where R is the number of time-varying regressors

 $\circ \ H_0: E[\eta_i|X_{i1},\ldots,X_{iT}]=0$ , i.e. RE and FE are consistent, but RE is more efficient

- The goal of policy evaluation is to obtain a causal effect of treatment on the outcome of interest
- Potential outcomes model
  - $\circ$  Each individual has 2 potential outcomes:  $Y_{1i}^st$  if treated and  $Y_{0i}^st$  if untreated
  - $\circ \Delta_i = Y_{1i}^* Y_{0i}^*$  individual effect of participating in treatment (not observed)
  - This is the fundamental problem of causal inference
- Treatment effects

$$ATE = E[\Delta] = E[Y_1^* - Y_0^*] = E[Y_1^*] - E[Y_0^*]$$

ATE is the effect for the full population

$$ATET = E[\Delta|D=1] = E[Y_1^* - Y_0^*|D=1] = E[Y_1^*|D=1] - E[Y_0^*|D=1]$$

ATET is the effect for individuals who actually received the treatment

 If there is self-selection into treatment, participation might not be independent of the potential outcomes, i.e. people with positive individual effects are more likely to participate

$$E[Y_1^*] 
eq E[Y_1^*|D=1] 
onumber \ E[Y_0^*] 
eq E[Y_0^*|D=0] 
onumber$$

- In this case
  - $\circ~E[Y_1^*|D=1]$  and  $E[Y_0^*|D=0]$  can be estimated
  - $\circ\ E[Y_1^*|D=0]$  and  $E[Y_0^*|D=1]$  can't be estimated
- A solution is to use randomized experiments
  - $\circ$  Treatment is assigned randomly, i.e  $(Y_{0i}^*,Y_{1i}^*)\perp D_i$
  - So we can assume the same expected effect for treated and untreated

$$\begin{split} E[Y_1^*] &= E[Y_1^*|D=1] = E[Y_1^*|D=0] \\ E[Y_0^*] &= E[Y_0^*|D=0] = E[Y_0^*|D=1] \end{split}$$

which implies ATE = ATET

• To estimate the treatment effect, let the observed outcome be

$$Y_i = D_i Y_{1i}^* + (1 - D_i) Y_{0i}^*$$

where  $D_i=1$  if a person received treatment

• Estimate the sample means to get the estimators

$$E[\hat{Y}_{1}^{*}|D=1] = rac{\sum_{i=1}^{N}D_{i}Y_{i}}{\sum_{i=1}^{N}D_{i}} \ E[\hat{Y}_{0}^{*}|D=0] = rac{\sum_{i=1}^{N}(1-D_{i})Y_{i}}{\sum_{i=1}^{N}(1-D_{i})} \ A\hat{T}E = A\hat{T}ET = rac{\sum_{i=1}^{N}D_{i}Y_{i}}{\sum_{i=1}^{N}D_{i}} - rac{\sum_{i=1}^{N}(1-D_{i})Y_{i}}{\sum_{i=1}^{N}(1-D_{i})} \ .$$

• Potential outcomes model is equivalent to the difference-in-means estimator

$$Y_i = \alpha + \delta D_i + U_i$$

where 
$$\delta = \hat{ATE} = \hat{ATET}$$

- Validity of experiments
  - Internal validity (no spill-over effects, no substitution, no Hawthorne effect) extent to which we can make causal inference
  - External validity how experimental results generalize
  - Stable unit treatment value assumption (SUTVA) treatment participation of one individual does not affect the potential outcomes of other individuals
- Field experiments used because randomized experiments are rare usually implemented as randomization in natural environment
- Types of field experiments
  - Oversubscription if there are more applicants than available slots, assign treatment by lottery
  - Phasing-in start low scale, expand later
  - Within-group only some subgroups get treatment, others not
  - Encouragement design randomly encourage subsample to participate
- DiNardo and Lee (2011) judge every method for evaluation on three criteria
  - Appropriate description of treatment assignment mechanism
  - Consistent with wide class of behavioral models
  - Yields testable implications

