Microeconometrics (Causal Inference) Week 2 - Linear regression (OLS) and bootstrapping

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What are we doing today?

- ► This week we will review linear regression (OLS) and its assumptions
 - ▶ We will also discuss how to interpret the results of a regression

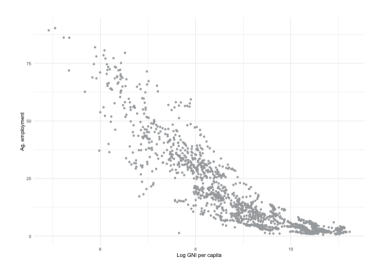
- ▶ Some of this will be review, but there will be some new stuff, too
 - ► For example, we will discuss bootstrapping on Thursday

What are we doing today?

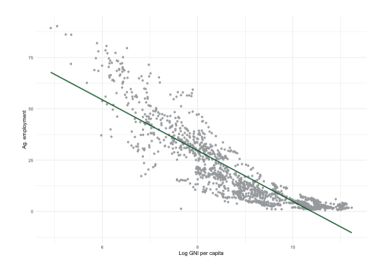
► I will be going through some econometric theory this week

- But we will also be using some data throughout the week
 - ► This data comes from the World Development Indicators database
 - ▶ I have uploaded the csv file to this week's GitHub folder

Fitting lines to data



Which line best fits the data?



Ordinary Least Squares (OLS)

- OLS is a method for fitting a line to data
 - It is the most common method for fitting lines to data
- \triangleright You have an outcome, let's call it y_i (where y is the outcome and i is an individual)
- You also have a set of covariates, let's call them x_{ik} (where k denotes different covariates)
- The relationship might look something like this, assuming linearity:

$$y_i = \beta_0 + \beta_1 x_{i,1} + \beta_2 x_{i,2} + \dots + \beta_k x_{i,k} + \epsilon_i$$
 (1)

Ordinary Least Squares (OLS)

- lacktriangle OLS is about finding the **average** relationship between Y and a set of covariates, X
 - ▶ The fit will never be perfect, which is why we have ϵ_i
 - $ightharpoonup \epsilon_i$ is the error term and shows that individuals will deviate from the average relationship
 - For example, Bill Gates is a high-school drop out but is still a billionaire; this is deviation from an average relationship between education and income

Minimizing the sum of squared errors

- ▶ In order to find a "line of best fit", we need an objective function
 - ▶ We want to minimize the distance between the line and the data, but how do we define that distance?
- ► We can define the distance as the sum of squared errors (SSE)
 - ► The SSE is the sum of the squared difference between the actual value and the predicted value
 - ► The predicted value is the value on the line of best fit
- In other words:

$$SSE = \sum_{i=1}^{n} (y_i - \hat{y}_i)^2$$
 (2)

Minimizing the sum of squared errors

$$SSE = \sum_{i=1}^{n} (y_i - \hat{y}_i)^2$$
 (3)

- \triangleright \hat{y}_i is the predicted value of y_i for individual i
- ▶ But how do we find \hat{y}_i ?
 - ► We need to find the line of best fit
 - \blacktriangleright We need to find the values of $\beta_0, \beta_1, \dots, \beta_k$ that minimize the SSE.
- We can do this by noting that $\hat{y}_i = \hat{\beta}_0 + \hat{\beta}_1 x_{i,1} + \hat{\beta}_2 x_{i,2} + \cdots + \hat{\beta}_k x_{i,k}$:

SSE =
$$\sum_{i=1}^{n} (y_i - \beta_0 - \hat{\beta}_1 x_{i,1} - \hat{\beta}_2 x_{i,2} - \dots - \hat{\beta}_k x_{i,k})^2$$
 (4)

In matrix form

- Let's rewrite the equation in matrix form:
- ightharpoonup Y is a vector of outcomes, X is a matrix of covariates, and β is a vector of coefficients:

$$SSE = (\mathbf{Y} - \mathbf{X}\beta)'(\mathbf{Y} - \mathbf{X}\beta)$$
 (5)

- ▶ It turns out that this becomes an optimization problem
- ▶ We know the **Y** and **X**, so we want to minimize this with respect to β :

$$\min_{\beta} (\mathbf{Y} - \mathbf{X}\beta)' (\mathbf{Y} - \mathbf{X}\beta) \tag{6}$$

The solution

► We know the solution to this:

$$\hat{\boldsymbol{\beta}} = (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'\mathbf{Y} \tag{7}$$

Seer this formula into your brain!

A problem: uncertainty

- We have a solution, but we have a problem: uncertainty
 - We generally do not have the population, which means $\hat{\beta}$ is just a sample estimate of the population parameters β
 - ▶ Note the use of hats to denote sample estimates

- ▶ We need to know how uncertain we are about our estimates
 - lacktriangle We need to know how much \hat{eta} will vary from sample to sample
 - ▶ Before we do this, we need to do a little bit of background on probability and statistics

Two important theorems

- ▶ There are two important theorems that we need to know about
 - ► The Law of Large Numbers
 - ► The Central Limit Theorem

► Let's go through these one at a time

Law of Large Numbers (LLN)

- ► The LLN says that as the sample size increases, the sample mean will converge to the population mean
 - ln other words, as $n \to \infty$, $\bar{x} \to \mu$
 - where n is the sample size, \bar{x} is the sample mean, μ is the true population mean

▶ You don't have to take my word for this, though. Let's see it in action.

