Simple Linear Regression: Inference

EC 320: Introduction to Econometrics

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Prologue

Housekeeping

Midterm

- Pick up your midterms from me in office hours (or send me an email to schedule an appt)
- Solution key will be posted to Canvas

Analytical Problem Set 4

- Due Friday, May 6th by 11:59pm.
- Review and some content we'll talk about today

Last Time

We discussed the classical assumptions of OLS:

- 1. **Linearity:** The population relationship is linear in parameters with an additive error term.
- 2. **Sample Variation:** There is variation in X.
- 3. **Exogeneity:** The X variable is exogenous (i.e., $\mathbb{E}(u|X)=0$).
- 4. **Homoskedasticity:** The error term has the same variance for each value of the independent variable (i.e., $Var(u|X) = \sigma^2$).
- 5. **Non-Autocorrelation:** Any pair of error terms share zero correlation due to having been independently drawn. (i.e., $\mathbb{E}(u_iu_j)=0 \ \forall i \ \mathrm{s.t.} \ i \neq j)$
- 6. **Normality:** The population error term is normally distributed with mean zero and variance σ^2 (i.e., $u \sim N(0, \sigma^2)$)

We restricted our attention to the first 5 assumptions.

Classical Assumptions

Last Time

1. We used the first 3 assumptions to show that OLS is unbiased:

$$\mathbb{E} \Big[\hat{eta} \Big] = eta$$

2. We used the first 5 assumptions to derive a formula for the **variance** of the OLS estimator: $Var(\hat{\beta}) = \frac{\sigma^2}{\sum_{i=1}^n (X_i - \bar{X})^2}$.

Classical Assumptions

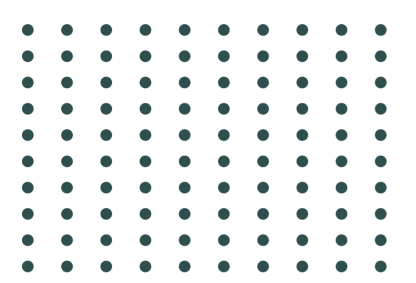
Today

We will use the sampling distribution of $\hat{\beta}$ to conduct hypothesis tests.

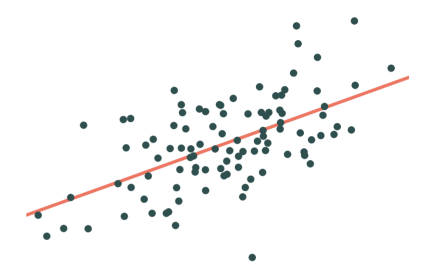
• Can use all 6 classical assumptions to show that OLS is normally distributed:

$$\hat{eta} \sim N \Bigg(eta, rac{\sigma^2}{\sum_{i=1}^n (X_i - ar{X})^2}\Bigg)$$

• We'll "prove" this using R.

