# Section 5: Inference and parallelization

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# 1 Admin

- 1. I posted the solutions for the first problem set on bCourses (Files/Problem Sets/Problem Set 1/problem Set1Solutions.pdf).
- 2. Monday is (technically) a holiday. Move or keep office hours?
- 3. Please take this survey (ASAP) to help me understand how section is helping and failing.

#### 1.1 Follow ups

#### 1.1.1 Redefining missing values

Someone asked about updating values of an existing data frame—for instance changing all .. to NA. As is generally the case in R, you have a few options.

Let's create a simple working example.

```
library(pacman)
p_load(dplyr)
x \leftarrow data.frame(a = c(1:4, NA), b = c("...", 1, 2, 3, "...")) %>% as tibble()
## # A tibble: 5 x 2
##
          a
##
     <int> <chr>
## 1
          1
## 2
         2
                1
## 3
         3
                2
## 4
         4
                3
## 5
        NA
```

**Option 1: Indexing and logical operators** We can use R's standard indexing in conjunction with logical operators to replace the unwanted values:

```
x <- data.frame(a = c(1:4, NA), b = c("...", 1, 2, 3, "...")) %>% as_tibble()
x[x == ".."] \leftarrow NA
Х
## # A tibble: 5 x 2
##
          а
     <int> <chr>
##
## 1
         1
            <NA>
## 2
         2
                1
                2
## 3
         3
                3
## 4
         4
        NA
```

Notice that the class of the b column is still factor, so we would still need to convert the column to numeric ( $x \leftarrow \text{mutate}(x, b = \text{as.numeric}(b))$ )

Option 2: replace and mutate You can combine the (base) replace() function with dplyr's mutate(). replace() wants three arguments: the data (vector), a list of the indices that should be replaced, and the new values. Using replace(), we can write a function that performs our desired replacement. I'll also use as.numeric() to convert result to numeric.

```
# Define our function
rep_na <- function(y) replace(y, which(y == ".."), NA)
# Make some data
x <- data.frame(a = c(1:4, NA), b = c("..", 1, 2, 3, "..")) %>% as_tibble()
```

```
# Replace .. with NA
x %>% mutate(b = rep_na(b) %>% as.numeric())
## # A tibble: 5 x 2
##
         а
     <int> <dbl>
##
## 1
         1
              NA
         2
                1
## 2
## 3
         3
                2
## 4
         4
                3
## 5
        NA
               NA
```

You can use mutate\_all() to apply the function to all columns:

```
# Make some data
x <- data.frame(a = c(1:4, NA), b = c("..", 1, 2, 3, "..")) %>% as_tibble()
# Replace .. with NA
x %>% mutate_all(rep_na) %>% mutate_all(as.numeric)
## # A tibble: 5 x 2
##
         а
##
     <dbl> <dbl>
## 1
         1
         2
## 2
               1
## 3
         3
               2
         4
               3
## 4
## 5
        NA
              NA
```

In addition to mutate\_all(), there are a bunch of other similar functions like mutate\_at(), mutate\_each(), and mutate\_if() (and similar suffixes for all of the standard functions in dplyr). Check them out.

**Note:** Multiple values to NA If you want to convert multiple values to NA (e.g., both ".." and ""), then use the logical operator  $\sin$ , e.g.,  $x[x \sin c("..", "")] <- NA$ .

# 1.1.2 Interactions and polynomials

For R's canned regression functions, you can specify interactions and polynomial terms, but the syntax is not exactly what you might guess:

- Polynomials: For higher-order terms, you need to use the I() function, i.e.,  $x + I(x^2) + I(x^3)$
- Interactions: Use: to specify the interaction between two variables (x1:x2). For those of use who are a bit lazy/efficient, you can use \* to specify the main effects and the interaction (x1\*x2 is equivalent to x1 + x2 + x1:x2).

### 1.1.3 Code folding

RStudio allows you to collapse/expand the current section or all sections. I don't think it has any other flexibility in collapsing/expanding.

#### 1.2 What you will need

# Packages:

• Previously used: dplyr, lfe, readr

• New: magrittr and parallel (parallel is likely already installed)

Data: The auto.csv file.

Review: It will also be helpful if you review the simulations from section 3 and section 4.

#### 1.3 Last week

In our previous section, we discussed logical operators, optional arguments to functions, the Frisch-Waugh-Lovell theorem (and its applications—omitted variable bias, bad controls), and R<sup>2</sup>/measures of fit.

#### 1.4 This week

Testing hypotheses in R! Plus simulation and parallelizing your code.

# 2 Testing hypotheses

This week we will focus on statistically testing hypotheses in R, which generally takes the form of testing whether a point estimate (e.g., a coefficient) is significantly different from some value (generally, zero).

The topic of hypothesis testing is  $super\ important^1$  but perhaps a little overlooked. We spend a lot of time cleaning and pruning our data, demonstrating the unbiasedness (or consistency or other properties) of various estimators, discussing identification strategies<sup>2</sup>, and writing models, and we can easily forget the original point of the exercise. In econometrics, the original point of the exercise is often something like: is there evidence of meaningful relationship between the variables y and x? The point estimate from the regression of y on x doesn't actually answer this question; it gives us a best guess at the slope of a line drawn between y and x. Hypothesis testing (and calculating standard errors) will tell us about the evidence for a relationship between y and x.

# 2.1 Setting up

As we discussed before, I find it helpful to have a section of my R script where I set settings, load libraries, define directories, read data.

```
# Settings
options(stringsAsFactors = F)
# Packages
library(pacman)
p_load(dplyr, lfe, magrittr, readr)
```

<sup>&</sup>lt;sup>1</sup>Not a technical term.