LLN as an example

Let's create a "population" of 100,000 random numbers from a normal distribution with mean 0 and standard deviation 1

```
# note that rnorm is a "random" number generator, so we need to set a seed to make sure we get the same results each time
set.seed(1304697)
# NOTE: just set the seed once, at the top of your script. Then run everything and you will get reproducible results
population <- rnorm(100000, mean = 0, sd = 1)
mean(population)</pre>
```

[1] -0.003674513

LLN as an example

- ▶ So we know the true mean is $\mu = -0.0036745$
- Let's see what happens as we take a sample of size 10:

```
sample <- population[sample(1:length(population), 10, replace = FALSE)]
mean(sample)
## [1] -0.07407166</pre>
```

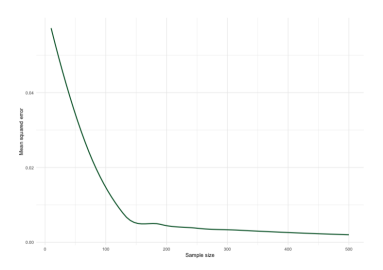
- Let's see what happens as we take a sample of size 100:

```
sample <- population[sample(1:length(population), 100, replace = FALSE)]
mean(sample)
## [1] -0.01106315</pre>
```

Suppose we did this a bunch of times...

- Let's say we took a sample of size 10, 100 times
 - ► We would get 100 different sample means
 - ▶ We can calculate how "far" each sample mean is from the true mean
 - ▶ We can do the same thing for a bunch of different sample sizes
- Let's then plot the average mean squared error (MSE) for each sample size
 - ▶ The MSE is the average squared difference between the sample mean and the true mean
 - ▶ In other words, it is the average of $(\bar{x} \mu)^2$

The results



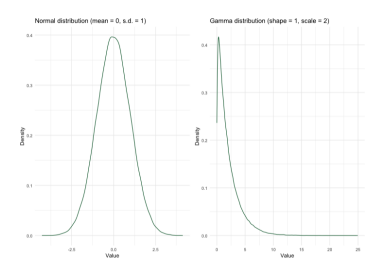
Non-normal distributions

- ► This doesn't just work with normal distributions
- Here's an example with a more skewed distribution
- Let's create a "population" of 100,000 random numbers from a gamma distribution with a shape (k) of 1 and a scale (θ) of 2

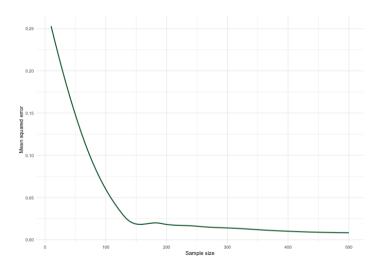
```
population2 <- rgamma(100000, shape = 1, scale = 2)
mean(population2)</pre>
```

[1] 2.005826

The two populations



The results



The takeaway from the LLN

- ► The main takeaway from the LLN is that as the sample size increases, the sample mean will converge to the population mean
 - ▶ In other words, as $n \to \infty$, $\bar{x} \to \mu$

- ► This holds for many different distributions!
 - lacktriangle Something we will return to is that this also holds for the distribution of $\hat{oldsymbol{eta}}$

The Central Limit Theorem

- ► The LLN is about the mean
 - ► The Central Limit Theorem (CLT) is about the **distribution**

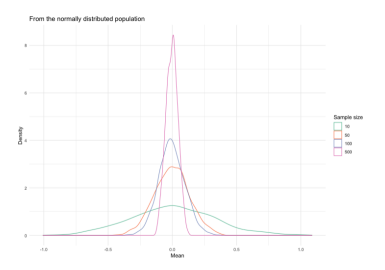
- ► The CLT says that as the sample size increases, the distribution of the sample mean will converge to a normal distribution
 - ln other words, as $n \to \infty$, $\bar{x} \sim N(\mu, \sqrt{\sigma^2/n})$
 - lacktriangle where μ is the true population mean and σ^2 is the true population variance

The CLT, empirically

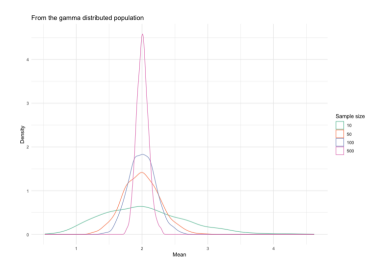
- We can do something similar to what we did with the LLN
 - Instead of looking at means, though we will look at distributions of the mean
 - In other words, we will density functions of the sample means
- ► I am going to take a sample and find the mean
 - ► Then I'm going to do it again
 - And again
 - ▶ 1,000 times

Then I will plot the density of the sample means, for four separate sample sizes (10, 50, 100, and 500)

The CLT with four sample sizes: 10, 50, 100, and 500 and 1,000 replications



The CLT with four sample sizes: 10, 50, 100, and 500 and 1,000 replications



The CLT lets us quantify uncertainty

- ► The key thing about the CLT is that the math behind us *lets us quantify the uncertainty!*
- For sample means, the CLT says that the standard error of the mean is:
 - $ightharpoonup \sigma_{\bar{x}} = \frac{\sigma}{\sqrt{n}}$
 - where σ is the population standard deviation and n is the sample size

▶ Of course, we don't know σ , but we can estimate it!

The CLT lets us quantify uncertainty

- \blacktriangleright We can estimate σ with s
 - s is the sample standard deviation
 - ► So we can estimate the standard error of the mean as:

$$ightharpoonup SE_{\bar{x}} = rac{s}{\sqrt{n}}$$

- ► This "standard error of the mean" is the standard deviation of the distribution of the sample mean
 - ► That distribution is called the sampling distribution of the mean

Uncertainty in OLS

Let's go back to our (population) regression equation:

$$\mathbf{Y} = \mathbf{X}\boldsymbol{\beta} + \boldsymbol{\epsilon} \tag{8}$$

- ► What drives uncertainty here?
 - ightharpoonup The error term. ϵ
 - Note that an error term is a *population* parameter
 - ► The sample analog is the **residual**

Uncertainty in OLS

We can see that the error term is responsible for uncertainty by thinking about the deviation of estimated regression coefficients $(\hat{\beta})$ from the true regression coefficients (β)

$$\mathbf{Y} = \beta \mathbf{X} + \epsilon \qquad \hat{\beta} = (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'\mathbf{Y}$$
$$\hat{\beta} = (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'(\beta \mathbf{X} + \epsilon)$$
$$\hat{\beta} = \beta + (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'\epsilon \qquad (9)$$