### Lecture 5: LATE and power analysis

- ullet If there is partial compliance in an experiment, initial treatment assignment is often not equal to actual treatment assignment, i.e.  $Z_i 
  eq D_i$
- If people self-select into treatment, treatment effect is heterogeneous

$$Y_i = lpha + \delta_i D_i + U_i$$

where  $\delta_i$  is individual (heterogeneous) effect

- In this case Imbens and Angrist (1994) suggest to study LATE imposing monotonicity assumption
  - If you get the treatment, you don't want to opt out
  - If you don't get the treatment, you want to get one

$$D_i(1) \geq D_i(0)$$

• When the instrument is binary, LATE is equal to

$$LATE = rac{E[Y|Z=1] - E[Y|Z=0]}{Pr[D=1|Z=1] - Pr[D=1|Z=0]}$$

### Lecture 5: LATE and power analysis

- If monotonicity holds, LATE is the average treatment effect for compliers
  - Without monotonicity difficult interpretation
- Randomization implies the same share of compliers, always takers, and never takers in treated and control groups
  - $\circ$  Compliers: D(1)=1 and D(0)=0
  - $\circ$  Always takers: D(1)=D(0)=1
  - $\circ$  Never takers: D(1)=D(0)=0
  - $\circ$  Defiers: D(1)=0 and D(0)=1 ruled out by monotonicity

### Lecture 5: LATE and power analysis

• What is the minimum effect we are able to detect, given the treatment intensity, the number of participants, and the power?

$$MDE = (t_{1-lpha/2} - t_{1-q}) \sqrt{rac{1}{p(1-p)} rac{\sigma^2}{n}} rac{1}{r_t - r_c}$$

• What is smallest sample size needed to run an experiment, given MDE, the treatment intensity, and the power?

$$n=\left(rac{t_{1-lpha/2}-t_{1-q}}{MDE}
ight)^2rac{\sigma^2}{p(1-p)}\left(rac{1}{r_t-r_c}
ight)^2.$$

where MDE - minimum detectable effect; n - sample size; p - treatment intensity;  $\sigma^2$  - variance;  $r_t$  - compliance rate in the treatment group;  $r_c$  - treatment intensity in the control group

- MDE can be based on
  - earlier literature
  - requirements from partner
  - cost-benefit analysis

#### Lecture 6: DiD

• To compare differences between treatment and control groups before and after the intervention, estimate the following model

$$Y_{gt} = lpha_t + \eta_g + \delta D_{gt} + U_{gt}$$

where  $lpha_t$  - common time trend,  $\eta_g$  - group specific effect

ullet If we have 2 groups and 2 time periods, we can rewrite that as

$$egin{aligned} Y_{T0} &= lpha_0 + \eta_T + U_{T0} \ Y_{T1} &= lpha_1 + \delta + \eta_T + U_{T1} \ Y_{C0} &= lpha_0 + \eta_C + U_{C0} \ Y_{C1} &= lpha_1 + \eta_C + U_{C1} \end{aligned}$$

Take the differences of expected values

$$egin{aligned} E[Y_{T1}] - E[Y_{T0}] &= (lpha_1 + \delta + \eta_T) - (lpha_0 + \eta_T) = lpha_1 + \delta - lpha_0 \ E[Y_{C1}] - E[Y_{C0}] &= (lpha_1 + \eta_C) - (lpha_0 + \eta_C) = lpha_1 - lpha_0 \ ext{DiD} &= (lpha_1 + \delta - lpha_0) - (lpha_1 - lpha_0) = \delta \end{aligned}$$

ullet DiD estimates ATET if there is a constant treatment effect  $\delta$  or there are only 2 periods

#### Lecture 6: DiD

- Key assumption: parallel trend assumption
- Parallel trend assumption is scale dependent
  - If prior trends are the same in the logarithm of wages, they are not equal in wage
     levels
- ullet Intervention should be random conditional on time and group specific effects, otherwise  $E[U_{q0}|D_q] 
  eq 0$  which violates exogeneity
  - Example: Ashenfelter dip treatment participants have a dip in outcomes just before entering the programme
- How to test the parallel trend assumption?
  - Check prior trends
  - Do placebo checks
- If there is no support for parallel trend assumption
  - Include time-varying covariates or group specific trends
  - o DDD
  - Synthetic control group
  - DiD with IV
  - Changes-in-Changes

#### Lecture 6: DiD

• If there are more than two periods, we can implement an event-study specification

$$Y_{gt} = lpha_t + \eta_g + \delta_{t- au_g} D_{gt} + U_{gt}$$

#### where

- $\circ$   $au_q$  is the moment of the treatment
- $\circ$  t is time period
- Impose the normalisation  $\delta_{-1}=0$ , which is the coefficient for the last period before the treatment, otherwise you get perfect multicollinearity
- If data sampling process or treatment is clustered, use clustered s.e.
  - Abadie et al. 2017 discuss clustering
- Do you want to discuss clustering s.e. more? Take the core course Advanced Econometrics II