<sup>&</sup>lt;sup>2</sup>If this phrase does not make sense, don't worry. Just wait until ARE 213.

```
# Directories
dir_data <- "/Users/edwardarubin/Dropbox/Teaching/ARE212/Section05/"</pre>
# Load the dataset from CSV
cars <- pasteO(dir_data, "auto.csv") %>% read_csv()
We should also load the functions that we will want to use:
# Functions ----
# Function to convert tibble, data.frame, or tbl_df to matrix
to_matrix <- function(the_df, vars) {</pre>
  # Create a matrix from variables in var
  new_mat <- the_df %>%
    # Select the columns given in 'vars'
    select_(.dots = vars) %>%
    # Convert to matrix
    as.matrix()
  # Return 'new mat'
  return(new_mat)
}
# Function for OLS coefficient estimates
b ols <- function(data, y var, X vars, intercept = TRUE) {</pre>
  # Require the 'dplyr' package
  require(dplyr)
  # Create the y matrix
  y <- to_matrix(the_df = data, vars = y_var)</pre>
  # Create the X matrix
  X <- to_matrix(the_df = data, vars = X_vars)</pre>
  # If 'intercept' is TRUE, then add a column of ones
  if (intercept == T) {
    \# Bind a column of ones to X
    X \leftarrow cbind(1, X)
    # Name the new column "intercept"
    colnames(X) <- c("intercept", X_vars)</pre>
  # Calculate beta hat
  beta hat <- solve(t(X) %*% X) %*% t(X) %*% y
  # Return beta hat
  return(beta hat)
}
```

#### 2.2 magrittr

Notice that we loaded a new package called magrittr. We'll talk about its functions shortly. magrittr offers is more flexibility for piping commands—similar to dplyr's pipe %>%. Fun facts: the package's name comes from the Belgian artist René Magritte. Specifically, the package is referencing Magritte's painting "The Treachery of Images," which shows a pipe and the caption "This is not a pipe." Get it?

#### **2.3** *t* tests

Before we get to the math, you should know that a Guinness brewer named William S. Gosset developed the *t* test to deal with the fact that his sample means did not *act* normally. More.

Okay. Now let's math it up.

Suppose we want to test whether the j<sup>th</sup> coefficient  $\beta_j$  in our model is is significantly different from some other value  $\bar{\gamma}$ , using the significance level  $\alpha$ . Following Max's notes, we can calculate the t statistic for the j<sup>th</sup> coefficient as

$$t_{j} = \frac{b_{j} - \bar{\gamma}}{\sqrt{s^{2} \cdot \left\{ \left(\mathbf{X}'\mathbf{X}\right)^{-1} \right\}_{jj}}} = \frac{b_{j} - \bar{\gamma}}{\operatorname{se}\left(b_{j}\right)}$$

where  $b_j$  is our estimate for  $\beta_j$ ,  $s^2$  is our estimate of  $\sigma^2$ , and  $\left\{ (\mathbf{X}'\mathbf{X})^{-1} \right\}_{jj}$  is the j<sup>th</sup> diagonal element of  $(\mathbf{X}'\mathbf{X})^{-1}$ .

Now let's write a function in R that will calculate this test statistic.

```
t_stat <- function(data, y_var, X_vars, gamma, intercept = T) {
  # Turn data into matrices
  y <- to_matrix(data, y_var)</pre>
  X <- to_matrix(data, X_vars)</pre>
  # Add intercept if requested
  if (intercept == T) X <- cbind(1, X)</pre>
  # Calculate n and k for degrees of freedom
  n \leftarrow nrow(X)
  k \leftarrow ncol(X)
  # Estimate coefficients
  b <- b ols(data, y var, X vars, intercept)
  # Calculate OLS residuals
  e <- y - X %*% b
  # Calculate s^2
  s2 <- (t(e) %*% e) / (n-k)
  # Force s2 to numeric
  s2 %<>% as.numeric()
  # Inverse of X'X
  XX_inv <- solve(t(X) %*% X)
  # Standard error
  se <- sqrt(s2 * diag(XX_inv))</pre>
  # Vector of _t_ statistics
  t_stats <- (b - gamma) / se
  # Return the _t_ statistics
  return(t_stats)
}
```

You could probably write the same function in a single (long) line of code. However, it would be (1) ugly and (2) difficult to debug. I recommend some intermediate steps—even if you are using pipes.

Now let's test drive our function t\_stat() on our car data. Specifically, let's regress price on an intercept, weight, and mpg and test the estimated coefficients against the null hypothesis that they are zero, i.e.,  $\bar{\gamma} = \begin{bmatrix} 0 & 0 & 0 \end{bmatrix}'$ .

We should probably compare our calculated t statistics to the t statistics calculated by the function felm(). For this task, we will grab the coefficients from the summary() function<sup>4</sup> after it is applied to felm(). To grab the coefficients from summary() we will use the %\$% operator from magnitur.

#### 2.3.1 %\$%

Officially, the %\$% pipe is called the *exposition* pipe operator because it *exposes* the names of the left-hand-side object to the right-hand-side expression. More simply: %\$% helps you avoid using too many \$s. For example, if you want to find the correlation between price and weight, you could type cor(cars\$price, cars\$weight) or you could go with cars %\$% cor(price, weight).

```
cor(cars$price, cars$weight)
## [1] 0.5386115
cars %$% cor(price, weight)
## [1] 0.5386115
```

For more on the magrittr package, type vignette("magrittr") into your console. Vignettes are helpful summaries of packages. To see all of the vignettes available, type browseVignettes().