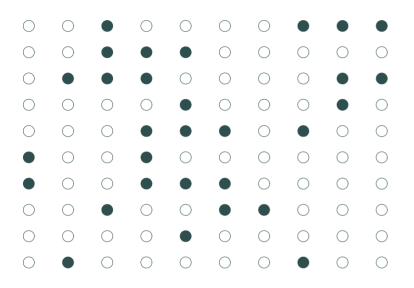


Population

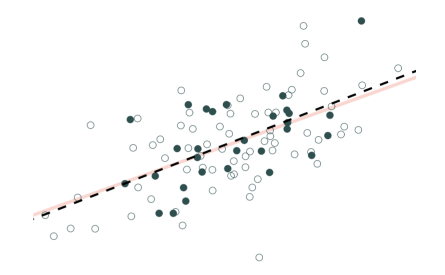


Population relationship

$$Y_i=2.53+0.57X_i+u_i$$
 $Y_i=eta_1+eta_2X_i+u_i$



Sample 1: 30 random individuals

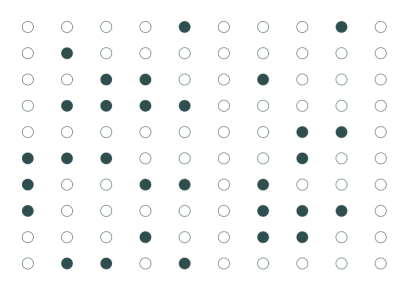


Population relationship

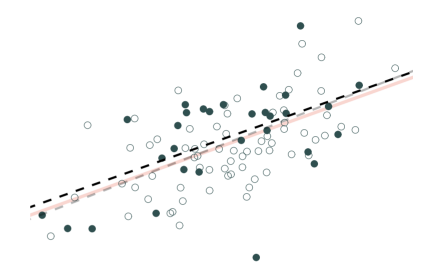
$$Y_i = 2.53 + 0.57X_i + u_i$$

Sample relationship

$$\hat{{Y}}_i = 2.36 + 0.61 X_i$$



Sample 2: 30 random individuals

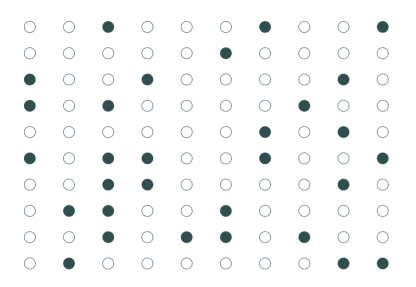


Population relationship

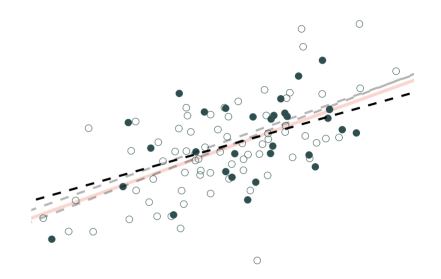
$$Y_i = 2.53 + 0.57Y_i + u_i$$

Sample relationship

$$\hat{Y}_i = 2.79 + 0.56 X_i$$



Sample 3: 30 random individuals



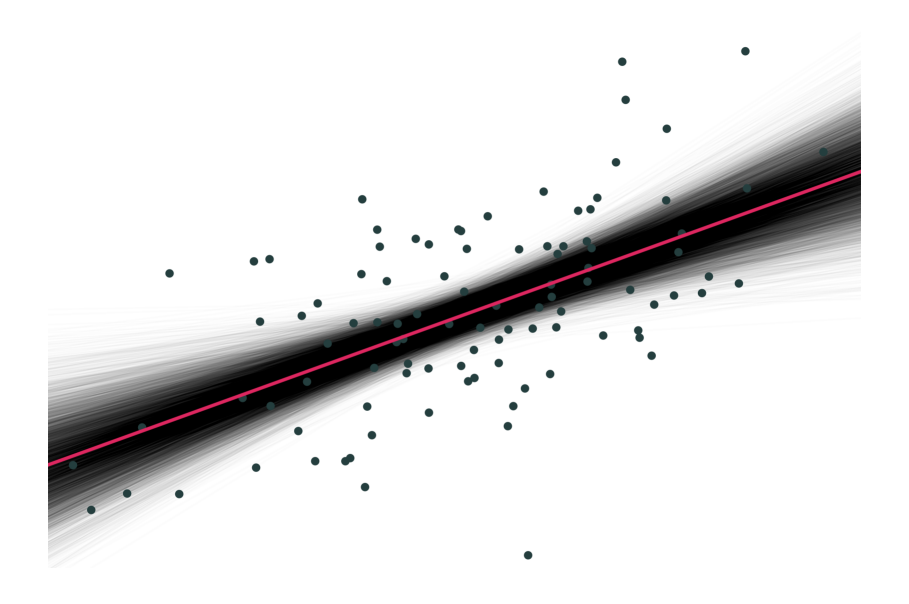
Population relationship

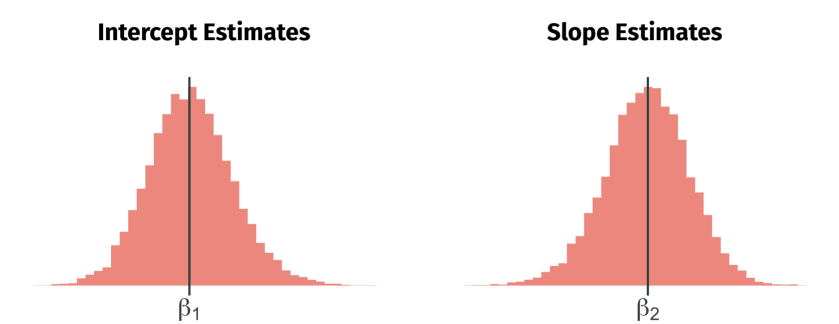
$$Y_i = 2.53 + 0.57X_i + u_i$$

Sample relationship

$$\hat{Y}_i = 3.21 + 0.45 X_i$$

Repeat **10,000 times** (Monte Carlo simulation).





