Practical Data Analysis for Political Scientists

Brenton Kenkel 2017-02-06

Contents

1	Abo	out This Book	2
2	Prin 2.1 2.2	Mrite Programs for People, Not Computers	3 4 5
3	Working with Data		10
	3.1	Loading	11
	3.2	Tidying	12
	3.3	Transforming and Aggregating	16
	3.4	Merging	19
	3.5	Appendix: Creating the Example Data	24
4	Data Visualization		
	4.1	Basic Plots	28
	4.2	Saving Plots	37
	4.3	Faceting	38
	4.4	Aesthetics	40
	4.5	Appendix: Creating the Example Data	44
5	Bivariate Regression		
	5.1	Probability Refresher	46
	5.2	The Linear Model	48
	5.3	Least Squares	50
	5.4	Properties	54
	5.5	Appendix: Regression in R	56
6	Matrix Algebra: A Crash Course 63		
	6.1	Vector Operations	63
	6.2	Matrix Operations	65
	6.3	Matrix Inversion	68
	6.4	Solving Linear Systems	70
	6.5	Appendix: Matrices in R	70

Chapter 1

About This Book

This book contains the course notes for Brenton Kenkel's course Statistics for Political Research II (PSCI 8357 at Vanderbilt University). It covers the basics of statistical modeling and programming with linear models, along with applications in R.

This book is written in R Markdown and published via Bookdown on GitHub Pages. You can find the R Markdown source files at https://github.com/brentonk/pdaps.

This work is licensed under a Creative Commons Attribution-NonCommercial-ShareAlike 4.0 International License.

Chapter 2

Principles of Programming

It may seem strange to begin a statistics class with two weeks on programming. It is strange. Here is why I have made this strange choice.

First, as a working social scientist, most of the time you spend on data analysis won't be on the *analysis* part. It'll be on obtaining and cleaning the data, to get it in a form that makes sense to analyze. Good programming skills will let you spend less time cleaning data and more time publishing papers.

Second, even if you don't want to develop good programming habits, journals are going to force you to. Every reputable political science journal requires that you provide replication scripts, and some of the best (e.g., *American Journal of Political Science*) have begun auditing the replication materials as a condition of publication. Better to learn The Right Way now when you have lots of time than to be forced to when you're writing a dissertation or on the market or teaching your own courses.

Third, while I feel embarrassed to invoke the cliché that is Big Data, that doesn't mean it's not a real thing. Political scientists have access to more data and more computing power than ever before. You can't collect, manage, clean, and analyze large quantities of data without understanding the basic principles of programming.

As Bowers (2011) puts it, "Data analysis is computer programming." By getting a PhD in political science, by necessity you're going to become a computer programmer. The choice before you is whether to be a good one or a bad one.

Wilson et al. (2014) list eight "best practices for scientific computing." The first two encapsulate most of what you need to know:

- 1. Write programs for people, not computers.
- 2. Let the computer do the work.

¹Or whatever other social science field.

2.1 Write Programs for People, Not Computers

The first two words here—write programs—are crucial. When you are doing analysis for a research project, you should be writing and running scripts, not typing commands into the R (or Stata) console. The console is ephemeral, but scripts are forever, at least if you save them.

Like the manuscripts you will write to describe your findings, your analysis scripts are a form of scientific communication. You wouldn't write a paper that is disorganized, riddled with grammatical errors, or incomprehensible to anyone besides yourself. Don't write your analysis scripts that way either.

Each script should be self-contained, ideally accomplishing one major task. Using an omnibus script that runs every bit of analysis is like writing a paper without paragraph breaks. A typical breakdown of scripts for a project of mine looks like:

- 0-download.r: downloads the data
- 1-clean.r: cleans the data
- 2-run.r: runs the main analysis
- 3-figs.r: generates figures

The exact structure varies depending on the nature of the project. Notice that the scripts are numbered in the order they should be run.

Within each script, write the code to make it as easy as possible for your reader to follow what you're doing. You should indent your code according to style conventions such as http://adv-r.had.co.nz/Style.html. Even better, use the Code -> Reindent Lines menu option in R Studio to automatically indent according to a sane style.

Another way to make your code readable—one that, unfortunately, cannot be accomplished quite so algorithmically—is to add explanatory comments. The point of comments is not to document how the language works. The following comment is an extreme example of a useless comment.

```
# Take the square root of the errors and assign them to
# the output variable
```

output <- sqrt(errors)</pre>

A better use for the comment would be to explain why you're taking the square root of the errors, at least if your purpose in doing so would be unclear to a hypothetical reader of the code.

My basic heuristic for code readability is If I got hit by a bus tomorrow, could one of my coauthors figure out what the hell I was doing and finish the paper?

2.2 Let the Computer Do the Work

Computers are really good at structured, repetitive tasks. If you ever find yourself entering the same thing into the computer over and over again, you are Doing It Wrong. Your job as the human directing the computer is to figure out the structure that underlies the repeated task and to program the computer to do the repetition.

For example, imagine you have just run a large experiment and you want to estimate effects by subgroups.² Your respondents differ across four variables—party ID (R or D), gender (male or female), race (white or nonwhite), and education (college degree or not)—giving you 16 subgroups. You *could* copy and paste your code to estimate the treatment effect 16 times. But this is a bad idea for a few reasons.

- Copy-paste doesn't scale. Copy-paste is managable (albeit misguided) for 16 iterations, but probably not for 50 and definitely not for more than 100.
- Making changes becomes painful. Suppose you decide to change how you calculate the estimate. Now you have to go back and individually edit 16 chunks of code.
- Copy-paste is error-prone, and insidiously so. If you do the calculation wrong all 16 times, you'll probably notice. But what if you screwed up for just one or two cases? Are you *really* going to go through and check that you did everything right in each individual case?

We're going to look at the most basic ways to get the computer to repeat structured tasks—functions and control flow statements. To illustrate these, we will use a result you discussed in Stat I: the central limit theorem.

The central limit theorem concerns the sampling distribution of the sample mean,

$$\bar{X} = \frac{1}{N} \sum_{n=1}^{N} X_n,$$

 $^{^2}$ There could be statistical problems with this kind of analysis, at least if the subgroups were specified post hoc. See https://xkcd.com/882/ ("Significant"). We're going to leave this issue aside for now, but we'll return to it later when we discuss the statistical crisis in science.

where each X_n is independent and identically distributed with mean μ and variance σ^2 . Loosely speaking, the CLT says that as N grows large, the sampling distribution of \bar{X} becomes approximately normal with mean μ and variance σ^2/N .

Here's what we would need to do to see the CLT in practice. We'd want to take a bunch of samples, each of size N, and calculate the sample mean of each. Then we'd have a sample of sample means, and we could check to verify that they are approximately normally distributed with mean μ and variance σ^2/N . This is a structured, repetitive task—exactly the kind of thing that should be programmed. We'll try it out with a random variable from a Poisson distribution with $\lambda = 3$, which has mean $\mu = 3$ and variance $\sigma^2 = 3$.

First things first. We can use the rpois function to draw a random sample of N numbers from the Poisson distribution.

```
samp <- rpois(10, lambda = 3)
samp</pre>
```

```
## [1] 2 3 8 3 5 4 3 4 2 2
```

To calculate the sample mean, we simply use the mean function.

```
mean(samp)
```

```
## [1] 3.6
```

We are interested in the distribution of the sample mean across many samples like this one. To begin, we will write a **function** that automates our core task—drawing a sample of N observations from Poisson(3) and calculating the sample mean. A function consists of a set of *arguments* (the inputs) and a *body* of code specifying which calculations to perform on the inputs to produce the output.

```
pois_mean <- function(n_obs) {
  samp <- rpois(n_obs, lambda = 3)
  ans <- mean(samp)
  return(ans)
}</pre>
```

This code creates a function called pois_mean. It has a single argument, called n_obs. It generates a random sample of n_obs draws from Poisson(3) and calculates its sample mean. It then returns the sample mean as the output.