#### 2.3.2 Back to t statistics!

Okay, so let's finally compare our function to felm()'s results.

First, our function:

```
t stat(cars,
  y_var = "price", X_vars = c("mpg", "weight"),
  gamma = 0, intercept = T)
##
                   price
## intercept 0.5410180
              -0.5746808
## mpg
               2.7232382
## weight
And now the t statistics from felm()<sup>5</sup>
felm(price ~ mpg + weight, cars) %>%
  summary() %$% (coefficients)[,3]
## (Intercept)
                        mpg
                                  weight
     0.5410180
                -0.5746808
                               2.7232382
```

<sup>&</sup>lt;sup>3</sup>Bad pun, I know.

<sup>&</sup>lt;sup>4</sup>The summary() function is an example of a generic function in R: when you call summary() on the results from felm(), it is actually using the function summary.felm(). summary() looks at the class of the object you feed it and then decides which function to use to create the summary.

<sup>&</sup>lt;sup>5</sup>The reason we use (coefficients)[,3] is that the coefficients object is actually a matrix with coefficient estimates, standard errors, t statistics, and p-values. The [,3] tells R we only want the third column of this matrix, which is the column with the t statistics.

Huzzah! They match! But unless you have a table of *t* statistics stored in your head, you will want more than a vector of *t* statistics. We want *p*-values.

#### 2.4 p-values

What do we need to do to turn our newly calculated t statistics into p-values? First off, we are going to stick with two-sided hypothesis tests where the null hypothesis is our parameter is zero. We need to find the percentage of the t distribution (with n-k degrees of freedom) that is *more extreme* than (the absolute value) of our test statistic. More formally, the p-value p for our j<sup>th</sup> t statistic t<sub>i</sub> is defined as

$$p = \Pr\left(t_{\mathrm{df}} > |t_i|\right) \times 2$$

Let's code up the calculation of a p-value. We will make use of the function pt(q, df), which gives the distribution function for a t-distributed random variable evaluated at q with df degrees of freedom. In our situation, we will make use of pt()'s optional argument lower.tail. The lower.tail argument defaults to TRUE, which means pt() is evaluating the CDF at q. If we set the lower.tail argument to FALSE, then we will get one minus the CDF, which is what we want for the p-value.

#### Example

```
# The default: lower.tail = TRUE
pt(q = 2, df = 15)
## [1] 0.9680275
# Setting lower.tail = TRUE
pt(q = 2, df = 15, lower.tail = T)
## [1] 0.9680275
# Setting lower.tail = FALSE
pt(q = 2, df = 15, lower.tail = F)
## [1] 0.0319725
```

This situation is a great example of how R plays very nicely with vectors. Imagine we have a vector of t statistics named t\_stats. We can calculate the p-values associated with the t statistics via

```
pt(q = abs(t_stats), df = n-k, lower.tail = F) * 2
```

#### 2.5 Put it all together

Finally, let's put together a nice-ish table like the one that felm() returns: coefficient estimates, standard errors, t statistics, and p-values. Because I'm so clever, we'll call the function that performs this task ols(). In addition, because many of our functions return matrices, we will use as .vector() to convert  $1 \times k$  or  $k \times 1$  matrices to vectors. We will also use the round() function to maintain a reasonable number of digits.

```
ols <- function(data, y_var, X_vars, intercept = T) {
   # Turn data into matrices
   y <- to_matrix(data, y_var)</pre>
```

<sup>&</sup>lt;sup>6</sup>These two assumptions will not always be true in econometrics—but they will be true 99.99999% of the time.

<sup>&</sup>lt;sup>7</sup>Because we are assuming the null hypothesis is always the parameter equals zero, our function with not have a gamma argument. Feel free to change this part.

```
X <- to_matrix(data, X_vars)</pre>
  # Add intercept if requested
  if (intercept == T) X <- cbind(1, X)</pre>
  # Calculate n and k for degrees of freedom
  n \leftarrow nrow(X)
  k <- ncol(X)</pre>
  # Estimate coefficients
  b <- b_ols(data, y_var, X_vars, intercept)</pre>
  # Calculate OLS residuals
  e <- y - X %*% b
  # Calculate s^2
  s2 <- (t(e) %*% e) / (n-k)
  # Update s2 to numeric
  s2 %<>% as.numeric()
  # Inverse of X'X
 XX inv <- solve(t(X) %*% X)
  # Standard error
  se <- sqrt(s2 * diag(XX_inv))</pre>
  # Vector of _t_ statistics
 t stats <- (b - 0) / se
  # Calculate the p-values
  p values = pt(q = abs(t stats), df = n-k, lower.tail = F) * 2
  # Nice table (data.frame) of results
  results <- data.frame(</pre>
    # The rows have the coef. names
    effect = rownames(b),
    # Estimated coefficients
    coef = as.vector(b) %>% round(3),
    # Standard errors
    std_error = as.vector(se) %>% round(3),
    # t statistics
    t_stat = as.vector(t_stats) %>% round(3),
    # p-values
    p_value = as.vector(p_values) %>% round(4)
  # Return the results
  return(results)
Finally, let's test our function and compare it to the output of felm() and the coefficients it outputs in
summary().
ols(data = cars,
 y_var = "price",
 X_vars = c("mpg", "weight"),
  intercept = T)
        effect
                   coef std_error t_stat p_value
## 1 intercept 1946.069 3597.050 0.541 0.5902
## 2
           mpg -49.512
                            86.156 -0.575 0.5673
```

```
0.641 2.723 0.0081
## 3
        weight
                  1.747
felm(price ~ mpg + weight, cars) %>%
  summary() %$% coefficients
                  Estimate
                             Std. Error
                                            t value
                                                       Pr(>|t|)
## (Intercept) 1946.068668 3597.0495988 0.5410180 0.590188628
                -49.512221
                             86.1560389 -0.5746808 0.567323727
## mpg
                              0.6413538 2.7232382 0.008129813
## weight
                  1.746559
Pretty cool, eh?
```