Can you spot the classical assumptions?

```
# Set population and sample sizes
n p \leftarrow 100
n s \leftarrow 30
# Generate population data
pop df ← tibble(
  x = rnorm(n p, mean = 5, sd = 1.5),
  e = rnorm(n_p, mean = 0, sd = 1),
  v = 2.53 + 0.57 * x + e
# Define simulation procedure
sim ols \leftarrow function(x, size = n s) {
  lm(y \sim x, data = pop_df \%>\% sample_n(size = size)) \%>\%
    tidv() %>%
    mutate(iteration = x)
# Run simulation
sim_df \leftarrow map_df(1:10000, \sim sim_ols(.x, size = n_s))
```

Inference

Motivation

What does statistical evidence say about existing theories?

We want to test hypotheses posed by politicians, economists, scientists, people with foil hats, etc.

- Does building a giant wall reduce crime?
- Does shutting down a government adversely affect the economy?
- Does legal cannabis reduce drunk driving or reduce opioid use?
- Do air quality standards improve health or reduce jobs?

While uncertainty exists, we can still conduct *reliable* statistical tests (rejecting or failing to reject a hypothesis).

Inference

We know OLS has some nice properties, and we know how to estimate an intercept and slope coefficient using OLS.

Our current workflow:

- Get data (points with X and Y values).
- ullet Regress Y on X.
- Plot the fitted values (i.e., $\hat{Y}_i = \hat{eta}_0 + \hat{eta}_1 X_i$) and report the estimates.

But how do we actually **learn** something from this exercise?

- Based upon our value of $\hat{\beta}_2$, can we rule out previously hypothesized values?
- How confident should we be in the precision of our estimates?

We need to be able to deal with uncertainty. Enter: Inference.

Inference

We use the standard error of $\hat{\beta}_2$, along with $\hat{\beta}_2$ itself, to learn about the parameter β_2 .

After deriving the distribution of $\hat{\beta}_2$, we have two (related) options for formal statistical inference (learning) about our unknown parameter β_2 :

- **Hypothesis tests:** Determine whether there is statistically significant evidence to reject a hypothesized value or range of values.
- **Confidence intervals:** Use the estimate and its standard error to create an interval that, when repeated, will generally^{††} contain the true parameter.

[†] Hint: It's normal with mean β_2 and variance $\frac{\sigma^2}{\sum_{i=1}^n (X_i - \bar{X})^2}$.

^{††} E.g., similarly constructed 95% confidence intervals will contain the true parameter 95% of the time.

OLS Variance

Hypothesis tests and confidence intervals require information about the variance of the OLS estimator:

$$ext{Var}({\hateta}_2) = rac{\sigma^2}{\sum_{i=1}^n (X_i - ar{X})^2}.$$

Problem

- The variance formula has a population parameter: σ^2 (a.k.a. error variance).
- We can't observe population parameters.
- **Solution:** Estimate σ^2 .

Estimating Error Variance

Learning from our (prediction) errors

We can estimate the variance of u_i (a.k.a. σ^2) using the sum of squared residuals:

$$s_u^2 = rac{\sum_i \hat{u}_i^2}{n-k}$$

where k gives the number of regression parameters.

- In a simple linear regression, k=2.
- s_u^2 is an unbiased estimator of σ^2 .

OLS Variance, Take 2

With
$$s_u^2 = rac{\sum_i \hat{u}_i^2}{n-k}$$
, we can calculate

$$\operatorname{Var}({\hat{eta}}_2) = rac{s_u^2}{\sum_{i=1}^n (X_i - ar{X})^2}.$$

Taking the square root, we get the **standard error** of the OLS estimator:

$$\hat{\mathrm{SE}}ig(\hat{eta}_2ig) = \sqrt{rac{s_u^2}{\sum_{i=1}^n (X_i - ar{X})^2}}.$$

Standard error = sample standard deviation of an estimator.