Let's try calling this function a few times, each with a sample size of N=30. Your output will differ slightly from what's printed here, since the function is generating random numbers.

```
pois_mean(n_obs = 30)
## [1] 3.0333
pois_mean(n_obs = 30)
## [1] 2.4667
```

```
pois_mean(n_obs = 30)
```

[1] 2.9667

Remember that what we're interested in is the *sampling distribution* of the sample mean—the distribution of the sample mean across every possible sample of N observations. We can approximate this distribution by running **pois_mean** many times (e.g., 1000 or more). This would be infeasible via copy-paste. Instead, we will use a **for loop**.

```
# Set up a vector to store the output
n_replicates <- 1000
sampling_dist <- rep(NA, n_replicates)

for (i in 1:n_replicates) {
   sampling_dist[i] <- pois_mean(n_obs = 30)
}</pre>
```

Here's how the for loop works. We specified i as the name of the index variable, with values 1:n_replicates. The for loop takes each value in the sequence, assigns it to the variable i, runs the given expression (in this case, assigning the output of pois_mean to the i'th element of sampling_dist), and then moves on to the next value in sequence, until it reaches the end.

Let's take a look at the results and compare them to our expectations.

```
mean(sampling_dist) # Expect 3

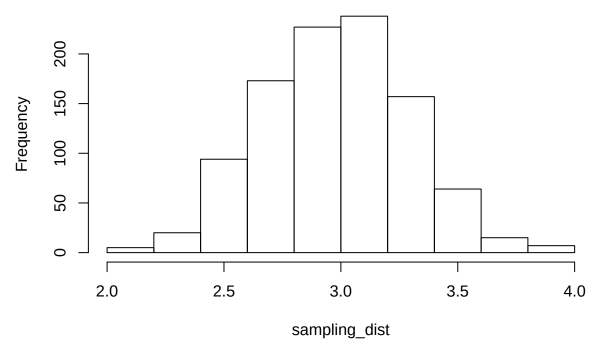
## [1] 2.9952

var(sampling_dist) # Expect 1/10

## [1] 0.096944

hist(sampling_dist) # Expect roughly normal
```

Histogram of sampling_dist



For loops are fun, but don't overuse them. Many simple operations are **vectorized** and don't require a loop. For example, suppose you want to take the square of a sequence of numbers. You could use a for loop ...

```
input <- c(1, 3, 7, 29)
output <- rep(NA, length(input))

for (i in 1:length(input)) {
  output[i] <- input[i]^2
}
output</pre>
```

[1] 1 9 49 841

... but it's faster (in terms of computational speed) and easier to just take advantage of vectorization:

```
input^2
```

[1] 1 9 49 841

Another useful piece of control flow is **if/else statements**. These check a logical condition—an expression whose value is TRUE or FALSE—and run different code depending on the value of the expression. (You may want to catch up on the comparison operators: ==, >, >=, <, <=, etc.)

Let's edit the pois_mean function to allow us to calculate the median instead of the mean. We'll add a second argument to the function, and implement the option using an if/else statement.

```
pois_mean <- function(n_obs, use_median = FALSE) {
  samp <- rpois(n_obs, lambda = 3)
  if (use_median) {
    ans <- median(samp)
  } else {
    ans <- mean(samp)
  }
  return(ans)
}</pre>
```

A couple of things to notice about the structure of the function. We use a comma to separate multiple function arguments. Also, we've specified FALSE as the *default* for the use_median argument. If we call the function without explicitly specifying a value for use_median, the function sets it to FALSE.

```
pois_mean(n_obs = 9)

## [1] 3.7778

pois_mean(n_obs = 9, use_median = TRUE)

## [1] 2

pois_mean(n_obs = 9, use_median = FALSE)

## [1] 2.6667
```

There is a vectorized version of if/else statements called, naturally, the ifelse function. This function takes three arguments, each a vector of the same length: (1) a logical condition, (2) an output value if the condition is TRUE, (3) an output value if the condition is FALSE.

```
x <- 1:10
big_x <- x * 100
small_x <- x * -100

ifelse(x > 5, big_x, small_x)
```

```
## [1] -100 -200 -300 -400 -500 600 700 800 900 1000
```

Functions, for loops, and if/else statements are just a few of the useful tools for programming in R.³ But even these simple tools are enough to allow you to do much more at scale than you could with a copy-paste philosophy.

³Others include the replicate function, the apply family of functions (sapply, lapply, tapply, mapply, ...), the **foreach** package, the **purrr** package, just to name a few of the most useful off the top of my head.

Chapter 3

Working with Data

Some material in this chapter is adapted from notes Matt DiLorenzo wrote for the Spring 2016 session of PSCI 8357.

Let me repeat something I said last week. In your careers as social scientists, starting with your dissertation research—if not earlier—you will probably spend more time collecting, merging, and cleaning data than you will on statistical analysis. So it's worth taking some time to learn how to do this well.

Best practices for data management can be summarized in a single sentence: Record and document everything you do to the data.

The first corollary of this principle is that raw data is sacrosanct. You should never edit raw data "in place". Once you download the raw data file, that file should never change.¹

In almost any non-trivial analysis, the "final" data—the format you plug into your analysis—will differ significantly from the raw data. It may consist of information merged from multiple sources. The variables may have been transformed, aggregated, or otherwise altered. The unit of observation may even differ from the original source. You must document every one of these changes, so that another researcher working from the exact same raw data will end up with the exact same final data.

The most sensible way to achieve this level of reproducibility is to do all of your data merging and cleaning in a script. In other words, no going into Excel and mucking around manually. Like any other piece of your analysis, your pipeline from raw data to final data should follow the principles of programming that we discussed last week.

Luckily for you,² the **tidyverse** suite of R packages (including **dplyr**, **tidyr**, and others) makes it easy to script your "data pipeline". We'll begin by loading the package.

¹Even if it's data you collected yourself, that data should still have a "canonical" representation that never gets overwritten. See Leek (2015) for more on distributing your own data.

²But not for me, because these tools didn't exist when I was a PhD student. Also, get off my lawn!

```
library("tidyverse")
```

3.1 Loading

The first step in working with data is to acquire some data. Depending on the nature of your research, you will be getting some or all of your data from sources available online. When you download data from online repositories, you should keep track of where you got it from. The best way to do so is—you guessed it—to script your data acquisition.

The R function download.file() is the easiest way to download files from URLs from within R. Just specify where you're getting the file from and where you want it to go. For the examples today, we'll use an "untidied" version of the World Development Indicators data from the World Bank that I've posted to my website.

Once you've got the file stored locally, use the utilities from the **readr** package (part of **tidyverse**) to read it into R as a data frame.³ We have a CSV file, so we will use **read_csv**. See help(package = "readr") for other possibilities.

```
untidy_data <- read_csv(file = "my-untidy-data.csv")</pre>
```

```
## Parsed with column specification:
## cols(
     country = col character(),
##
##
     gdp.2005 = col double(),
##
     gdp.2006 = col double(),
     gdp.2007 = col double(),
##
##
     gdp.2008 = col_double(),
     pop.2005 = col double(),
##
##
     pop.2006 = col double(),
##
     pop.2007 = col_double(),
     pop.2008 = col double(),
##
##
     unemp.2005 = col double(),
##
     unemp.2006 = col double(),
##
     unemp.2007 = col double(),
##
     unemp.2008 = col double()
## )
```

Remember that each column of a data frame might be a different type, or more formally *class*, of object. read csv and its ilk try to guess the type of data each column contains: character,

³More precisely, the **readr** functions produce output of class "tbl_df" (pronounced "tibble diff," I'm told), which are like data frames but better. See help(package = "tibble") for what can be done with tbl_dfs.

integer, decimal number ("double" in programming-speak), or something else. The readout above tells you what guesses it made. If it gets something wrong—say, reading a column as numbers that ought to be characters—you can use the col_types argument to set it straight.