# 3 F tests

As Max has likely told you, there will be situations where you don't care as much about testing the significance of individual coefficients as you care about testing *joint* significance. For example, is there evidence that either  $\beta_1$  or  $\beta_2$  differ significantly from zero? Enter the F test!<sup>8</sup>

# 3.1 The formal part

Assume our data-generating process is

$$\mathbf{y} = \beta_0 + \mathbf{x}_1 \beta_1 + \mathbf{x}_2 \beta_2 + \mathbf{x}_3 \beta_3 + \varepsilon$$

To test whether  $\beta_1$ ,  $\beta_2$ , and  $\beta_3$  are jointly different from zero, we need two more objects:  $\mathbf{R}$  and  $\mathbf{r}$ .  $\mathbf{R}$  is a  $j \times k$  matrix where j is the number of joint hypotheses we will test and k is the number of coefficients in our model (inclusive of the intercept). The  $\mathbf{R}$  matrix creates linear combinations of the coefficients for each of individual hypotheses that we will jointly test. The  $j \times 1$  vector  $\mathbf{r}$  gives the right-hand side of the hypotheses that we want to test.

For example, if we want to jointly test  $\beta_1 = 1$ ,  $\beta_2 = 0$  and  $\beta_3 = 7$ , our matrices would be<sup>9</sup>

$$\mathbf{R} = \begin{bmatrix} 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \end{bmatrix} \text{ and } \mathbf{r} = \begin{bmatrix} 1 \\ 0 \\ 7 \end{bmatrix}$$

If we want to jointly test that  $\beta_1$ ,  $\beta_2$ , and  $\beta_3$  are jointly different from zero, the matrices become

$$\mathbf{R} = \begin{bmatrix} 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \end{bmatrix} \text{ and } \mathbf{r} = \begin{bmatrix} 0 \\ 0 \\ 0 \end{bmatrix}$$

As you saw in lecture, you can write the statistic in the F test as

<sup>&</sup>lt;sup>8</sup>Joint significance is a little bit of a strange idea. We are basically asking if there is evidence that at least one of the coefficients is significantly different from zero.

<sup>&</sup>lt;sup>9</sup>If you are having trouble seeing how  $\bf R$  and  $\bf r$  work, write out the multiplication of  $\bf R\beta = \bf r$  with  $\beta$ s. It should give you a better sense of what is going on here.

$$F = \frac{(\mathbf{R}\mathbf{b} - \mathbf{r})' \left[ \mathbf{R} \left( \mathbf{X}' \mathbf{X} \right)^{-1} \right] (\mathbf{R}\mathbf{b} - \mathbf{r})/J}{s^2} = \frac{(\mathbf{R}\mathbf{b})' \left[ \mathbf{R} \left( \mathbf{X}' \mathbf{X} \right)^{-1} \right] (\mathbf{R}\mathbf{b})/J}{s^2}$$

where the equality comes from the special (though common) case of  $\mathbf{r} = \mathbf{0}$ . In our case, J = 3 because we are jointly testing three hypotheses (also called imposing three restrictions).

#### 3.2 F in R

Now we will write up a function that jointly tests all of the coefficients (except the intercept). Here, I will assume the joint hypotheses in which we are interested take the from  $\beta_i=0$ . You should feel free to write more complex functions that take an arbitrary  ${\bf R}$  and  ${\bf r}$ . I'm also going to assume that we want an intercept.

```
joint_test <- function(data, y_var, X_vars) {</pre>
  # Turn data into matrices
  y <- to_matrix(data, y_var)</pre>
  X <- to_matrix(data, X_vars)</pre>
  # Add intercept
  X \leftarrow cbind(1, X)
  # Name the new column "intercept"
  colnames(X) <- c("intercept", X vars)</pre>
  # Calculate n and k for degrees of freedom
  n \leftarrow nrow(X)
  k \leftarrow ncol(X)
  # J is k-1
  J < -k - 1
  # Create the R matrix: bind a column of zeros
  # onto a J-by-J identity matrix
  R <- cbind(0, diag(J))</pre>
  # Estimate coefficients
  b <- b_ols(data, y_var, X_vars)</pre>
  # Calculate OLS residuals
  e <- y - X %*% b
  # Calculate s^2
  s2 <- (t(e) %*% e) / (n-k)
  # Force s2 to numeric
  s2 %<>% as.numeric()
  # Create the inner matrix R(X'X)^{(-1)}R'
  RXXR <- R %*% solve(t(X) %*% X) %*% t(R)
  # Calculate the F stat
  f_stat <- t(R %*% b) %*% solve(RXXR) %*% (R %*% b) / J / s2
  # Calculate the p-value
  p_value \leftarrow pf(q = f_stat, df1 = J, df2 = n-k, lower.tail = F)
  # Create a data.frame of the f stat. and p-value
  results <- data.frame(</pre>
    f stat = f stat %>% as.vector(),
```

```
p_value = p_value %>% as.vector())
return(results)
}
```

Now run the F test jointly testing the significance of the coefficients on miles per gallon and weight (with the outcome variable price).