- Sharp RDD
  - $\circ$  Treatment assignment is sharp at the cutoff point  $ar{S}$

$$D_i = I(S_i > ar{S})$$

• Use crossing the cutoff, to estimate the marginal treatment effect

$$MTE(ar{S}) = \mathrm{lim}_{s\downarrow ar{S}} E[Y_i|S_i = s] - \mathrm{lim}_{s\uparrow ar{S}} E[Y_i|S_i = s]$$

• The model for sharp RDD to estimate the effect of a treatment on the outcome

$$Y_i = lpha + \delta D_i + K(S_i - ar{S}) + U_i$$

- RDD and RKD exploit local randomization
  - $\circ$  Interpretation: ATE for people who change status when moving from just below to just above  $ar{S}$  LATE

- Fuzzy RDD
  - $\circ$  Treatment assignment is discontinuous at the cutoff point  $ar{S}$

$$\lim_{s\downarrow ar{S}} P(D_i=1|S_i=s) 
eq \lim_{s\uparrow ar{S}} P(D_i=1|S_i=s)$$

 Use crossing the cutoff as a locally valid instrument, to estimate the marginal treatment effect

$$MTE(ar{S}) = rac{\lim_{s\downarrow ar{S}} E[Y_i|S_i=s] - \lim_{s\uparrow ar{S}} E[Y_i|S_i=s]}{\lim_{s\downarrow ar{S}} P[D_i=1|S_i=s] - \lim_{s\uparrow ar{S}} P[D_i=1|S_i=s]}$$

- The model for fuzzy RDD uses two stages as in IV
  - $\circ$  First-stage: estimate the effect of crossing  $ar{S}$  on the probability to get the treatment

$$egin{aligned} D_i &= \gamma_0 + \gamma_1 I(S_i > ar{S}) + G(S_i - ar{S}) + V_i \ &\Longrightarrow \; \hat{D}_i = \hat{\gamma}_0 + \hat{\gamma}_1 I(S_i > ar{S}) + \hat{G}(S_i - ar{S}) \end{aligned}$$

 $\circ$  Second-stage: use  $\hat{D}_i$  to estimate the effect of the treatment on the outcome

$$Y_i = lpha + \delta \hat{D}_i + K(S_i - ar{S}) + U_i$$

- RKD is similar to RDD, but instead of a jump in the intercept, the slope changes at the threshold
- The model for RKD uses "two stages"
  - $\circ$  First-stage: estimate the effect of crossing  $ar{S}$  on the probability to get the treatment

$$D_i=\gamma_0+\gamma_1(S_i-ar{S})I(S_i$$

 $\circ$  Second-stage: estimate the effect of crossing  $ar{S}$  on the changes in the slope

$$Y_i = eta_0 + \delta_1(S_i - ar{S})I(S_i < ar{S}) + \delta_2(S_i - ar{S})I(S_i \geq ar{S}) + U_i$$

 $\circ$  Estimate the causal effect of  $D_i$  on  $Y_i$  at  $S_i = ar{S}$ 

$$\frac{\delta_2-\delta_1}{\gamma_2-\gamma_1}$$

There is a treatment effect if

$$\lim_{S_i\uparrow ar{S}}rac{\partial E[Y_i|S_i]}{\partial S_i}=\delta_1
eq \delta_2=\lim_{S_i\downarrow ar{S}}rac{\partial E[Y_i|S_i]}{\partial S_i}$$

- Good practices
  - $\circ$  Use the McCrary test to check for continuity of density of  $S_i$  around the cutoff  $ar{S}$
  - Choose different bandwidths to check sensitivity
  - Choose different functional forms but don't use higher-order polynomials
  - Check if other characteristics are balanced around the discontinuity
  - Use controls as placebo tests
  - Try to use local-linear regression instead of polynomials
- Threats to validity
  - Treatment assignment rule may be public knowledge, which may trigger behavioral responses
  - Possible manipulation of the treatment variable

#### Courses

- Advanced Econometrics I
- Advanced Econometrics II
- Advanced Microeconometrics
- Applied Microeconometrics

# Final thoughts

• Good luck:)