FYI, you could also run read csv() directly on a URL, as in:

```
read_csv("http://bkenkel.com/data/untidy-data.csv")
```

However, in analyses intended for publication, it's usually preferable to download and save the raw data. What's stored at a URL might change or disappear, and you'll need to have a hard copy of the raw data for replication purposes.

Now let's take a look at the data we've just loaded in.

```
head(untidy_data)
```

```
## # A tibble: 6 × 13
##
     country gdp.2005 gdp.2006 gdp.2007 gdp.2008
                                                                pop.2006
                                                     pop.2005
                                    <dbl>
##
       <chr>
                 <dbl>
                          <dbl>
                                              <dbl>
                                                        <dbl>
                                                                   <dbl>
## 1
          AD
                3.8423
                         4.0184
                                   4.0216
                                            3.6759
                                                     0.081223
                                                                0.083373
## 2
          AE 253.9655 278.9489 287.8318 297.0189
                                                     4.481976
                                                                5.171255
## 3
          AF
                9.7630
                        10.3052
                                  11.7212
                                           12.1445 24.399948 25.183615
## 4
          AG
                1.1190
                         1.2687
                                   1.3892
                                            1.3902
                                                     0.082565
                                                                0.083467
## 5
                9.2684
                         9.7718
                                  10.3483
                                           11.1275
                                                     3.011487
                                                                2.992547
          AL
               7.6678
## 6
          AM
                         8.6797
                                   9.8731
                                           10.5544
                                                     3.014917
                                                                3.002161
     ... with 6 more variables: pop.2007 <dbl>, pop.2008 <dbl>,
## #
       unemp.2005 <dbl>, unemp.2006 <dbl>, unemp.2007 <dbl>, unemp.2008 <dbl>
```

We have a country variable giving country abbreviations. The other variables are numerical values: the country's GDP in 2005, 2006, 2007, and 2008; then the same for population and unemployment. Let's get this into a format we could use for analysis.

3.2 Tidying

Wickham (2014) outlines three qualities that make data "tidy":

- 1. Each variable forms a column.
- 2. Each observation forms a row.
- 3. Each type of observational unit forms a table.

For one thing, this means that whether a dataset is tidy or not depends—at least in part (some data collections are messy from any angle)—on the purpose it's being analyzed for.

Each row of untidy_data is a country. In observational studies in comparative politics and international relations, more commonly the unit of observation is the country-year.⁴ How

⁴Insert lame joke about how Americanists haven't heard of other countries. But, seriously, if you're

can we take untidy_data and easily make it into country-year data?

We'll use the **tidyr** package (again, part of **tidyverse**) to clean up this data. The biggest problem right now is that each column, besides the country identifier, really encodes two pieces of information: the year of observation and the variable being observed. To deal with this, we'll have to first transform the data from one untidy format to another. We're going to use the **gather()** function to make each row a country-year-variable.

What gather() does is make a row for each entry from a set of columns. It's probably easiest to understand it by seeing it in practice:

```
## # A tibble: 6 × 3
##
     country variable
                         number
##
       <chr>
                 <chr>
                          <dbl>
## 1
          AD gdp.2005
                         3.8423
## 2
          AE gdp.2005 253.9655
## 3
          AF gdp.2005
                         9.7630
          AG gdp.2005
## 4
                         1.1190
## 5
          AL gdp.2005
                         9.2684
## 6
          AM gdp.2005
                         7.6678
```

With the first argument, we told gather() to use the untidy_data data frame. With the last argument, we told it the set of columns to "gather" together into a single column. The key column specifies the name of the variable to store the "key" (original column name) in, and the value column specifies the name of the variable to store the associated value. For example, the second row of long_data encodes what we previously saw as the gdp.2005 column of untidy data.

Now we have a new problem, which is that variable encodes two pieces of information: the variable and the year of its observation. tidyr provides the separate() function to solve that, splitting a single variable into two.

```
long data <- separate(long data,</pre>
                        col = variable,
                        into = c("var", "year"))
head(long data)
## # A tibble: 6 × 4
##
     country
                var year
                             number
##
       <chr> <chr> <chr>
                              <dbl>
## 1
          AD
                     2005
                             3.8423
                gdp
```

confused because you haven't heard of other countries, just think of "state-years".

```
## 2
                      2005 253.9655
           ΑE
                gdp
## 3
           ΑF
                      2005
                              9.7630
                gdp
## 4
           AG
                      2005
                              1.1190
                gdp
## 5
           AL
                      2005
                              9.2684
                gdp
## 6
           AM
                gdp
                      2005
                              7.6678
```

A tibble: 2 × 3

So now we have country-year-variable data, with the year and variable conveniently stored in different columns. To turn this into country-year data, we can use the **spread()** function, which is like the inverse of **gather()**. **spread()** takes a key column and a value column, and turns each different key into a column of its own.

```
clean data <- spread(long data,</pre>
                     key = var,
                     value = number)
head(clean data)
## # A tibble: 6 × 5
##
     country year
                                   pop unemp
                         gdp
##
       <chr> <chr>
                       <dbl>
                                 <dbl> <dbl>
## 1
                      3.8423 0.081223
          AD
              2005
                                          NA
## 2
          AD
              2006
                      4.0184 0.083373
                                          NA
## 3
          AD 2007
                      4.0216 0.084878
                                          NA
                      3.6759 0.085616
## 4
          AD
              2008
                                          NA
              2005 253.9655 4.481976
## 5
          ΑE
                                         3.1
## 6
          ΑE
              2006 278.9489 5.171255
                                         3.3
```

When using spread() on data that you didn't previously gather(), be sure to set the fill argument to tell it how to fill in empty cells. A simple example:

```
test data
## # A tibble: 3 × 3
##
          id
                  k
##
       <chr> <chr> <dbl>
## 1 brenton
                       10
                  a
## 2 brenton
                       20
                  b
                        5
## 3 patrick
                  b
spread(test data, key = k, value = v)
## # A tibble: 2 × 3
##
          id
                        b
                  а
## *
       <chr> <dbl> <dbl>
## 1 brenton
                 10
                       20
## 2 patrick
                 NA
                        5
spread(test_data, key = k, value = v, fill = 100)
```

```
## id a b
## * <chr> <dbl> <dbl>
## 1 brenton 10 20
## 2 patrick 100 5
```

One more important note on **tidyverse** semantics. It includes a fabulous feature called the *pipe*, %>%, which makes it easy to string together a truly mind-boggling number of commands.

In pipe syntax, x %% f() is equivalent to f(x). That seems like a wasteful and confusing way to write f(x), and it is. But if you want to string together a bunch of commands, it's much easier to comprehend

```
x %>%
f() %>%
g() %>%
h() %>%
i()
```

than i(h(g(f(x)))).

You can pass function arguments using the pipe too. For example, f(x, bear = "moose") is equivalent to x % f(bear = "moose").

The key thing about the **tidyverse** functions is that each of them takes a data frame as its first argument, and returns a data frame as its output. This makes them highly amenable to piping. For example, we can combine all three steps of our tidying above with a single command, thanks to the pipe:⁵

```
## # A tibble: 860 × 5
##
     country
               year
                                   pop unemp
                          gdp
## *
       <chr> <chr>
                       <dbl>
                                 <dbl> <dbl>
## 1
          AD
               2005
                      3.8423 0.081223
                                           NA
## 2
               2006
                      4.0184 0.083373
          AD
                                           NA
## 3
          AD
               2007
                      4.0216 0.084878
                                           NA
## 4
          AD
               2008
                      3.6759 0.085616
                                           NA
## 5
               2005 253.9655 4.481976
          ΑE
                                          3.1
```

⁵If you are reading the PDF copy of these notes (i.e., the ones I hand out in class), the line breaks are eliminated, making the piped commands rather hard to read. I am working on fixing this. For now, you may find the online notes at http://bkenkel.com/pdaps easier to follow.

```
## # ... with 855 more rows
```

Without the pipe, if we wanted to run all those commands together, we would have to write:

Sad!