# 3.3 Warnings/messages

What if you want to incorporate this F statistic and joint test p-value in your ols() function? You might face a few challenges.

- 1. We created a nice table at the end of the ols() function. Where would we put this new output from the F test? One option is to use a list: you could return a list where the first element is our nice results table and the second element is the information from our F test.
- 2. The second challenge that you face is that our ols() function allows the user to select whether or not to include an intercept, but we need an intercept for our F test. Solution: warning and error messages in R.

For an extreme solution, you could write an if statement that checked whether the user specified intercept = F. If intercept is indeed FALSE, you could stop the function in its tracks and print the error message "No intercept!". In code, this would look like

```
if (intercept == F) stop("No intercept!")
```

However, this solution is a bit extreme: you will not be able to get your results for the other parts of your function if intercept = F. Instead, you might want to use the warning() function to warn the user that because she/he requested no intercept, you will not return an F statistic or joint p-value. Let's do it.

Below is a function ols\_joint() that returns a list of results and issues a warning if the user specifies no intercept.

```
ols_joint <- function(data, y_var, X_vars, intercept = T) {
    # Run the ols() function
    ols_results <- ols(data, y_var, X_vars, intercept)
    # If intercept is T, run the joint_test() function
    # Otherwise, define joint_results to be NA and
    # issue a warning
    if (intercept == T) {</pre>
```

```
joint_results <- joint_test(data, y_var, X_vars)</pre>
  } else {
    warning("No intercept: will not perform F test.")
    joint_results <- data.frame(</pre>
      f stat = NA,
      p_value = NA)
  }
  # Create the results list
  results <- list(ols_results, joint_results)</pre>
  # Return the results list
  return(results)
}
Now let's see what happens when we include and exclude an intercept.
With an intercept:
ols_joint(data = cars,
 y_var = "price",
 X_vars = c("mpg", "weight"),
  intercept = T)
## [[1]]
##
        effect
                   coef std_error t_stat p_value
## 1 intercept 1946.069 3597.050 0.541 0.5902
           mpg -49.512
                         86.156 -0.575 0.5673
## 3
        weight
                  1.747
                            0.641 2.723 0.0081
##
## [[2]]
       f_stat
                   p_value
## 1 14.73982 4.424878e-06
Without an intercept:
ols_joint(data = cars,
 y_var = "price",
 X_vars = c("mpg", "weight"),
  intercept = F)
## Warning in ols_joint(data = cars, y_var = "price", X_vars = c("mpg",
## "weight"), : No intercept: will not perform F test.
## [[1]]
## effect coef std_error t_stat p_value
        mpg -5.479
                      28.122 -0.195 0.8461
## 1
                       0.199 10.431 0.0000
## 2 weight 2.076
##
## [[2]]
## f_stat p_value
## 1 NA
                 NA
```

# 4 Simulation

I think it's time for a simulation. Let's simulate the power (probability that we reject the null when the null is false) of a simple linear regression. We will also incorporate sample size into this simulation. First, let's write a few functions.

#### 4.1 Functions

The first function will generate data for the regression, given a sample size sample\_size. As I mentioned above, we will stick with a simple linear regression

$$\mathbf{y} = \beta_0 + \beta_1 \mathbf{x} + \boldsymbol{\varepsilon}$$

Let's assume  $\beta_0 = 7$  and  $\beta_1 = 0.5$ . We will draw x and  $\varepsilon$  from the standard normal distribution. <sup>10</sup>

```
# Function to generate data
gen_data <- function(sample_size) {
    # Create data.frame with random x and error
    data_df <- data.frame(
        x = rnorm(sample_size),
        e = rnorm(sample_size))
    # Calculate y = 7 + 0.5 x + e; drop 'e'
    data_df %<>% mutate(y = 7 + 0.5 * x + e) %>%
        select(-e)
    # Return data_df
    return(data_df)
}
```

Now we will write a function that takes the output of  $gen_data()$  and feeds it to ols(), which gives us the coefficient estimates and p-value from testing the coefficients against zero.

```
one_sim <- function(sample_size) {
    # Estimate via OLS

    ols_est <- ols(data = gen_data(sample_size),
        y_var = "y", X_vars = "x")
# Grab the estimated coefficient on x
# (the second element of 'coef')
b1 <- ols_est %$% coef[2]
# Grab the second p-value
# (the first p-value is for the intercept)
p_value <- ols_est %$% p_value[2]
# Return a data.frame with b1 and p_value</pre>
```

<sup>&</sup>lt;sup>10</sup>Below you will notice a new pipe operator, %<>%. This operator is also from the magrittr package. This pipe does exactly what our old friend %>% does, but it then assigns the final value to the object on the left-hand side of %<>%. For instance, tmp %<>% mean() takes the object tmp, calculates its mean, and then replaces tmp with the calculated mean. Without using %<>%, one would write tmp <- tmp %>% mean() or tmp <- mean(tmp). The %<>% operator is especially helpful when chaining together several commands with the standard pipe %>%.

```
return(data.frame(b1, p_value))
}
```

Finally, we need to write a function that will run the simulation n\_sims times. One option we used previously is the lapply() function. I'll provide you with an alternative here that uses replicate(). replicate() needs two arguments: the number of replications (n) and the expression (expr) you would like to replicate. You can also give replicate() an argument called simplify which tells the function how to format the results (which type of class). I prefer setting simplify = F so that I know I will get a list. We can then use bind\_rows() on the list to create a nice data frame.