Standard Errors

R's lm() function estimates standard errors out of the box:

I won't ask you to estimate standard errors by hand!

Null hypothesis (H_0): $\beta_2 = 0$

Alternative hypothesis (H_a): $\beta_2 \neq 0$

There are four possible outcomes of our test:

- 1. We fail to reject the null hypothesis and the null is true.
- 2. We **reject** the null hypothesis and the null is false.
- 3. We **reject** the null hypothesis, but the null is actually true (**Type I error**).
- 4. We **fail to reject** the null hypothesis, but the null is actually false (**Type** II error).

Goal: Make a statement about β_2 using information on $\hat{\beta}_2$.

 \hat{eta}_2 is random: it could be anything, even if $eta_2=0$ is true.

- ullet But if $eta_2=0$ is true, then \hat{eta}_2 is unlikely to take values far from zero.
- As the standard error shrinks, we are even less likely to observe "extreme" values of $\hat{\beta}_2$ (assuming $\beta_2=0$).

Our test should take extreme values of $\hat{\beta}_2$ as evidence against the null hypothesis, but it should also weight them by what we know about the variance of $\hat{\beta}_2$.

Null hypothesis

$$\mathsf{H}_0:eta_2=0$$

Alternative hypothesis

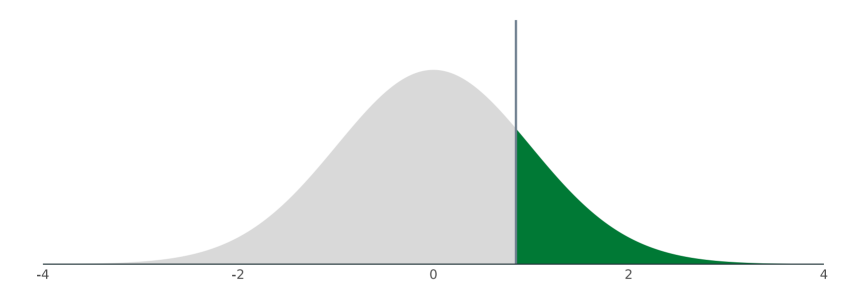
$$\mathsf{H}_\mathsf{a}\!\!:eta_2
eq 0$$

To conduct the test, we calculate a *t*-statistic:

$$t = rac{\hat{eta}_2 - eta_2^0}{\hat{ ext{SE}} \Big(\hat{eta}_2\Big)}$$

- Distributed according to a t-distribution with n-2 degrees of freedom.
- β_2^0 is the value of β_2 in our null hypothesis (e.g., $\beta_2^0=0$).

Next, we use the t-statistic to calculate a p-value.



Describes the probability of seeing a t-statistic as extreme as the one we observe if the null hypothesis is actually true.

But...we still need some benchmark to compare our p-value against.

We worry mostly about false positives, so we conduct hypothesis tests based on the probability of making a Type I error.

How? We select a **significance level** α that specifies our tolerance for false positives. This is the probability of Type I error we choose to live with.



We then compare α to the p-value of our test.

- If the p-value is less than α , then we **reject the null hypothesis** at the $\alpha \cdot 100$ percent level.
- If the p-value is greater than α , then we **fail to reject the null hypothesis**.
- **Note:** Fail to reject \neq accept.

Example: Are campus police associated with campus crime?

$$H_0$$
: $\beta_{Police} = 0$ v.s. H_a : $\beta_{Police} \neq 0$

Significance level: lpha=0.05 (i.e., 5 percent test)

Test Condition: Reject H_0 if p < lpha

p=0.18. Do we reject the null hypothesis?

p-values are difficult to calculate by hand.

Alternative: Compare t-statistic to critical values from the t-distribution.



Notation: $t_{1-\alpha/2,n-2}$ or t_{crit} .

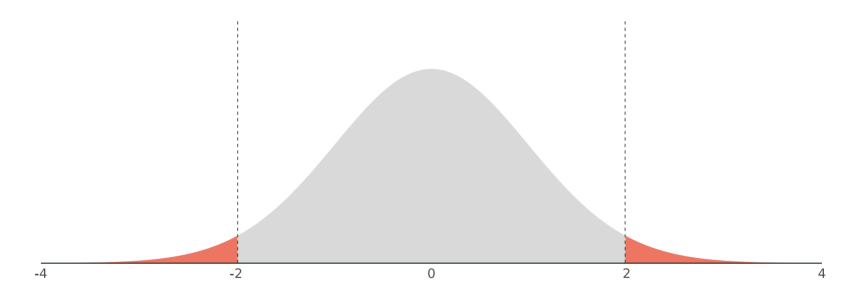
• Find in a t table using the significance level α and n-2 degrees of freedom.