- ightharpoonup eta is fixed, but ϵ is random
 - ightharpoonup So \hat{eta} is random, too
 - ightharpoonup That randomness is driven by the error term, ϵ

Variance of $\hat{\beta}$

- ▶ Variance is about how much a random variable varies from its mean
 - ► It's a second-order moment: it's about how much a random variable varies from its mean (true value!)
 - ▶ Note how we use expectations here, since we are talking about population parameters

$$Var(\hat{\boldsymbol{\beta}}|\boldsymbol{X}) = (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'\mathbb{E}(\epsilon'\epsilon|\boldsymbol{X})\mathbf{X}(\mathbf{X}'\mathbf{X})^{-1}$$
(10)

- ► Well that doesn't look very good, does it?
 - lacktriangle We can simplify this. . . with an assumption on $\mathbb{E}(\epsilon'\epsilon|\pmb{X})$, which we can denote Ω

Homoskedasticity

▶ If we are willing to assume structure on the error term, we can simplify the variance

Assumption: Homoskedasticity

 $\Omega = \sigma^2 \mathbf{I}$, where \mathbf{I} is the identity matrix (ones on the diagonal and zeros elsewhere).

This simplifies the variance to:

$$Var(\hat{\boldsymbol{\beta}}_{homoskedasticity}) = \sigma^2 (\mathbf{X}'\mathbf{X})^{-1}$$
 (11)

Homoskedasticity

- ► What, exactly, is homoskedasticity?
- ▶ It means that the variance of the error term is constant
 - ▶ In other words, the variance of the error term does not depend on the covariates
 - ▶ This is a very strong assumption and is often violated in practice

Estimating the variance under homoskedasticity

- ▶ Even though it may not be reasonable, let's assume homoskedasticity
 - ▶ Then, the variance of $\hat{\beta}_{homoskedasticity}$ is:

$$\sigma^2(\mathbf{X}'\mathbf{X})^{-1} \tag{12}$$

- ▶ There's a problem, though: we don't know σ^2
 - We can estimate it, just like with the mean
 - ► We can estimate it with the residual sum of squares (RSS)
 - ► The RSS is the sum of the squared *residuals*

Estimating the variance under homoskedasticity

- We can estimate σ^2 as $\frac{RSS}{n-k-1}$, where n is the sample size and k is the number of covariates
 - ► The extra 1 is due to the intercept
- ► We estimate the variance as:

$$Var(\hat{\boldsymbol{\beta}}_{homoskedasticity}) = \hat{\sigma}^2 (\mathbf{X}'\mathbf{X})^{-1}$$
(13)

$$=\frac{RSS}{n-k-1}(\mathbf{X}'\mathbf{X})^{-1} \tag{14}$$

$$= \frac{\sum_{1}^{n} (y_{i} - \hat{y}_{i})^{2}}{n - k - 1} (\mathbf{X}'\mathbf{X})^{-1}$$
 (15)

What does this equation tell us?

$$Var(\hat{\boldsymbol{\beta}}_{homoskedasticity}) = \frac{\sum_{1}^{n} (y_i - \hat{y}_i)^2}{n - k - 1} (\mathbf{X}'\mathbf{X})^{-1}$$
 (16)

- ► There are a couple things to point out here
 - First, the variance of $\hat{\beta}$ is a function of the variance of the error term
 - The more unexplained variation in the outcome, the larger the variance of our estimated coefficients
 - ightharpoonup Second, the variance of \hat{eta} is also a function of the variance of the covariates
 - The more variation in the covariates, the smaller the variance of our estimated coefficients

Why do we care about variance?

- We care about variance because it allows us to do hypothesis testing
 - ▶ We can test whether or not a coefficient is different from zero (or any other value)
 - ▶ We can test whether or not two coefficients are different from each other
- ▶ We only have a *sample*, which means we cannot say anything with certainty
 - We can only say something with a certain degree of confidence
- ► Taking into account variance allows us to say something about *the population from* which the sample is drawn
 - ► We generally don't care about the sample itself, only what it tells us about something larger

Confidence intervals

- ightharpoonup Suppose we have an estimated coefficient, \hat{eta}
- Let's create a simulation exercise with a "population"

```
# some variable, x, randomly distributed between 0 and 5
x <- runif(100000, min = 0, max = 5)
# y is a function of x, plus some random error
y <- 3*x + rnorm(100000, mean = 0, sd = 1)
# put it into a data frame
df <- as_tibble(cbind(y, x))

# what is the "true" value of beta in our population of 100,000 people?
lm(y - x, data = df)
beta <- coef(lm(y - x, data = df))[2]
beta</pre>
```

Confidence intervals

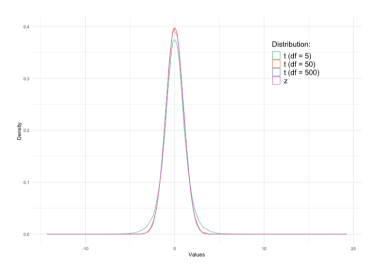
Now let's take a sample of just 50. In a smaller sample, the coefficient can be quite different sometimes.

```
sample <- df[sample(1:nrow(df), 50, replace = FALSE),]</pre>
summary(lm(v ~ x, data = sample))
##
## Call:
## lm(formula = v \sim x. data = sample)
##
## Residuals:
       Min
                 10 Median
                                           Max
## -1.95124 -0.67324 0.05356 0.61053 2.52386
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 0.3640
                           0.3214 1.133
                                          0.263
## v
                2 9222
                           0 1045 27 971 <2e-16 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 1.026 on 48 degrees of freedom
## Multiple R-squared: 0.9422, Adjusted R-squared: 0.941
## F-statistic: 782.4 on 1 and 48 DF. p-value: < 2.2e-16
```

Confidence intervals

- lacktriangle We want to put a confidence interval around our estimate of \hat{eta}
 - ► This will give us some idea of what the "true" value is (we know it here, but we generally do not!)
- ▶ You hopefully remember two separate distributions from earlier classes:
 - z distribution
 - t distribution
- In theory, you should always use a t distribution if you do not know the true population σ^2 (which we usually don't)
 - ► We will always use t distributions