The function will require two arguments: n\_sims, the number of simulations that we want to run, and n, the sample size drawn within each simulation. The function will also accept a third optional argument seed, which is the seed we will use in the simulation. The default value for seed will be 12345.

```
ols_sim <- function(n_sims, sample_size, seed = 12345) {
    # Set the seed
    set.seed(seed)
    # Run one_sim n_sims times; convert results to data.frame
    sim_df <- replicate(
        n = n_sims,
        expr = one_sim(sample_size),
        simplify = F
        ) %>% bind_rows()
    # Return sim_df
    return(sim_df)
}
```

Finally, let's run the ols\_sim() function for two different sample sizes: 10 and 100. For each sample size, we will run the simulation 1,000 times.

**Warning**: Running two-thousand iterations of a simulation can take some time.

```
# Run ols_sim for sample size of 10
sim10 <- ols_sim(n_sims = 1e3, sample_size = 10)
# Run ols_sim for sample size of 100
sim100 <- ols_sim(n_sims = 1e3, sample_size = 100)</pre>
```

# 4.2 Graphs

Let's look at our results. I'm going to use the ggplot2 package. Don't worry about the coding syntax for the moment: we will cover ggplot2 next section. For now, just look at the (pretty) pictures. 12

```
p_load(ggplot2, ggthemes)
```

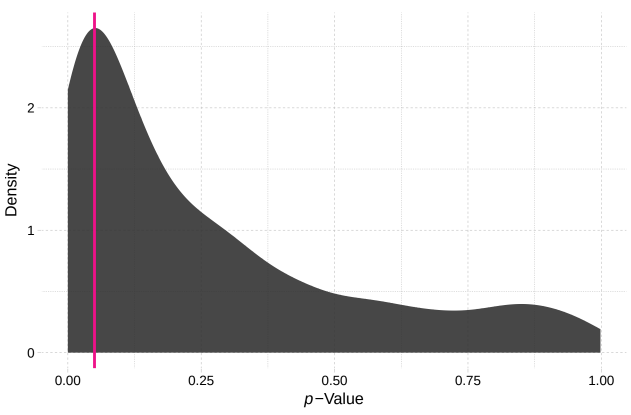
Density plot of the *p*-values for the sample-size 10 simulation

```
ggplot(data = sim10, aes(x = p_value)) +
  stat_density(fill = "grey20", alpha = 0.9) +
```

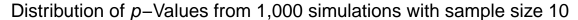
<sup>&</sup>lt;sup>11</sup>You are free to use either function—or an entirely different route. replicate() is a bit simpler, but lapply() has a parallelized version. Your choice. The lapply() call would look something like lapply(X = rep(sample\_size, n\_sims), FUN = one\_sim)

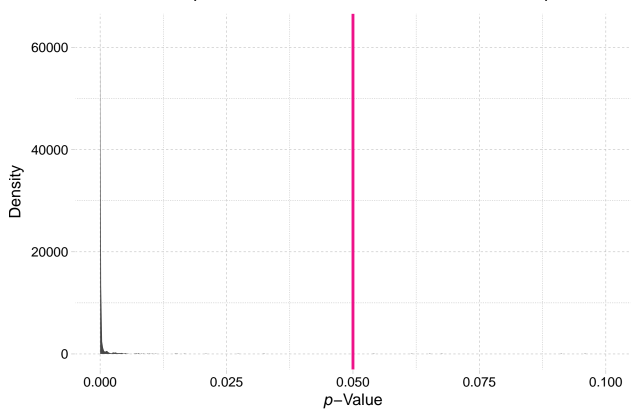
<sup>&</sup>lt;sup>12</sup>If you want to run the code, you will need to install the packages ggplot2 and ggthemes.

# Distribution of *p*–Values from 1,000 simulations with sample size 10



Density plot of the *p*-values for the sample-size 100 simulation

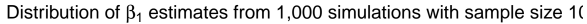


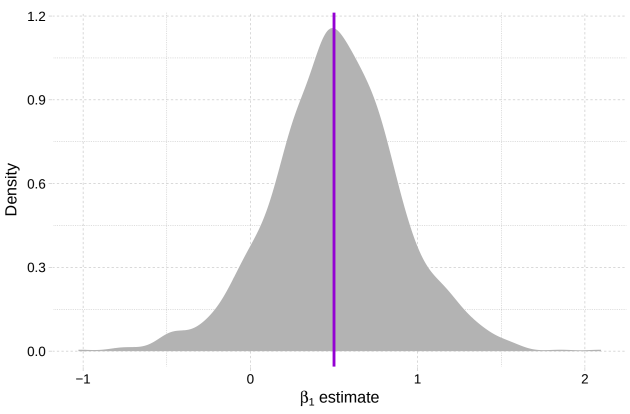


So what is the takeaway here? Power increases with sample size. Why does the second plot look so strange? Nearly all of the *p*-values are approximately zero.

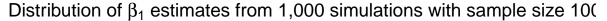
A related topic: How does sample size affect the distribution of estimates for  $\beta_1$ ?