3.3 Transforming and Aggregating

Tidying the data usually isn't the end of the process. If you want to perform further calculations on the raw, that's where the tools in **dplyr** (part of, you guessed it, the **tidyverse**) come in.

Perhaps the simplest **dplyr** function (or "verb", as the R hipsters would say) is **rename()**, which lets you rename columns.

```
clean_data %>%
  rename(gross_domestic_product = gdp)
```

```
## # A tibble: 860 × 5
     country year gross domestic product
##
                                                pop unemp
## *
       <chr> <chr>
                                     <dbl>
                                              <dbl> <dbl>
## 1
          AD
              2005
                                    3.8423 0.081223
                                                        NA
## 2
          AD 2006
                                    4.0184 0.083373
                                                        NA
## 3
          AD 2007
                                    4.0216 0.084878
                                                        NA
## 4
          AD
              2008
                                    3.6759 0.085616
                                                       NA
              2005
                                  253.9655 4.481976
          ΑE
                                                       3.1
## # ... with 855 more rows
```

The **dplyr** functions, like the vast majority of R functions, do not modify their inputs. In other words, running rename() on clean_data will return a renamed copy of clean_data, but won't overwrite the original.

clean data

```
## 2
          AD
              2006
                     4.0184 0.083373
                                        NA
## 3
          AD 2007
                     4.0216 0.084878
                                        NA
## 4
          AD
              2008
                     3.6759 0.085616
                                        NA
## 5
          ΑE
              2005 253.9655 4.481976
                                       3.1
## # ... with 855 more rows
```

If you wanted to make the change stick, you would have to run:

```
clean_data <- clean_data %>%
  rename(gross_domestic_product = gdp)
```

select() lets you keep a couple of columns and drop all the others. Or vice versa if you use minus signs.

```
clean data %>%
  select(country, gdp)
## # A tibble: 860 × 2
     country
##
                  gdp
## *
       <chr>
                <dbl>
## 1
          AD
               3.8423
## 2
               4.0184
          AD
## 3
          AD
               4.0216
## 4
          AD
               3.6759
## 5
          AE 253.9655
## # ... with 855 more rows
clean_data %>%
  select(-pop)
## # A tibble: 860 × 4
##
     country year
                         gdp unemp
## *
       <chr> <chr>
                      <dbl> <dbl>
## 1
          AD 2005
                     3.8423
                                NA
## 2
          AD 2006
                     4.0184
                                NA
## 3
          AD 2007
                     4.0216
                                NΑ
## 4
          AD 2008
                                NA
                     3.6759
                               3.1
## 5
          AE 2005 253.9655
## # ... with 855 more rows
```

mutate() lets you create new variables that are transformations of old ones.

gdp

##

country year

pop unemp gdppc log_gdppc

2

3

2006

2007

34.9

```
<dbl> <dbl>
##
       <chr> <chr>
                       <dbl>
                                             <dbl>
                                                         <dbl>
## 1
          AD
              2005
                      3.8423 0.081223
                                          NA 47.305
                                                        3.8566
## 2
          AD
              2006
                      4.0184 0.083373
                                          NA 48.198
                                                        3.8753
## 3
          AD
              2007
                      4.0216 0.084878
                                          NA 47.381
                                                        3.8582
              2008
                      3.6759 0.085616
                                          NA 42.935
                                                        3.7597
## 4
          AD
## 5
          ΑE
              2005 253.9655 4.481976
                                         3.1 56.664
                                                        4.0371
## # ... with 855 more rows
```

filter() cuts down the data according to the logical condition(s) you specify.

```
clean_data %>%
  filter(year == 2006)
## # A tibble: 215 × 5
##
     country year
                                    pop unemp
                         gdp
##
       <chr> <chr>
                       <dbl>
                                  <dbl> <dbl>
              2006
                      4.0184
## 1
          AD
                              0.083373
                                           NA
## 2
          ΑE
              2006 278.9489
                              5.171255
                                          3.3
## 3
          AF
              2006
                     10.3052 25.183615
                                          8.8
              2006
## 4
          AG
                      1.2687
                              0.083467
                                           NA
## 5
          AL
              2006
                      9.7718
                              2.992547
                                         12.4
## # ... with 210 more rows
```

summarise() calculates summaries of the data. For example, let's find the maximum unemployment rate.

```
clean data %>%
  summarise(max unemp = max(unemp, na.rm = TRUE))
## # A tibble: 1 × 1
##
     max_unemp
##
         <dbl>
## 1
          37.6
```

This seems sort of useless, until you combine it with the group by() function. If you group the data before summarise-ing it, you'll calculate a separate summary for each group. For example, let's calculate the maximum unemployment rate for each year in the data.

```
clean_data %>%
  group_by(year) %>%
  summarise(max unemp = max(unemp, na.rm = TRUE))
## # A tibble: 4 × 2
##
      year max_unemp
     <chr>
               <dbl>
##
      2005
                37.3
## 1
                36.0
```

```
## 4 2008 37.6
```

summarise() produces a "smaller" data frame than the input—one row per group. If you want to do something similar, but preserving the structure of the original data, use mutate in combination with group_by.

```
clean_data %>%
  group_by(year) %>%
  mutate(max unemp = max(unemp, na.rm = TRUE),
         unemp over max = unemp / max unemp) %>%
  select(country, year, contains("unemp"))
## Source: local data frame [860 x 5]
## Groups: year [4]
##
##
     country year unemp max unemp unemp over max
       <chr> <chr> <dbl>
                              <dbl>
##
                                              <dbl>
## 1
          AD 2005
                               37.3
                       NA
                                                 NA
## 2
          AD 2006
                               36.0
                       NA
                                                 NΑ
          AD 2007
                               34.9
## 3
                       NA
                                                 NA
## 4
          AD
              2008
                       NA
                               37.6
                                                 NA
              2005
                               37.3
                                            0.08311
## 5
          ΑE
                      3.1
## # ... with 855 more rows
```

This gives us back the original data, but with a max_unemp variable recording the highest unemployment level that year. We can then calculate each individual country's unemployment as a percentage of the maximum. Whether grouped mutate or summarise is better depends, of course, on the purpose and structure of your analysis.

Notice how I selected all of the unemployment-related columns with contains("unemp"). See ?select helpers for a full list of helpful functions like this for select-ing variables.

3.4 Merging

Only rarely will you be lucky enough to draw all your data from a single source. More often, you'll be merging together data from multiple sources.

The key to merging data from separate tables is to have consistent identifiers across tables. For example, if you run an experiment, you might have demographic data on each subject in one table, and each subject's response to each treatment in another table. Naturally, you'll want to have a subject identifier that "links" the records across tables, as in the following hypothetical example.

```
subject_data
## # A tibble: 3 × 4
## id gender loves bernie does yoga
```

3

4

5

AD

AD

AD

2006

2007

2008

... with 1,065 more rows

NA

NA

NA

```
##
     <dbl>
            <chr>
                           <chr>>
                                      <chr>
## 1
      1001
              male
                             yes
                                         no
## 2 1002 female
                                        yes
                              no
## 3 1003
              male
                              no
                                         no
subject_response_data
## # A tibble: 6 × 3
##
        id treatment response
##
     <dbl>
                <chr>>
                          <chr>
## 1 1001 read book
                            sad
## 2 1001 watch tv
                            sad
## 3 1002 read book
                          happy
## 4 1002 watch tv
                            sad
## 5 1003 read book
                            sad
## 6 1003 watch_tv
                         happy
Let's practice merging data with our cleaned-up country-year data. We'll take two datasets
from my website: a country-level dataset with latitudes and longitudes, and a country-year-
level dataset with inflation over time.
latlong data <- read_csv("http://bkenkel.com/data/latlong.csv")</pre>
latlong_data
## # A tibble: 245 × 3
##
     country latitude longitude
##
       <chr>
                 <dbl>
                            <dbl>
                42.546
## 1
          AD
                           1.6016
## 2
          ΑE
                23.424
                          53.8478
                         67.7100
## 3
          AF
                33.939
## 4
          AG
                17.061
                        -61.7964
## 5
          AΙ
                18.221
                        -63.0686
## # ... with 240 more rows
inflation_data <- read_csv("http://bkenkel.com/data/inflation.csv")</pre>
inflation_data
## # A tibble: 1,070 × 3
##
     country year inflation
##
       <chr> <int>
                         <dbl>
             2004
## 1
          AD
                            NA
## 2
          AD
              2005
                            NA
```

For your convenience, both of these datasets use the same two-letter country naming scheme

as the original data. Unfortunately, out in the real world, data from different sources often use incommensurate naming schemes. Converting from one naming scheme to another is part of the data cleaning process, and it requires careful attention.

dplyr contains various _join() functions for merging. Each of these take as arguments the two data frames to merge, plus the names of the identifier variables to merge them on. The one I use most often is left_join(), which keeps every row from the first ("left") data frame and merges in the columns from the second ("right") data frame.