Different distributions



Critical values

- ▶ I'll assume you all remember critical values from earlier metrics classes
 - ► If not, ask now!
 - ightharpoonup Confidence level (e.g. 0.05 for 95% confidence): lpha

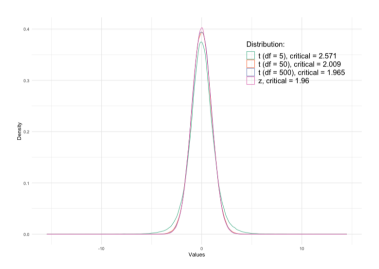
Critical values

- ▶ I'll assume you all remember critical values from earlier metrics classes
 - ► If not, ask now!
 - ightharpoonup Confidence level (e.g. 0.05 for 95% confidence): α
- ▶ We construct our CI as:

$$CI = \left(\hat{\beta} - t_{n-k-1}^{1-\frac{\alpha}{2}} \times \sqrt{Var(\hat{\beta})}, \quad \hat{\beta} + t_{n-k-1}^{1-\frac{\alpha}{2}} \times \sqrt{Var(\hat{\beta})}\right)$$
(17)

- At smaller sample sizes, the t distribution has fatter tails
 - This means that we need to be more uncertain about the true value of β and the critical value can be quite different

Critical values for the different distributions

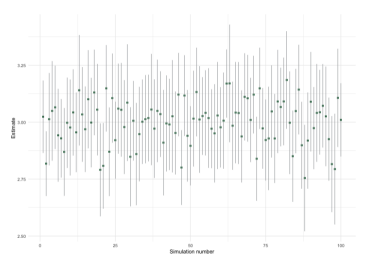


Understanding the confidence interval

- Let's go back to our example with the sample of 50
- ▶ What, exactly, does the confidence interval represent?

- ▶ Suppose we took a bunch of samples of size 50, let's say 100 of them
 - ightharpoonup We would get a bunch of different estimates of \hat{eta}
 - ▶ Let's plot them along with the confidence intervals

Degrees of freedom: n - k - 1 = 50 - 1 - 1 = 48



[1] "Coverage rate is defined as the proportion of samples for which the CI contains the true value: 0.94"

Confidence intervals

- ► Confidence intervals are a frequentist idea
 - ▶ They are about the probability of the true value being in the interval
- We usually can't know for sure whether or not the true value is in the interval
 - ► The probability is only about "averages" over many samples
- ► We can change our confidence level:
 - ightharpoonup lpha = 0.01 (confidence level of 99%) gives us a wider interval
 - ightharpoonup lpha= 0.10 (confidence level of 90%) gives us a narrower interval

Hypothesis testing

- ▶ We can also explicitly test whether or not a coefficient is different from some value
 - ▶ What is the most common value we are interested in?

Hypothesis testing

- We can also explicitly test whether or not a coefficient is different from some value
 - ▶ What is the most common value we are interested in?
 - Zero!
- ▶ We can do this with a t-test
 - Null hypothesis (H_0) : what we assume is true
 - Alternative hypothesis (H_1) : not the null
- Common example with a regression coefficient:
 - ► $H_0: \beta = 0$
 - $H_1: \beta \neq 0$
- ▶ NOTE: We are testing for the POPULATION parameter, not the sample statistics

Hypothesis testing

t-test:

$$\hat{t} = \frac{\hat{\beta} - H_0}{SE(\hat{\beta})},\tag{18}$$

where $SE(\hat{\beta})$ is the standard error of $\hat{\beta}$, or $\sqrt{Var(\hat{\beta})}$. I use \hat{t} here to underline that this comes from our sample statistics.

- ► We can then compare this to a critical value
 - ightharpoonup If $|\hat{t}|>t_{n-k-1}^{1-rac{lpha}{2}}$, then we reject the null
 - lacksquare If $|\hat{t}| < t_{n-k-1}^{1-rac{lpha}{2}}$, then we fail to reject the null

Type 1 and type 2 errors

- ► Type 1 error: we reject the null when it is true
 - $ightharpoonup \alpha$ is the probability of a type 1 error

- ► Type 2 error: we fail to reject the null when it is false
 - ▶ We do not know the probability of a type 2 error

Interpreting the output

```
# using feols (from fixest package) instead of lm now
feols(agemployment ~ log(gnipc) + lfp + log(population) + urbanpop, data = data)
## OLS estimation, Dep. Var.: agemployment
## Observations: 1,420
## Standard-errors: IID
                 Estimate Std. Error t value Pr(>|t|)
## (Intercept)
              105.201328 2.731881 38.50875 < 2.2e-16 ***
## log(gnipc) -10.112581 0.239763 -42.17745 < 2.2e-16 ***
## lfp
               ## log(population) 0.376890 0.129339 2.91398 0.0036246 **
                ## urbanpop
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## RMSE: 7.33605 Adi. R2: 0.834605
```