Compare the the critical value to your *t*-statistic:

- If $|t|>|t_{1-lpha/2,n-2}|$, then **reject the null**.
- If $|t|<|t_{1-lpha/2,n-2}|$, then fail to reject the null.

Two-Sided Tests

Based on a critical value of $t_{1-\alpha/2,n-2}=t_{0.975,100}=1.98$, we can identify a **rejection region** on the *t*-distribution.



If our t statistic is in the rejection region, then we reject the null hypothesis at the 5 percent level.

Two-Sided Tests

R defaults to testing hypotheses against the null hypothesis of zero.

$$\mathsf{H}_0$$
: $\beta_2=0$ vs. H_a : $\beta_2\neq 0$

Significance level: lpha=0.05 (i.e., 5 percent test)

 $t_{
m stat}=7.15$ and $t_{0.975,\,28}=2.05$, which implies that p<0.05.

Therefore, we **reject** H_0 at the 5% level.

Two-Sided Tests

Example: Are campus police associated with campus crime?

$$\mathsf{H}_0$$
: $eta_{\mathrm{Police}} = 0$ v.s. H_a : $eta_{\mathrm{Police}}
eq 0$

Significance level: lpha=0.1 (i.e., 10 percent test)

Test Condition: Reject H_0 if $|t| > t_{\mathrm{crit}}$

t=1.35 and $t_{
m crit}=1.66$. Do we reject the null hypothesis?

One-Sided Tests

Sometimes we are confident that a parameter is non-negative or non-positive.

A **one-sided** test assumes that values on one side of the null hypothesis are impossible.

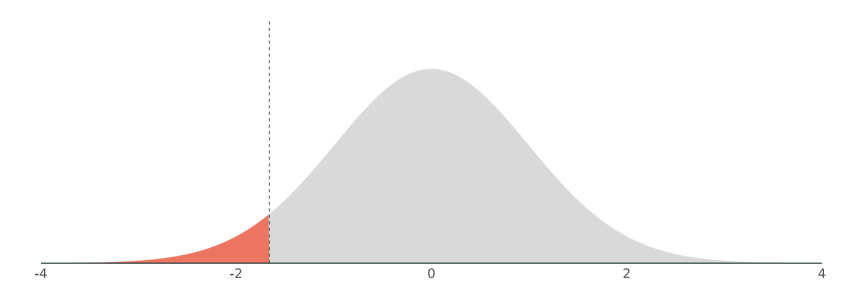
- Option 1: H_0 : $eta_2=0$ vs. H_a : $eta_2>0$
- Option 2: H_0 : $eta_2=0$ vs. H_a : $eta_2<0$

If this assumption is reasonable, then our rejection region changes.

• Same α .

One-Sided Tests

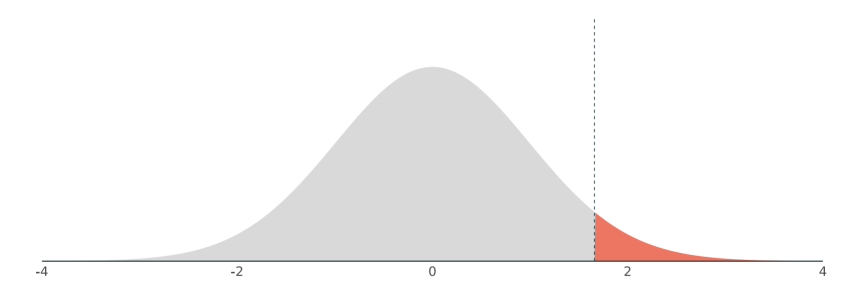
Left-tailed: Based on a critical value of $t_{1-\alpha,n-2}=t_{0.95,100}=1.66$, we can identify a **rejection region** on the *t*-distribution.



If our t statistic is in the rejection region, then we reject the null hypothesis at the 5 percent level.