For example, let's merge the latitude and longitude data for each country into clean_data.

```
## # A tibble: 860 × 7
##
     country year
                                   pop unemp latitude longitude
                         gdp
##
       <chr> <chr>
                       <dbl>
                                 <dbl> <dbl>
                                                 <dbl>
                                                           <dbl>
## 1
          AD
              2005
                      3.8423 0.081223
                                                42.546
                                                          1.6016
                                          NA
## 2
          AD
              2006
                      4.0184 0.083373
                                          NA
                                                42.546
                                                          1.6016
## 3
          AD 2007
                      4.0216 0.084878
                                          NA
                                                42.546
                                                          1.6016
## 4
              2008
                      3.6759 0.085616
                                          NA
                                                42.546
                                                          1.6016
          AD
## 5
          AF.
              2005 253.9655 4.481976
                                         3.1
                                                23.424
                                                         53.8478
     ... with 855 more rows
```

Since latlong_data is country-level, the value is the same for each year. So the merged data contains redundant information. This is one reason to store data observed at different levels in different tables—with redundant observations, it is easier to make errors yet harder to catch them and fix them.

We can also merge data when the identifier is stored across multiple columns, as in the case of our country-year data. But first, a technical note.⁶ You might notice that the year column of clean_data is labeled <chr>>, as in character data. Yet the year column of inflation_data is labeled <int>>, as in integer data. We can check that by running class() on each respective column.

```
class(clean_data$year)

## [1] "character"

class(inflation_data$year)
```

```
## [1] "integer"
```

From R's perspective, the character string "1999" is a very different thing than the integer number 1999. Therefore, if we try to merge clean_data and inflation_data on the year variable, it will throw an error.

⁶This won't be the case if you got clean_data by loading it in directly from clean-data.csv on my website, since read_csv() will have correctly encoded year as an integer.

Error in eval(substitute(expr), envir, enclos): Can't join on 'year' x 'year' because of

To fix this, let's use mutate() to convert the year column of clean_data to an integer. We probably should have done this in the first place—after all, having the year encoded as a character string would have thrown off plotting functions, statistical functions, or anything else where it would be more natural to treat the year like a number.

```
## # A tibble: 860 × 5
     country year
##
                         gdp
                                   pop unemp
##
       <chr> <int>
                       <dbl>
                                 <dbl> <dbl>
## 1
          AD
              2005
                      3.8423 0.081223
                                          NA
## 2
          AD
              2006
                      4.0184 0.083373
                                          NA
## 3
          AD
             2007
                      4.0216 0.084878
                                          NA
## 4
          AD
              2008
                      3.6759 0.085616
                                          NA
## 5
              2005 253.9655 4.481976
          ΑE
                                         3.1
## # ... with 855 more rows
```

Looks the same as before, except with an important difference: year is now labeled <int>.

Now we can merge the two datasets together without issue. Notice how we use a vector in the by argument to specify multiple columns to merge on.

```
## # A tibble: 860 × 6
##
     country year
                         gdp
                                   pop unemp inflation
##
       <chr> <int>
                                 <dbl> <dbl>
                                                  <dbl>
                       <dbl>
                      3.8423 0.081223
## 1
          AD
              2005
                                          NA
                                                     NA
## 2
          AD
              2006
                      4.0184 0.083373
                                          NA
                                                     NA
## 3
          AD
              2007
                      4.0216 0.084878
                                          NA
                                                     NA
## 4
          AD
              2008
                      3.6759 0.085616
                                          NA
                                                     NA
## 5
          ΑE
              2005 253.9655 4.481976
                                         3.1
                                                     NA
## # ... with 855 more rows
```

You might remember that inflation_data contained some country-years not included in the original data (namely, observations from 2004). If we want the merged data to use the observations from inflation_data rather than clean_data, we can use the right_join() function.

5

ΑE

... with 855 more rows

2005 253.9655 4.481976

```
right_join(clean data,
           inflation data,
           by = c("country", "year"))
## # A tibble: 1,070 \times 6
##
     country year
                       gdp
                                 pop unemp inflation
##
                               <dbl> <dbl>
                                                 <dbl>
       <chr> <int>
                     <dbl>
## 1
          AD 2004
                         NA
                                   NA
                                         NA
                                                    NA
## 2
          AD 2005 3.8423 0.081223
                                         NA
                                                    NA
## 3
          AD 2006 4.0184 0.083373
                                         NA
                                                    NA
## 4
          AD 2007 4.0216 0.084878
                                         NA
                                                    NA
          AD 2008 3.6759 0.085616
## 5
                                         NA
                                                    NA
## # ... with 1,065 more rows
One last common issue in merging is that the identifier variables have different names in the
two datasets. If it's inconvenient or infeasible to correct this by renaming the columns in
one or the other, you can specify the by argument as in the following example.
inflation data <- rename(inflation data,
                           the country = country,
                           the_year = year)
inflation_data
## # A tibble: 1,070 \times 3
##
     the country the year inflation
##
            <chr>>
                     <int>
                                <dbl>
## 1
                       2004
               AD
                                    NA
## 2
               AD
                      2005
                                    NA
## 3
               AD
                      2006
                                    NΑ
## 4
                       2007
               AD
                                    NA
## 5
               AD
                      2008
                                    NA
## # ... with 1,065 more rows
left_join(clean data,
          inflation data,
          by = c("country" = "the country", "year" = "the year"))
## # A tibble: 860 × 6
##
     country year
                                    pop unemp inflation
                          gdp
##
       <chr> <int>
                       <dbl>
                                 <dbl> <dbl>
                                                   <dbl>
## 1
          AD 2005
                      3.8423 0.081223
                                           NA
                                                      NA
## 2
          AD 2006
                      4.0184 0.083373
                                           NA
                                                      NA
## 3
          AD 2007
                      4.0216 0.084878
                                           NA
                                                      NA
               2008
                      3.6759 0.085616
## 4
          AD
                                           NA
                                                      NA
```

3.1

NA

3.5 Appendix: Creating the Example Data

I used the same tools this chapter introduces to create the untidy data. I may as well include the code to do it, in case it helps further illustrate how to use the **tidyverse** tools (and, as a bonus, the **WDI** package for downloading World Development Indicators data).