Something really nice about fixest (feols)

```
# using feels instead of lm now
reg1 <- feols(agemployment ~ log(gnipc) + lfp + log(population) + urbanpop, data = data)
reg2 <- feols(servicesemployment ~ log(gnipc) + lfp + log(population) + urbanpop, data = data)
etable(reg1, reg2,
       se.below = TRUE, se.row = FALSE.
      digits = 3.
      digits.stats = 3.
      fitstat = c("r2", "ar2", "f", "n")
##
                       reg1
                                       reg2
## Dependent Var.: agemployment servicesemployment
##
## Constant
                105.2***
                                   -7.05**
##
                 (2.73)
                                 (2.18)
## log(gnipc)
                 -10.1***
                                  7.35***
##
                 (0.240)
                                   (0.191)
## lfp
                 0.242***
                               -0.048***
##
                  (0.015)
                                   (0.012)
## log(population) 0.377** -0.873***
                  (0.129)
                              (0.103)
##
## urbanpop
                  -0.208***
                            0.267***
##
                  (0.016)
                               (0.013)
## R2
                      0.835
                                      0.849
## Adj. R2
                  0.835
                                    0.849
                1,791.1
                                  1,995.2
## F-test
                  1,420
                                    1,420
## Observations
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

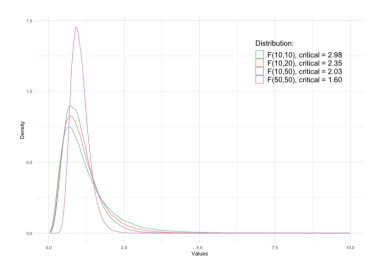
Some important statistics

- $ightharpoonup R^2$: the proportion of variation in the outcome explained by the model
 - $ightharpoonup R^2$ is always between 0 and 1
 - $ightharpoonup R^2$ is a measure of *fit*, but you don't really want to use it to judge how "good" your model is
 - ightharpoonup Adding more variables to your model can NEVER decrease R^2
- Adjusted R^2 (\bar{R}^2): penalizes R^2 for adding more variables
 - ightharpoonup Adjusted R^2 is always between 0 and 1
 - Adjusted R^2 is a measure of fit, but you don't really want to use it to judge how "good" your model is
 - ightharpoonup Adding more variables to your model can decrease adjusted R^2

$$\bar{R}^2 = 1 - (1 - R^2) \frac{n - 1}{n - k - 1} \tag{19}$$

- ► The F distribution is important in linear regression
 - It is the ratio of two chi-squared distributions, so it has two different degrees of freedom
 - We sometimes refer to these as the "numerator" and "denominator" degrees of freedom
 - An F statistic is always positive
- ▶ In the previous regression, the F at the bottom is a test of whether or not all of the coefficients are equal to zero
 - ► This is a joint test of the coefficients
 - ▶ The null hypothesis is that *all* of the coefficients are equal to zero
 - ▶ The alternative hypothesis is that at least one of the coefficients is not equal to zero
- In practice, we usually just use the p-value because of the fact that critical values of F distributions can be quite different depending on the d.f.

F distributions with different degrees of freedom



- t-tests are the workhorse for inference on a *single* coefficient
- But what if we want to test multiple coefficients at once?
 F-tests
- Consider the following regression:

$$y_i = \beta_0 + \beta_1 x_{1,i} + \beta_2 x_{2,i} + \beta_3 x_{3,i} + \epsilon_i$$
 (20)

▶ How might we think about testing whether $\beta_1 = \beta_2 = 0$?

$$y_i = \beta_0 + \beta_1 x_{1,i} + \beta_2 x_{2,i} + \beta_3 x_{3,i} + \epsilon_i$$
 (21)

- If we assume $\beta_1 = \beta_2 = 0$, that is the same as estimating:

$$y_i = \tilde{\beta}_0 + \tilde{\beta}_1 x_{1,i} + \tilde{\epsilon}_i \tag{22}$$

- Intuitively, we want to know whether the first specification is "better" than the second. We can calculate the estimated variance of the residuals in each specification:

$$\hat{\sigma}^2 = \sum_{1}^{n} (y_i - \hat{y}_i)^2 \tag{23}$$

$$\tilde{\sigma}^2 = \sum_{i=1}^{n} (y_i - \tilde{y}_i)^2 \tag{24}$$

$$\hat{\sigma}^2 = \sum_{1}^{n} (y_i - \hat{y}_i)^2 \tag{25}$$

$$\tilde{\sigma}^2 = \sum_{1}^{n} (y_i - \tilde{y}_i)^2 \tag{26}$$

- We can then calculate the F statistic:

$$F = \frac{(\tilde{\sigma}^2 - \hat{\sigma}^2)/q}{\hat{\sigma}^2/(n-k-1)},\tag{27}$$

where q is the number of restrictions we are testing. In this case, q=2 (two restricted coefficients).

$$F = \frac{(\tilde{\sigma}^2 - \hat{\sigma}^2)/q}{\hat{\sigma}^2/(n-k-1)} \tag{28}$$

- ▶ The numerator $((\tilde{\sigma}^2 \hat{\sigma}^2)/q)$ is distributed as χ^2 with q degrees of freedom
- ▶ The denominator $(\hat{\sigma}^2/(n-k-1))$ is distributed as χ^2 with n-k-1 degrees of freedom
- ▶ Their ratio is distributed as F with q and n k 1 degrees of freedom:

$$F = \frac{(\tilde{\sigma}^2 - \hat{\sigma}^2)/q}{\hat{\sigma}^2/(n-k-1)} \sim F(q, n-k-1)$$
 (29)

Now you see why they are sometimes referred to as "numerator" and "denominator" degrees of freedom!

F-test by hand - note the relationship with the t-test for a single coefficient!