One-Sided Tests

Right-tailed: Based on a critical value of $t_{1-\alpha,n-2}=t_{0.95,100}=1.66$, we can identify a **rejection region** on the t-distribution.



If our t statistic is in the rejection region, then we reject the null hypothesis at the 5 percent level.

One-Sided Tests

Example: Do campus police deter campus crime?

$$H_0$$
: $eta_{
m Police} = 0$ v.s. H_a : $eta_{
m Police} < 0$

Significance level: lpha=0.1 (i.e., 10 percent test)

Test Condition: Reject H_0 if $t < -t_{
m crit}$

t=1.35 and $t_{
m crit}=1.29$. Do we reject the null hypothesis?

Until now, we have considered **point estimates** of population parameters.

• Sometimes a range of values is more interesting/honest.

We can construct $(1-lpha)\cdot 100$ -percent level confidence intervals for eta_2

$$\hat{eta}_2 \pm t_{1-lpha/2,n-2} \; \hat{\mathrm{SE}} \Big(\hat{eta}_2 \Big)$$

 $t_{1-lpha/2,n-2}$ denotes the 1-lpha/2 quantile of a t distribution with n-2 degrees of freedom.

Q: Where does the confidence interval formula come from?

A: The confidence interval formula comes from the rejection condition of a two-sided test.

Reject
$$\mathsf{H}_0$$
 if $|t| > t_{\mathrm{crit}}$

The test condition implies

Fail to reject H_0 if $|t| \leq t_{\mathrm{crit}}$

which is equivalent to

Fail to reject H_0 if $-t_{\mathrm{crit}} \leq t \leq t_{\mathrm{crit}}$.

Replacing t with its formula gives

Fail to reject
$$\mathsf{H}_0$$
 if $-t_{\mathrm{crit}} \leq rac{\hat{eta}_2 - eta_2^0}{\hat{\mathrm{SE}}\left(\hat{eta}_2
ight)} \leq t_{\mathrm{crit}}.$

Standard errors are always positive, so the inequalities do not flip when we multiply by $\hat{SE}(\hat{\beta}_2)$:

Fail to reject
$$H_0$$
 if $-t_{\mathrm{crit}}\,\hat{\mathrm{SE}}ig(\hat{eta}_2ig) \leq \hat{eta}_2 - eta_2^0 \leq t_{\mathrm{crit}}\,\hat{\mathrm{SE}}ig(\hat{eta}_2ig).$

Subtracting $\hat{\beta}_2$ yields

Fail to reject
$$\mathsf{H}_0$$
 if
$$-\hat{eta}_2 - t_{\mathrm{crit}} \, \hat{\mathrm{SE}} \Big(\hat{eta}_2 \Big) \leq -\beta_2^0 \leq -\hat{eta}_2 + t_{\mathrm{crit}} \, \hat{\mathrm{SE}} \Big(\hat{eta}_2 \Big).$$

Multiplying by -1 and rearranging gives

Fail to reject H_0 if $\hat{eta}_2 - t_{\mathrm{crit}} \, \hat{\mathrm{SE}} \Big(\hat{eta}_2 \Big) \leq eta_2^0 \leq \hat{eta}_2 + t_{\mathrm{crit}} \, \hat{\mathrm{SE}} \Big(\hat{eta}_2 \Big).$

Replacing eta_2^0 with eta_2 and dropping the test condition yields the interval

$$\hat{eta}_2 - t_{
m crit}\,\hat{
m SE}ig(\hat{eta}_2ig) \leq eta_2 \leq \hat{eta}_2 + t_{
m crit}\,\hat{
m SE}ig(\hat{eta}_2ig)$$

which is equivalent to

$$\hat{eta}_2 \pm t_{
m crit} \,\, {
m SE} \Big(\hat{eta}_2 \Big).$$

Insight: A confidence interval is related to a two-sided hypothesis test.

- If a 95 percent confidence interval contains zero, then we fail to reject the null hypothesis at the 5 percent level.
- If a 95 percent confidence interval does not contain zero, then we reject the null hypothesis at the 5 percent level.
- **Generally:** A $(1 \alpha) \cdot 100$ percent confidence interval embeds a two-sided test at the $\alpha \cdot 100$ level.