First I load the necessary packages.

```
library("tidyverse")
library("WDI")
library("countrycode")
library("stringr")
```

Next, I download the relevant WDI data. I used the WDIsearch() function to locate the appropriate indicator names.

```
country year NY.GDP.MKTP.KD SP.POP.TOTL SL.UEM.TOTL.ZS
##
     iso2c
## 1
        1A Arab World 2005
                                1.6428e+12
                                             313430911
                                                               12.1402
## 2
        1A Arab World 2006
                                1.7629e+12
                                             320906736
                                                               11.3296
                                1.8625e+12
                                             328766559
## 3
        1A Arab World 2007
                                                               10.8961
        1A Arab World 2008
## 4
                                1.9799e+12
                                             336886468
                                                               10.5060
## 5
        1W
                World 2005
                                5.7703e+13 6513959904
                                                                6.1593
                World 2006
                                6.0229e+13 6594722462
                                                                5.9000
## 6
        1W
```

I want to get rid of the aggregates, like the "Arab World" and "World" we see here. As a rough tack at that, I'm going to exclude those so-called countries whose ISO codes don't appear in the **countrycode** package data.⁷

```
dat_countries <- dat_raw %>%
  filter(iso2c %in% countrycode_data$iso2c)
```

Let's check on which countries are left. (I cut it down to max six characters per country name for printing purposes.)

```
dat_countries$country %>%
  unique() %>%
  str_sub(start = 1, end = 6)
```

⁷**countrycode** is a very useful, albeit imperfect, package for converting between different country naming/coding schemes.

```
[1] "Andorr" "United" "Afghan" "Antigu" "Albani" "Armeni" "Angola"
##
##
     [8] "Argent" "Americ" "Austri" "Austra" "Aruba"
                                                      "Azerba" "Bosnia"
##
    [15] "Barbad" "Bangla" "Belgiu" "Burkin" "Bulgar" "Bahrai" "Burund"
    [22] "Benin"
                  "Bermud" "Brunei" "Bolivi" "Brazil" "Bahama" "Bhutan"
##
    [29] "Botswa" "Belaru" "Belize" "Canada" "Congo," "Centra" "Congo,"
##
    [36] "Switze" "Cote d" "Chile"
                                    "Camero" "China"
                                                      "Colomb" "Costa "
##
    [43] "Cuba"
                  "Cabo V" "Curaca" "Cyprus" "Czech " "German" "Djibou"
##
    [50] "Denmar" "Domini" "Domini" "Algeri" "Ecuado" "Estoni" "Egypt,"
##
    [57] "Eritre" "Spain" "Ethiop" "Finlan" "Fiji"
                                                      "Micron" "Faroe "
##
    [64] "France" "Gabon" "United" "Grenad" "Georgi" "Ghana"
##
                                                               "Gibral"
##
    [71] "Greenl" "Gambia" "Guinea" "Equato" "Greece" "Guatem" "Guam"
    [78] "Guinea" "Guyana" "Hong K" "Hondur" "Croati" "Haiti"
##
                                                               "Hungar"
    [85] "Indone" "Irelan" "Israel" "Isle o" "India"
##
                                                      "Iraq"
                                                               "Iran, "
    [92] "Icelan" "Italy"
                           "Jamaic" "Jordan" "Japan"
                                                      "Kenya"
                                                               "Kyrgyz"
##
    [99] "Cambod" "Kiriba" "Comoro" "St. Ki" "Korea," "Korea," "Kuwait"
##
## [106] "Cayman" "Kazakh" "Lao PD" "Lebano" "St. Lu" "Liecht" "Sri La"
## [113] "Liberi" "Lesoth" "Lithua" "Luxemb" "Latvia" "Libya"
                                                               "Morocc"
## [120] "Monaco" "Moldov" "Monten" "St. Ma" "Madaga" "Marsha" "Macedo"
## [127] "Mali"
                  "Myanma" "Mongol" "Macao " "Northe" "Maurit" "Malta"
## [134] "Maurit" "Maldiv" "Malawi" "Mexico" "Malays" "Mozamb" "Namibi"
## [141] "New Ca" "Niger" "Nigeri" "Nicara" "Nether" "Norway" "Nepal"
## [148] "Nauru" "New Ze" "Oman"
                                    "Panama" "Peru"
                                                      "French" "Papua "
## [155] "Philip" "Pakist" "Poland" "Puerto" "West B" "Portug" "Palau"
## [162] "Paragu" "Qatar" "Romani" "Serbia" "Russia" "Rwanda" "Saudi "
## [169] "Solomo" "Seyche" "Sudan"
                                    "Sweden" "Singap" "Sloven" "Slovak"
## [176] "Sierra" "San Ma" "Senega" "Somali" "Surina" "South " "Sao To"
## [183] "El Sal" "Sint M" "Syrian" "Swazil" "Turks " "Chad"
## [190] "Thaila" "Tajiki" "Timor-" "Turkme" "Tunisi" "Tonga" "Turkey"
## [197] "Trinid" "Tuvalu" "Tanzan" "Ukrain" "Uganda" "United" "Urugua"
## [204] "Uzbeki" "St. Vi" "Venezu" "Britis" "Virgin" "Vietna" "Vanuat"
## [211] "Samoa" "Yemen," "South " "Zambia" "Zimbab"
```

With that out of the way, there's still some cleaning up to do. The magnitudes of GDP and population are too large, and the variable names are impenetrable. Also, the country variable, while helpful, is redundant now that we're satisfied with the list of countries remaining.

dat_countries_wide <- dat_countries_long %>%
 spread(key = var_year, value = value)

```
head(dat countries)
##
     country year
                       gdp
                                 pop unemp
## 1
          AD 2005
                    3.8423 0.081223
                                         NA
## 2
          AD 2006
                    4.0184 0.083373
                                         NA
## 3
          AD 2007
                    4.0216 0.084878
                                        NΑ
          AD 2008
                    3.6759 0.085616
## 4
                                        NA
## 5
          AE 2005 253.9655 4.481976
                                       3.1
          AE 2006 278.9489 5.171255
## 6
                                       3.3
Now I convert the data to "long" format.
dat countries long <- dat countries %>%
  gather(key = variable,
         value = value,
         gdp:unemp)
head(dat countries long)
##
     country year variable
                              value
## 1
          AD 2005
                        gdp
                              3.8423
## 2
          AD 2006
                        gdp 4.0184
## 3
          AD 2007
                        gdp
                              4.0216
## 4
          AD 2008
                        gdp
                              3.6759
          AE 2005
## 5
                        gdp 253.9655
## 6
          AE 2006
                        gdp 278.9489
I then smush variable and year into a single column, and drop the individual components.
dat countries long <- dat countries long %>%
  mutate(var year = paste(variable, year, sep = ".")) %>%
  select(-variable, -year)
head(dat_countries_long)
##
     country
               value var_year
## 1
          AD
               3.8423 gdp.2005
## 2
          AD
               4.0184 gdp.2006
## 3
          AD
               4.0216 gdp.2007
## 4
          AD
               3.6759 gdp.2008
## 5
          AE 253.9655 gdp.2005
          AE 278.9489 gdp.2006
## 6
Finally, I "widen" the data, so that each var_year is a column of its own.
```

head(dat countries wide)

```
##
     country gdp.2005 gdp.2006 gdp.2007 gdp.2008 pop.2005
                                                            pop.2006
## 1
          AD
               3.8423
                        4.0184
                                 4.0216
                                          3.6759
                                                  0.081223
                                                            0.083373
## 2
          AE 253.9655 278.9489 287.8318 297.0189
                                                  4.481976
                                                            5.171255
## 3
          ΑF
               9.7630
                      10.3052 11.7212 12.1445 24.399948 25.183615
               1.1190
                       1.2687
                                 1.3892
                                          1.3902 0.082565
## 4
          AG
                                                            0.083467
## 5
          AL
               9.2684
                        9.7718 10.3483 11.1275 3.011487
                                                            2.992547
## 6
          ΑM
               7.6678
                        8.6797
                                 9.8731 10.5544 3.014917
                                                            3.002161
     pop.2007 pop.2008 unemp.2005 unemp.2006 unemp.2007 unemp.2008
##
     0.084878 0.085616
                                 NA
                                            NA
                                                       NA
## 1
                                                                  NA
## 2 6.010100 6.900142
                                3.1
                                           3.3
                                                      3.4
                                                                 4.0
## 3 25.877544 26.528741
                                8.5
                                           8.8
                                                      8.4
                                                                 8.9
## 4 0.084397 0.085350
                                 NA
                                            NA
                                                       NA
                                                                  NA
## 5 2.970017
               2.947314
                               12.5
                                          12.4
                                                     13.5
                                                                13.0
## 6 2.988117
               2.975029
                               27.8
                                          28.6
                                                     28.4
                                                                16.4
```