```
# full specification
full <- feols(agemployment ~ log(gnipc) + lfp + log(population) + urbanpop, data = data)
# restricted
restricted <- feols(agemployment ~ log(gnipc) + lfp + urbanpop, data = data)
sigma full <- sum((full$resid)^2) # sigma for full
sigma_restricted <- sum((restricted$resid)^2) # sigma for restricted
F <- ((sigma restricted - sigma full)/1)/(sigma full/(full$nobs - 4 - 1)) # calculate F statistic
## [1] 8.49128
full
## OLS estimation, Dep. Var.: agemployment
## Observations: 1.420
## Standard-errors: IID
                  Estimate Std. Error t value Pr(>|t|)
## (Intercept) 105.201328 2.731881 38.50875 < 2.2e-16 ***
## log(gnipc) -10.112581 0.239763 -42.17745 < 2.2e-16 ***
## lfp
               ## log(population) 0.376890 0.129339 2.91398 0.0036246 **
            ## urbanpop
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## RMSE: 7.33605 Adi. R2: 0.834605
```

- For a *single* coefficient, $F = t^2$

We've been assuming homoskedasticity

▶ In the previous examples, we've been assuming *homoskedasticity*

$$\mathbb{E}(\epsilon'\epsilon|\mathbf{X}) = \mathbf{\Omega} = \sigma^2 \mathbf{I} \tag{30}$$

- In practice, this assumption is unlikely to be true
 - You can test for it, but it's not really worth it. Just assume it's not true.
 - lack In other words, assume $\mathbf{\Omega} \neq \sigma^2 \mathbf{I}$
- ► Homoskedasticity can take many forms
 - lt simply means that the variance is not constant across observations
 - Let's talk about a general estimator under heteroskedasticity and another one under clustering

Sandwich estimators

- ► There is a broad class of estimators that are called "sandwich estimators" because of their form
- ► The baseline *heteroskedasticity-consistent* standard errors are:

$$Var(\hat{\boldsymbol{\beta}}) = (\mathbf{X}'\mathbf{X})^{-1} \left(\sum_{i}^{n} X_{i} X_{i}' \sigma_{i}^{2}\right) (\mathbf{X}'\mathbf{X})^{-1}$$
(31)

 \triangleright Of course we don't know σ_i^2 , so we calculate a feasible estimate using the residuals:

$$Var(\hat{\boldsymbol{\beta}})^{HC0} = (\mathbf{X}'\mathbf{X})^{-1} \left(\sum_{i}^{n} X_{i} X_{i}' \hat{e}_{i}^{2}\right) (\mathbf{X}'\mathbf{X})^{-1}$$
(32)

► This is sometimes referred to as *HC*0 standard errors, or **Eicker-White** standard errors

Bias in HC0

- ▶ But there's a problem: it turns out that $\sum_{i} \hat{e}_{i}^{2}$ is a biased estimator for σ_{i}^{2} !
 - ▶ It's biased towards zero ("attenuated") so the standard errors are a bit too small
- ► The next step is to use a scaling factor to "correct" the standard errrors
 - ▶ This is sometimes referred to as *HC*1 standard errors

$$Var(\hat{\beta})^{HC1} = \frac{n}{n - k - 1} (\mathbf{X}'\mathbf{X})^{-1} \left(\sum_{i}^{n} X_{i} X_{i}' \hat{e}_{i}^{2} \right) (\mathbf{X}'\mathbf{X})^{-1}$$
(33)

- ▶ This is a bit ad hoc, but is generally preferred (Hanson, 2022).
 - ► This is currently the default in Stata (with the option robust)

➤ To confuse you more, there are two additional alternatives:

$$Var(\hat{\beta})^{HC2} = (\mathbf{X}'\mathbf{X})^{-1} \left(\sum_{i}^{n} (1 - h_{ii})^{-1} X_{i} X_{i}' \hat{\mathbf{e}}_{i}^{2} \right) (\mathbf{X}'\mathbf{X})^{-1}$$
(34)

$$Var(\hat{\boldsymbol{\beta}})^{HC3} = (\mathbf{X}'\mathbf{X})^{-1} \left(\sum_{i=1}^{n} (1 - h_{ii})^{-2} X_i X_i' \hat{\mathbf{e}}_i^2 \right) (\mathbf{X}'\mathbf{X})^{-1}$$
 (35)

- h_{ii} has to do with a diagonal element of an orthogonal projection matrix. We're going to ignore this for now.
- These are sometimes referred to as HC2 and HC3 standard errors
- ▶ When I refer to HC standard errors, I will be talking about HC1 unless I say otherwise

Clustering

- One of the most common types of heteroskedasticity is clustering
 - ► This is when the error term is correlated within groups
 - For example, if we have data on students within schools, the error term for students in the same school is likely to be correlated
- \triangleright Let K denote the number of clusters and k a specific cluster
 - ▶ We can then calculate the cluster-robust variance estimator as:

$$Var(\hat{\beta})^{cluster} = (\mathbf{X}'\mathbf{X})^{-1} \sum_{k=1}^{K} \left(\sum_{i=1}^{n_k} X_{ik} \hat{\mathbf{e}}_{ik} \right) \left(\sum_{l=1}^{n_k} X_{lk} \hat{\mathbf{e}}_{lk} \right)' (\mathbf{X}'\mathbf{X})^{-1}$$
(36)

- ► The basic intuition is that the more correlated the observations in a cluster is, the larger the variance of the estimated coefficients
 - ► You can actually show this with some algebra and some assumptions
 - ightharpoonup This intra-cluster correlation is often referred to as ρ

Clustering

- \triangleright Suppose you have no clustering at all. Then your effective sample size is n.
- Suppose all of your observations are perfectly correlated with the cluster ($\rho = 1$). Then your effective sample size is just the number of clusters.
- ▶ Of course, neither assumption is likely to ever be true. We are usually somewhere in between, but hopefully this helps with intuition.
- ▶ All of these alternative standard error (variance) calculations only affect the standard errors. **Coefficients do not change.**

Clustering and standard errors

```
##
                     reg1
                               reg2
## Dependent Var.: agemployment agemployment
##
## Constant
              105.2*** 105.2***
               (2.73)
                         (9.40)
## log(gnipc)
                -10.1*** -10.1***
##
               (0.240) (0.713)
## lfp
                0.242*** 0.242***
                (0.015) (0.041)
##
## log(population) 0.377** 0.377
##
                (0.129) (0.462)
## urbanpop
                 -0.208*** -0.208***
##
                (0.016)
                         (0.047)
## R2
                    0.835
                         0.835
## Adi. R2
                0.835
                         0.835
## F-test
              1.791.1 178.5
             1.420
## Observations
                          1.420
## ---
## Signif, codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

What level do we cluster at?

- ► This is a hard question to answer in many cases
 - ► With panel data, we usually cluster at the unit level (e.g. individuals)
 - ► In the previous example, this was the country level
- Always cluster at the level of randomization
 - ► If the data is collected using a complex survey design, we cluster at the the "enumeration area" level
- ► We do not know the "correct" level at which to cluster except in rare situations.