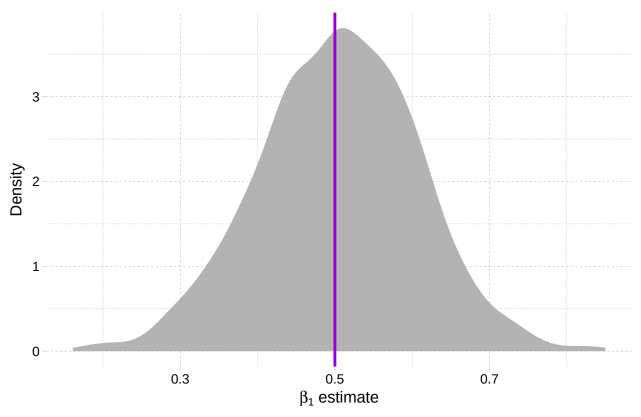
Density plot of the estimates for  $\beta_1$  for sample-size 10 simulation





Density plot of the estimates for  $\beta_1$  for sample-size 100 simulation





So what is going on here? We are seeing the distribution of the estimates for  $\beta_1$  get tighter around the true parameter value as the sample size increases (make sure you note the differences on the x-axis).

# 5 Parallelization

Depending on your computer, the simulations above may have taken a little bit of time. Simulations can sometimes take a long time—you are, after all, asking your computer to repeat a tasks thousands of times. So how can you speed up this process?

- 1. Write more efficient code.
- 2. Parallelize where possible.

Let's assume your/my code is as efficient as it can be (it rarely is). It's time to parallelize. Parallelization in R is fairly straightforward—especially if you are working on a Mac or in a Linux environment (sorry, Windows folks). I will first cover parallelization for Mac/Linux machines. Then I will cover parallelization on Windows machines.

#### 5.1 On Mac and Linux

You will need to install the parallel package. The parallel package offers a number of functions to parallelize your code, but today we will focus on the parallelized version of lapply(): mclapply(). After loading the parallel package, enter ?mclapply into the R console to see the function description. As you can see,

mclapply() starts with the same two arguments as lapply()—X and FUN—and then offers a number of other options. The most important option for us at the moment is mc.cores, which is the option that allows you to choose how many processes to run simultaneously.

```
p_load(parallel)
```

While infinity would be a nice number to enter in the mc.cores argument, you should probably stick with a number less than or equal to the number of cores on your computer. To see how many cores are available on your machine, type detectCores() into the R console.

```
detectCores()
## [1] 4
Looks like I have four.<sup>13</sup>
```

We are now going to parallelize our function ols\_sim(). For this parallelization, we will replace the replicate() function with a similar use of lapply(). However, instead of using plain old lapply(), we will use mclapply() with four cores.

```
ols_sim_mc <- function(n_sims, sample_size, seed = 12345) {
    # Require the parallel package
    require(parallel)
    # Set the seed
    set.seed(seed)
    # Run one_sim n_sims times; convert results to data.frame
    sim_df <- mclapply(
        X = rep(x = sample_size, times = n_sims),
        FUN = one_sim,
        # Specify that we want 4 cores
        mc.cores = 4
        ) %>% bind_rows()
    # Return sim_df
    return(sim_df)
}
```

Now let's run the simulation again—both parallelized and non-parallelized. This time around, let's go for 10,000 iterations. And let's time everything (using proc.time() to get the time at which the simulation starts and stops).

# Not parallelized:

```
# Run ols_sim for sample size of 10
start1 <- proc.time()
sim10 <- ols_sim(n_sims = 1e4, sample_size = 10)
stop1 <- proc.time()
# Run ols_sim for sample size of 100
start2 <- proc.time()
sim100 <- ols_sim(n_sims = 1e4, sample_size = 100)
stop2 <- proc.time()</pre>
```

<sup>&</sup>lt;sup>13</sup>You can sometimes get performance increases by using a number a little larger than your number of cores. I think this has something to do with multi-threaded cores, but we are now getting outside of my area of expertise, so....

#### Parallelized:

```
# Run ols_sim_mc for sample size of 10
start3 <- proc.time()
sim10_mc <- ols_sim_mc(n_sims = 1e4, sample_size = 10)
stop3 <- proc.time()
# Run ols_sim_mc for sample size of 100
start4 <- proc.time()
sim100_mc <- ols_sim_mc(n_sims = 1e4, sample_size = 100)
stop4 <- proc.time()</pre>
```

What are the times? We can check them using the difference between the stop time and the start time for each specification:

```
stop1 - start1
     user system elapsed
   93.131
            1.745 96.832
stop2 - start2
##
     user system elapsed
## 103.799
            2.165 128.594
stop3 - start3
     user
           system elapsed
            2.582 48.605
   99.074
stop4 - start4
     user system elapsed
   98.854
            2.781 51.239
```

The column we care about here is the elapsed column (we can talk about the other columns another time). Comparing the non-parallelized times (the suffixes 1 and 2) and the parallelized times (the suffixes 3 and 4) reveals that parallelization in this setting (using four cores) cuts processing time approximately in half.

# 5.2 On Windows

The idea of parallelization does not change with Windows machines—what we covered above still applies—but the Windows installation of R requires a slightly different method of parallelizing your code. You can use the Windows-style parallelization we cove below on Mac or Linux computers, but you cannot use the Mac/Linux-style parallelization on Windows computers.<sup>14</sup> The only reason I am teaching both is that the Mac/Linux methods are a bit easier.

We will still use the parallel package on Windows machines, but we will cover a few more functions necessary for setting up the parallelization, and we will use the function parLapply() instead of mclapply().

Let's get started. As above, you can check the number of cores available to you using the detectCores() function from the parallel package.