Now we have some ugly data. I save the output to upload to my website.

```
write_csv(dat_countries_wide, path = "untidy-data.csv")
```

And here's how I made the second country-year dataset used in the merging section. The country dataset with latitudes and longitudes is from https://developers.google.com/public-data/docs/canonical/countries_csv.

```
dat_2 <-
WDI(country = "all",
    indicator = "FP.CPI.TOTL.ZG",
    start = 2004,
    end = 2008) %>%
as_data_frame() %>%
select(country = iso2c,
        year,
        inflation = FP.CPI.TOTL.ZG) %>%
mutate(year = as.integer(year)) %>%
filter(country %in% clean_data$country) %>%
arrange(country, year)
write_csv(dat_2, path = "inflation.csv")
```

Chapter 4

Data Visualization

Visualization is most important at the very beginning and the very end of the data analysis process. In the beginning, when you've just gotten your data together, visualization is perhaps the easiest tool to explore each variable and learn about the relationships among them. And when your analysis is almost complete, you will (usually) use visualizations to communicate your findings to your audience.

We only have time to scratch the surface of data visualization. This chapter will cover the plotting techniques I find most useful for exploratory and descriptive data analysis. We will talk about graphical techniques for presenting the results of regression analyses later in the class—once we've, you know, learned something about regression.

4.1 Basic Plots

We will use the **ggplot2** package, which is part of—I'm as tired of it as you are—the **tidyverse**.

```
library("tidyverse")
```

For the examples today, we'll be using a dataset with statistics about the fifty U.S. states in 1977,¹ which is posted on my website.

```
state_data <- read_csv("http://bkenkel.com/data/state-data.csv")
state_data</pre>
```

```
## # A tibble: 50 × 12
##
          State Abbrev Region Population Income Illiteracy LifeExp Murder
##
           <chr>
                  <chr>
                          <chr>>
                                      <dbl>
                                              <dbl>
                                                          <dbl>
                                                                  <dbl>
                                                                          <dbl>
## 1
        Alabama
                      AL
                          South
                                       3615
                                               3624
                                                            2.1
                                                                  69.05
                                                                           15.1
## 2
                                        365
         Alaska
                      AK
                           West
                                               6315
                                                            1.5
                                                                  69.31
                                                                           11.3
```

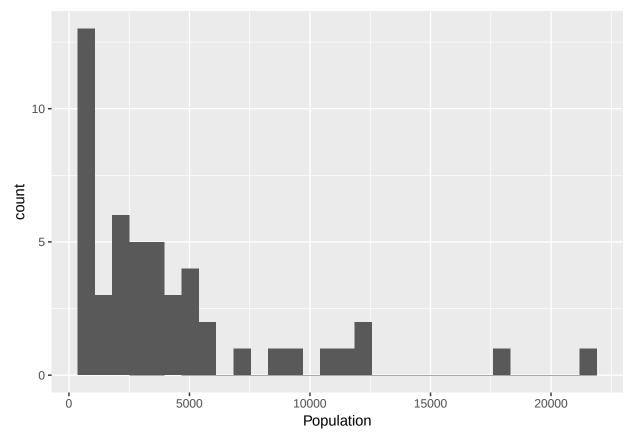
¹Why 1977? Because it was easily available. See the appendix to this chapter.

```
## 3
        Arizona
                     AZ
                                      2212
                                              4530
                                                           1.8
                                                                 70.55
                                                                           7.8
                          West
## 4
       Arkansas
                     AR
                         South
                                      2110
                                              3378
                                                           1.9
                                                                 70.66
                                                                          10.1
## 5 California
                     CA
                          West
                                     21198
                                              5114
                                                           1.1
                                                                 71.71
                                                                          10.3
    ... with 45 more rows, and 4 more variables: HSGrad <dbl>, Frost <dbl>,
       Area <dbl>, IncomeGroup <chr>>
## #
```

When I obtain data, I start by looking at the univariate distribution of each variable via a histogram. The following code creates a histogram in ggplot.

```
ggplot(state_data, aes(x = Population)) +
  geom_histogram()
```

`stat_bin()` using `bins = 30`. Pick better value with `binwidth`.



Let's walk through the syntax there. In the first line, we call <code>ggplot()</code>, specifying the data frame to draw from, then in the <code>aes()</code> command (which stands for "aesthetic") we specify the variable to plot. If this were a bivariate analysis, here we would have also specified a <code>y</code> variable to put on the y-axis. If we had just stopped there, we would have a sad, empty plot. The <code>+</code> symbol indicates that we'll be adding something to the plot. <code>geom_histogram()</code> is the command to overlay a histogram.

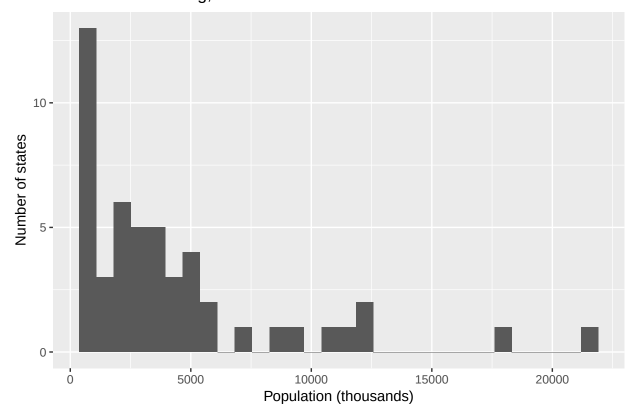
We'll only be looking at a few of the ggplot commands today. I recommend taking a look at the online package documentation at http://docs.ggplot2.org to see all of the many features available.

When you're just making graphs for yourself to explore the data, you don't need to worry about things like axis labels as long as you can comprehend what's going on. But when you prepare graphs for others to read (including those of us grading your problem sets!) you need to include an informative title and axis labels. To that end, use the xlab(), ylab(), and ggtitle() commands.

```
ggplot(state_data, aes(x = Population)) +
  geom_histogram() +
  xlab("Population (thousands)") +
  ylab("Number of states") +
  ggtitle("Some states are big, but most are small")
```

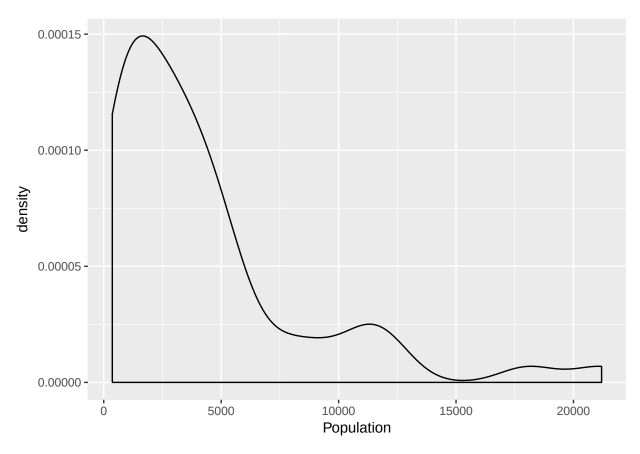
`stat_bin()` using `bins = 30`. Pick better value with `binwidth`.