Some complications

- What if the size of clusters is quite different?
 - ► Variance is calculated as the sum of errors within clusters, so different cluster sizes means differences in variance
 - ► This is basically heteroskedasticity
 - ► The key takeaway from this is that smaller samples may show bias (finite-sample bias)
 - ▶ Which leads to...
- ▶ What if *K* is small, i.e. we have relatively few clusters?
 - ightharpoonup One way to think about it is that our effective sample size is closer to K than n
 - Unless we are willing to assume normality of the error term (which we already know we can't), small samples can be problematic
 - ▶ What to do? We will return to this in the next section on bootstrapping.

Before moving on... the regression "assumptions"

Have you seen these before?

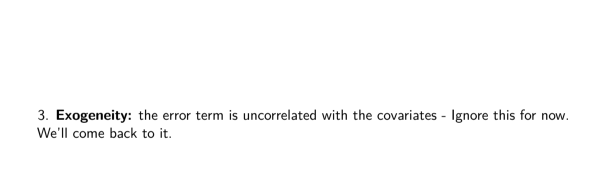
- Linearity: the relationship between the outcome and covariates is linear
- Full rank: the matrix X has full rank
 - Another way to say this: no perfect collinearity
- Exogeneity: the error term is uncorrelated with the covariates
- Momoskedasticity: the variance of the error term is constant
- Normality: the error term is normally distributed

Linearity

1. **Linearity:** the relationship between the outcome and covariates is linear - Regardless of the "true" relationship, OLS will always give you the best linear approximation

Full rank of X

2. **Full rank:** the matrix \mathbf{X} has full rank, or no perfect collinearity - This is required to estimate the regression. - If there is perfect collinearity, then the matrix $\mathbf{X}'\mathbf{X}$ is not invertible - Stata and R will automatically drop variables to make this happen



Full rank of X

Homoskedasticity

4. **Homoskedasticity:** the variance of the error term is constant - Obviously not required! We've already talked about this. There are corrections.

Normality

5. **Normality:** the error term is normally distributed - This is not required for OLS! - The only time this is important is with small samples. - With small samples, we need to assume normality to do hypothesis testing. - With large samples, we can use the central limit theorem to get around non-normality. - A note: the more "non-normal" the error term is, the larger the sample size we need for appropriate inferences.

Bootstrapping (resampling)

- ▶ It turns out that there is an alternative to calculting standard errors using formulas
 - ► It's called bootstrapping
 - ► It's a resampling method
 - ▶ I'll abuse terminology and will generally refer to all of these as "bootstrapping"
- ▶ Bootstrapping originated with Efron (1970)
 - Lots of work since then
 - ► There are many different types of bootstrapping

Resampling with replacement

- Bootstrapping is about sampling from your data
 - ► Something you might not be used to: it is sampling with replacement
- \blacktriangleright Imagine a vector with 10 elements: $\{1, 2, 3, 4, 5, 6, 7, 8, 9, 10\}$
 - Imagine we sample a randomly selected element. Let's say we get 5.
 - Now, we want to take another randomly selected element. The question: do we put the 5 back in or not?
 - ▶ With replacement: put the 5 back in
 - Without replacement: don't put the 5 back in
- Bootstrapping is with replacement
 - ▶ Put the 5 back in!

Resampling with replacement

```
## # A tibble: 10 x 2
     country year
      <chr>>
              <dbl>
    1 AUT
               2003
    2 BGD
               2003
    3 BEL
               2003
    4 BEN
               2003
   5 BRA
               2003
   6 BGR
               2003
## 7 KHM
               2003
   8 CAN
               2003
## 9 CHL
               2003
## 10 COL
               2003
```

Resampling 10 observations with replacement

► The key is that we sample rows — i.e. observations. We do not sample columns. We take an observation and all its values.

```
## # A tibble: 10 x 2
      country
               vear
      <chr>
              <db1>
    1 CHL
               2003
   2 COL
               2003
   3 KHM
               2003
   4 REN
               2003
   5 AUT
               2003
   6 AUT
               2003
   7 BRA
               2003
   8 CAN
               2003
   9 BEN
               2003
## 10 CAN
               2003
```

- ► Suppose we are interested in the variance of a coefficient estimate, beta₁
 - ▶ We can use bootstrapping to estimate the variance of beta₁
 - ► Keeping it simple, here is our regression:

$$y_i = \beta_0 + \beta_1 x_{1,i} + \beta_2 x_{2,i} + \beta_3 x_{3,i} + \beta_4 x_{4,i} + \epsilon_i$$
(37)

- ightharpoonup Imagine a single bootstrap sample, which we will denote with b
 - We are going to call the estimate we are interested in $\hat{\theta}_b$ (which in this case is a coefficient)
- Let's calculate the variance using our bootstrap as follows:

$$Var(\hat{\theta}) = \frac{1}{B-1} \sum_{b=1}^{B} (\hat{\theta}_b - \bar{\theta})^2, \tag{38}$$

where

$$\bar{\theta} = \frac{1}{B} \sum_{b=1}^{B} \hat{\theta}_b \tag{39}$$

- ► In words:
 - ► We take a bunch of bootstrap samples (more is better, but we'll do 1,000)
 - We estimate the regression in each bootstrap sample and save the coefficient we are interested in
 - ► After all 1,000 bootstrap samples, we calculate the variance as the mean (almost) of the squared differences between each bootstrap estimate and the mean of the bootstrap estimates

```
# empty container to hold the estimates
bootvec <- c()
# 1000 samples
for (b in 1:1000){
    # take a sample with replacement (same size as data)
    sample <- data[sample(1:nrow(data), nrow(data), replace = TRUE),]
# estimate regression
    reg <- feols(agemployment - log(gnipc) + lfp + log(population) + urbanpop, data = sample)
# add coefficient to vector
    bootvec <- c(bootvec, reg$coefficients[2])
}
# find variance
var <- sum((bootvec - mean(bootvec))^2)/999
# estimated standard error is the square root
sqrt(var)</pre>
```