<sup>&</sup>lt;sup>14</sup>The most glaring issue is the fact that the Windows version of mclapply() does not allow mc.cores to be greater than one—*i.e.*, it does not allow parallelization.

```
p_load(parallel)
detectCores()
## [1] 4
```

Assuming you are ready to run some code in parallel, we will now set up a cluster using the makeCluster(), clusterExport(), and clusterEvalQ() functions.

I recommend that you do set the number inside of makeCluster() to be less than or equal to the number of cores that you detected above. You should also know that if you set the number of cores in makeCluster() to be equal to the number of cores you detected, your computer may get really slow while you wait for your parallelized code to finish (e.g., no Youtube).

Let's make a cluster with four cores.

```
# Make the cluster
cl <- makeCluster(4)</pre>
```

Your new cluster is entirely blank. The functions clusterExport() and clusterEvalQ() pass objects from your current environment (what you see when you enter ls() in the R console) to the new clusters. You'll often want to use some of the data/functions/packages you have already loaded to the cluster. What I mean here is even if you have loaded a few packages like dplyr and magrittr, they are not loaded on the cluster.

We can load packages using clusterEvalQ()—let's load dplyr and magrittr.

```
# Load packages to cluster
clusterEvalQ(cl = cl, {
    # Load packages
    library(dplyr)
    library(magrittr)
})
```

Notice that clusterEvalQ() is actually just evaluating the code that we give it. We could do other things here than just loading functions.

We also need to load any user-defined functions that we want to use on the cluster (the functions we wrote ourselves) to the cluster. For this task, we feed a character vector with the function/object names to the function clusterExport(). We have five functions that we will use the in simulation, so let's load them: 15

```
# Load custom functions
clusterExport(cl = cl, c("b_ols", "gen_data",
    "one_sim", "ols", "to_matrix"))
```

You can also load data to the cluster using clusterExport(), for example, clusterExport(cl, "cars"). Alternatively, you could load everything in memory to the cluster using clusterExport(cl, ls()), but this route is a bit lazy and probably will slow your simulations down.

We are finally ready to run our parallelized function. I am going to write it as a function and name the function ols\_sim\_par(). All we are really doing is replacing the replicate() with parLapply(). Note that we need to tell parLapply() which cluster to use (cl) and that fun is no longer capitalized as it was with lapply(). After defining this new simulation function ols\_sim\_par, we need to export it to the cluster, as we did with our other user-defined functions.

<sup>&</sup>lt;sup>15</sup>Now you are seeing why I like the Mac/Linux option for parallelization.

```
# Define the function
ols_sim_par <- function(n_sims, sample_size, seed = 12345) {</pre>
  # Require the parallel package
  require(parallel)
 # Set the seed
  set.seed(seed)
  # Run one sim n sims times; convert results to data.frame
  sim_df <- parLapply(</pre>
    cl = cl,
    X = rep(x = sample_size, times = n_sims),
    fun = one_sim
    ) %>% bind_rows()
  # Return sim_df
  return(sim_df)
# Send it to the cluster
clusterExport(cl, "ols_sim_par")
```

Finally, we can run the simulation. As we did above in the Mac/Linux section, let's time the parallelized and non-parallelized versions and run them for 10,000 iterations.

#### Parallelized:

```
# Run ols sim par for sample size of 10
start5 <- proc.time()</pre>
sim10_par <- ols_sim_par(n_sims = 1e4, sample_size = 10)</pre>
stop5 <- proc.time()</pre>
# Run ols_sim_par for sample size of 100
start6 <- proc.time()</pre>
sim100_par <- ols_sim_par(n_sims = 1e4, sample_size = 100)</pre>
stop6 <- proc.time()</pre>
Not parallelized:
# Run ols_sim for sample size of 10
start7 <- proc.time()</pre>
sim10 <- ols_sim(n_sims = 1e4, sample_size = 10)</pre>
stop7 <- proc.time()</pre>
# Run ols_sim for sample size of 100
start8 <- proc.time()</pre>
sim100 <- ols_sim(n_sims = 1e4, sample_size = 100)</pre>
stop8 <- proc.time()</pre>
When you finish the calculations using your cluster, you need to stop it:
# Stop the cluster
stopCluster(cl)
```

Putting it all together:

And finally, we can compare the time taken for the parallelized and non-parallelized versions of our functions:

```
stop5 - start5
```

```
##
            system elapsed
      user
##
    88.017
             1.801 92.378
stop6 - start6
##
      user
            system elapsed
             1.503 86.924
    83.945
stop7 - start7
##
            system elapsed
      user
##
     0.106
             0.081 32.826
stop8 - start8
            system elapsed
##
      user
##
     0.113
             0.082 31.515
```

The result? Parallelizing your simulation cuts the time in half!<sup>16</sup>

#### 5.3 Notes

The gains from parallelizing your code can be huge—especially when you have access to huge research clusters (the university, Amazon, and Google all host cheap-to-free possibilities for students). However, writing efficient code can give you just as great of gains—and can also be less taxing on your computer (plus you can combine efficient code with parallelization for even bigger gains). Also, the parallelization we covered today will not work if your functions depend on previous outcomes from the simulation. This limitation generally not an issue for econometricians, but you should probably still be aware of it.

#### 5.4 Resources

There are a lot of resources—tutorials, packages, scripts—about parallelizing your R code. A decent place to start is ParallelR.

 $<sup>^{16}</sup>$ We are being a little inaccurate in the calculation of the Windows-style processing times: we should probably include the time that it takes to set up the cluster—not just the time that it takes to run ols\_sim\_par().