Example

#> [1] 1.984467

95% confidence interval for β_2 is $0.567 \pm 1.98 \times 0.0793 = [0.410, \, 0.724]$

We have a confidence interval for β_2 , i.e., [0.410, 0.724].

What does it mean?

Informally: The confidence interval gives us a region (interval) in which we can place some trust (confidence) for containing the parameter.

More formally: If we repeatedly sample from our population and construct confidence intervals for each of these samples, then $(1 - \alpha) \cdot 100$ percent of our intervals (e.g., 95%) will contain the population parameter somewhere in the interval.

Now back to our simulation...

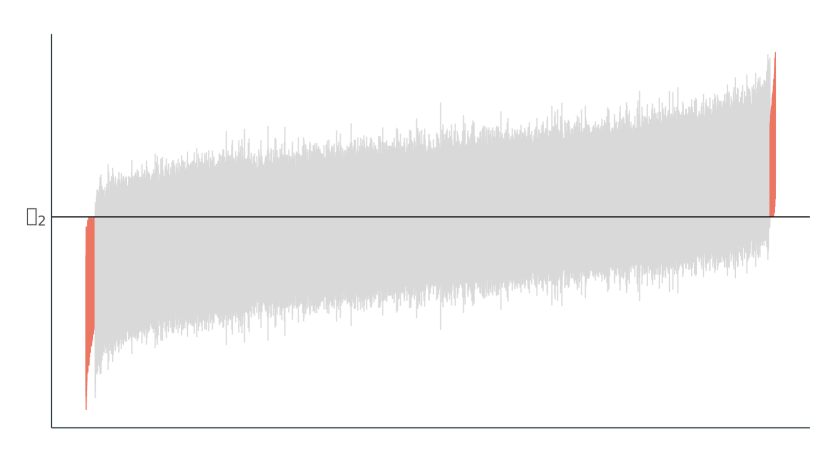
We drew 10,000 samples (each of size n=30) from our population and estimated our regression model for each sample:

$$Y_i = \hat{eta}_1 + \hat{eta}_2 X_i + \hat{u}_i$$

(repeated 10,000 times)

Now, let's estimate 95% confidence intervals for each of these intervals...

From our previous simulation: 97.9% of 95% confidence intervals contain the true parameter value of β_2 .

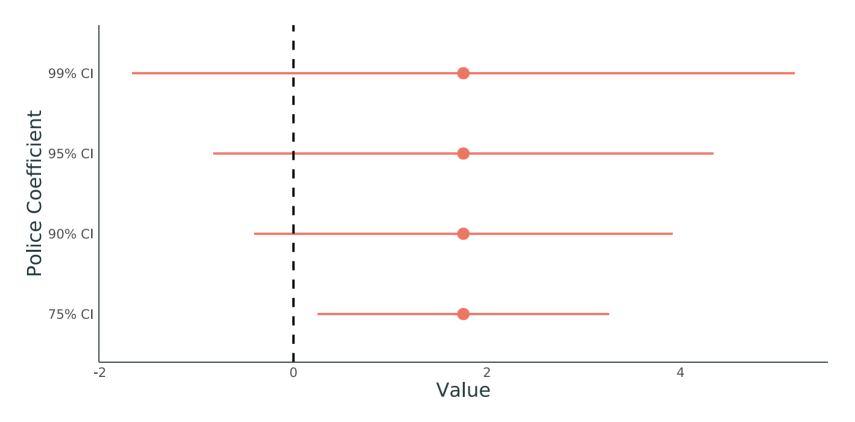


Example: Association of police with crime

You can instruct tidy to return a 95 percent confidence interval for the association of campus police with campus crime:

```
lm(crime ~ police, data = campus) %>% tidy(conf.int = TRUE, conf.level = 0.95)
\#> \# A \text{ tibble: } 2 \times 7
    term estimate std.error statistic p.value conf.low conf.high
#>
    <chr>
                   <dbl>
                             <dbl>
                                       <dbl>
                                                < dbl >
                                                        <dbl>
                                                                  <dbl>
#>
                                        7.75 1.06e-11 13.7
#> 1 (Intercept)
                   18.4
                              2.38
                                                                  23.1
#> 2 police
                    1.76
                              1.30
                                        1.35 1.81e- 1 -0.830
                                                                   4.34
```

Example: Association of police with crime



Four confidence intervals for the same coefficient.