[1] 0.2373331

- ► The estimated standard error is the square root of the variance, and it is quite close to the standard error we got from the regression
- ▶ In reality, there's no reason to do this for a single coefficient like this
 - ▶ But it's a good way to understand the intuition behind bootstrapping
- ▶ Where this becomes really powerful is where there is no closed-form solution for the variance
 - For example, what if we wanted to calculate $\frac{\hat{\beta}_2}{\hat{\beta}_2}$?
 - ► To be clear, this is a non-sense example, but it's just to illustrate the point

Our non-sense example

```
# empty container to hold the estimates
bootvec <- c()
# 1000 samples
for (b in 1:1000) {
  # take a sample with replacement (same size as data)
  sample <- data[sample(1:nrow(data), nrow(data), replace = TRUE),]</pre>
  # estimate regression
  reg <- feols(agemployment ~ log(gnipc) + lfp + log(population) + urbanpop, data = sample)
  # add coefficient to vector
  bootvec <- c(bootvec, reg$coefficients[2]/reg$coefficients[3])</pre>
# find variance
var <- sum((bootvec - mean(bootvec))^2)/999
# what's the mean?
mean(bootvec)
## [1] -41.87397
# estimated standard error is the square root
sgrt(var)
## [1] 2.857473
```

Clustered bootstrap

► There's a problem with the previous example: we are assuming that the observations are independent

- What if we think there is clustering?
 - We can use a clustered bootstrap, where we randomly sample CLUSTERS, not observations
 - ▶ In this case, clusters are countries, so we will randomly sample countries

Clustered bootstrap

```
# empty container to hold the estimates
feols(agemployment ~ log(gnipc) + lfp + log(population) + urbanpop, data = data. cluster = "country")
## OLS estimation, Dep. Var.: agemployment
## Observations: 1,420
## Standard-errors: Clustered (country)
                  Estimate Std. Error
                                      t value Pr(>|t|)
## (Intercept) 105.201328 9.400891 11.190570 < 2.2e-16 ***
## log(gnipc) -10.112581 0.713338 -14.176414 < 2.2e-16 ***
                0.241674 0.041227 5.861974 3.0986e-08 ***
## lfp
## log(population) 0.376890 0.462084 0.815631 4.1609e-01
## urbanpop
               ## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## RMSE: 7.33605 Adi. R2: 0.834605
```

```
# empty container to hold the estimates
bootvec <- c()
# get list of countries
countries <- unique(data$country)
# This takes longer to run, so just 500 as an example (try to use more)
for (b in 1:500){
  # take a sample with replacement (same size as data)
 samplecountries <- countries[sample(1:length(countries), length(countries), replace = TRUE)]</pre>
  # go through sample countries and get data, appending
 sample <- c()
 for (c in samplecountries){
    sample <- rbind(sample, data %>% filter(data$country==paste0(c)))
  # estimate regression
 reg <- feols(agemployment ~ log(gnipc) + lfp + log(population) + urbanpop, data = sample)
  # add coefficient to vector
 bootvec <- c(bootvec, reg$coefficients[2])
```

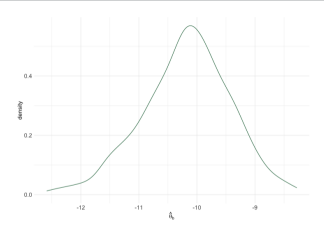
[1] "Standard error: 0.759"

Using the bootstrap vector for confidence intervals

- We can use our bootstrap vector to calculate confidence intervals
 - ▶ We can use the 2.5th and 97.5th percentiles of the bootstrap vector to get a 95% confidence interval:

$$CI = \left(\hat{\theta}_{\alpha/2}, \hat{\theta}_{\alpha/2}\right) \tag{40}$$

Using the bootstrap vector for confidence intervals



- ## [1] "Mean: -10.2"
- ## [1] "Lower: -11.9"
- ## [1] "Upper: -8.75"

Non-parametric

- ► This type of bootstrap is non-parametric
 - ▶ That is, we are not assuming anything about the distribution of the data
 - ► The previous kernel density plot wasn't normally distributed, it was skewed, etc.
 - ► This flexibility is one of the main advantages of bootstrapping
- ► The main downside to bootstrapping is computation
 - ► The previous example takes about 5 minutes to run on my computer with just 500 samples (and it's a good computer)
 - ► More elaborate estimation can take much longer

An actual example from one of my papers

- ► Brummund and Merfeld (2022)
 - ► We want to estimate marginal revenue product of labor (MRPL)
- ► Here is the specification:

$$\ln R_{iht} = \alpha_h + I(Farm = 1) \times \left(\sum_{j} \beta_j \ln \gamma_{jiht} + \frac{1}{2} \sum_{j} \sum_{k} \beta_{jk} \ln \gamma_{jiht} \ln \gamma_{kiht} + \delta C_{iht} + D_{dt} + \eta_m\right) + \sum_{j} \beta_j \ln \gamma_{jiht} + \frac{1}{2} \sum_{j} \sum_{k} \beta_{jk} \ln \gamma_{jiht} \ln \gamma_{kiht} + \delta C_{iht} + D_{dt} + \eta_m$$

$$+ I(Farm = 1) + \varepsilon_{iht}$$

$$(41)$$

An actual example from one of my papers

This is what we need to estimate, using coefficients from the previous equation:

$$\frac{\partial R^f}{\partial L^f} = \frac{\hat{R}_{iht}^f}{L_{iht}^f} \left[\beta_L^f + \beta_{LL}^f \log L_{iht}^f + \beta_{LA} \log A_{iht} + \beta_{LF} \log F_{iht} \right]$$
(42)

$$\frac{\partial R^{nf}}{\partial L^{nf}} = \frac{\hat{R}_{iht}^{nf}}{L_{iht}^{nf}} \left[\beta_L^{nf} + \beta_{LL}^{nf} \log L_{iht}^{nf} + \beta_{LC} \log C_{iht} \right]$$
(43)

- It's not possble to get closed-form solutions for the variance of these
 - So what did we do? Bootstrapped!
 - Household panel data, so clustered bootstrap at the household level