Some states are big, but most are small



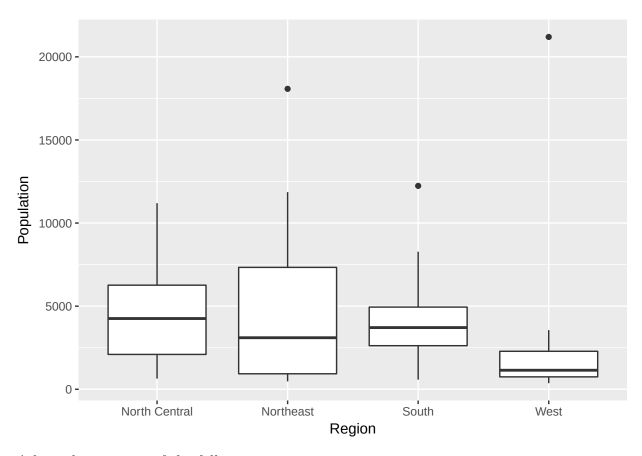
The density plot is a close relative of the histogram. It provides a smooth estimate of the probability density function of the data. Accordingly, the area under the density plot integrates to one. Depending on your purposes, this can make the y-axis of a density plot easier or (usually) harder to interpret than the count given by a histogram.

```
ggplot(state_data, aes(x = Population)) +
  geom_density()
```



The box plot is a common way to look at the distribution of a continuous variable across different levels of a categorical variable.

```
ggplot(state_data, aes(x = Region, y = Population)) +
  geom_boxplot()
```

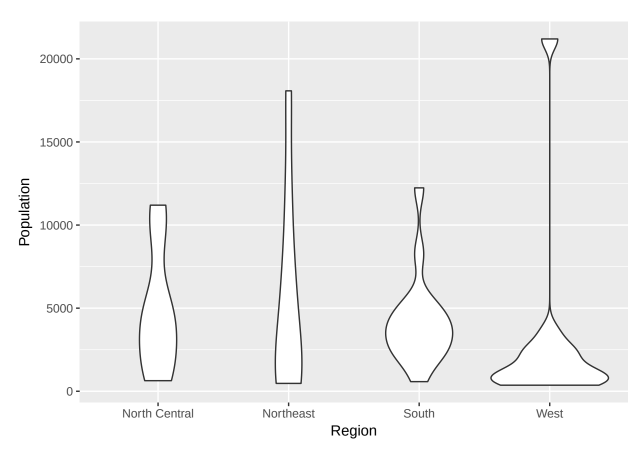


A box plot consists of the following components:

- Center line: median of the data
- Bottom of box: 25th percentile
- Top of box: 75th percentile
- \bullet Lower "whisker": range of observations no more than 1.5 IQR (height of box) below the 25th percentile
- Upper "whisker": range of observations no more than 1.5 IQR above the 75th percentile
- Plotted points: any data lying outside the whiskers

If you want to skip the summary and plot the full distribution of a variable across categories, you can use a violin plot.

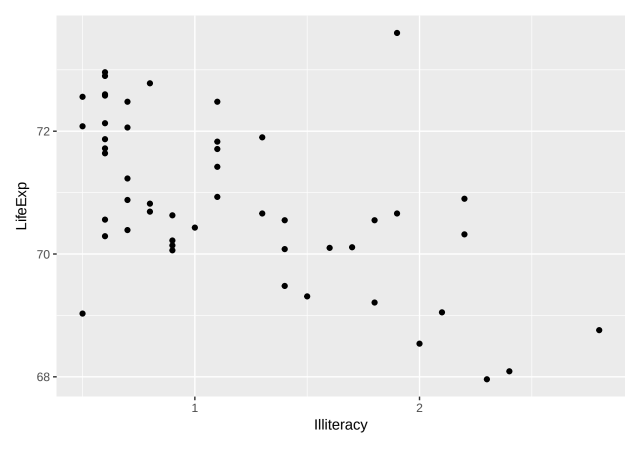
```
ggplot(state_data, aes(x = Region, y = Population)) +
  geom_violin()
```



Technically, violin plots convey more information than box plots since they show the full distribution. However, readers aren't as likely to be familiar with a violin plot. It's harder to spot immediately where the median is (though you could add that to the plot if you wanted). Plus, violin plots look goofy with outliers—see the "West" column above—whereas box plots handle them easily.

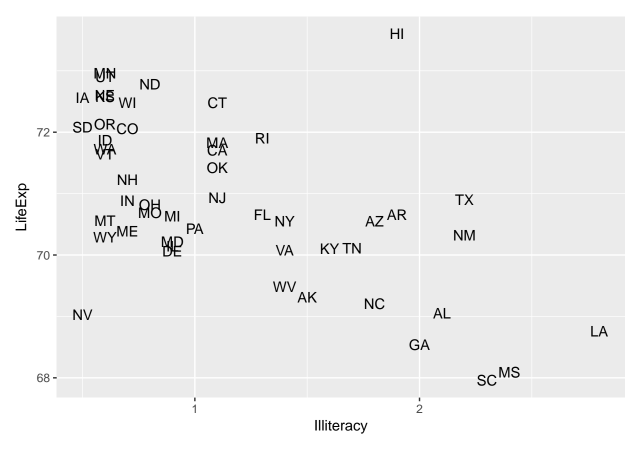
For visualizing relationships between continuous variables, nothing beats the scatterplot.

```
ggplot(state_data, aes(x = Illiteracy, y = LifeExp)) +
  geom_point()
```



When you're plotting states or countries, a hip thing to do is plot abbreviated names instead of points. To do that, you can use <code>geom_text()</code> instead of <code>geom_point()</code>, supplying an additional aesthetic argument telling ggplot where to draw the labels from.

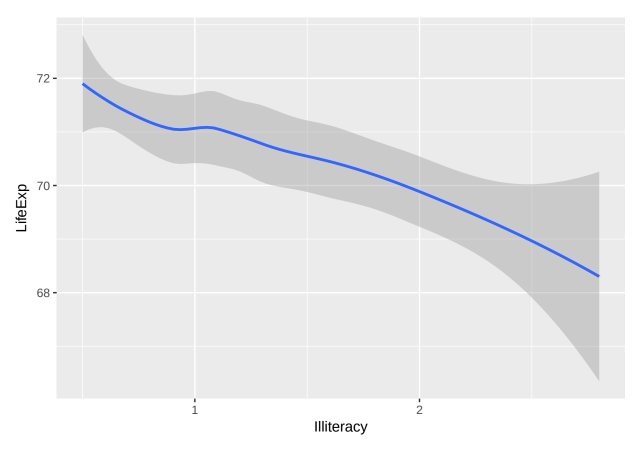
```
ggplot(state_data, aes(x = Illiteracy, y = LifeExp)) +
geom_text(aes(label = Abbrev))
```



Maybe it's overwhelming to look at all that raw data and you just want a summary. For example, maybe you want an estimate of expected LifeExp for each value of Illiteracy. This is called the *conditional expectation* and will be the subject of much of the rest of the course. For now, just now that you can calculate a smoothed conditional expectation via geom_smooth().

```
ggplot(state_data, aes(x = Illiteracy, y = LifeExp)) +
  geom_smooth()
```

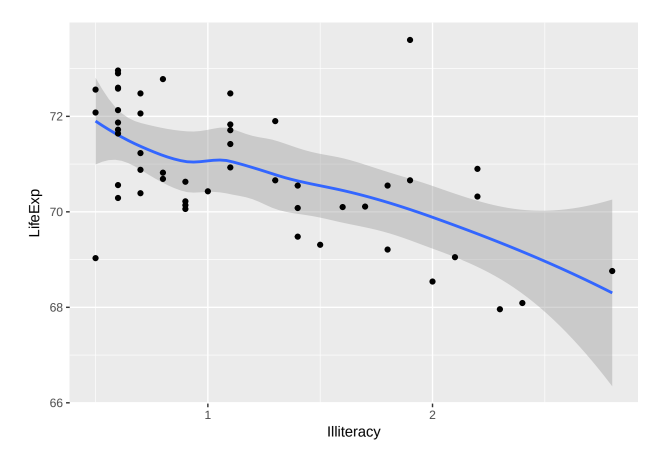
`geom_smooth()` using method = 'loess'



And if you're the kind of overachiever who likes to have the raw data and the summary, you can do it. Just add them both to the ggplot() call.

```
ggplot(state_data, aes(x = Illiteracy, y = LifeExp)) +
  geom_smooth() +
  geom_point()
```

```
## `geom_smooth()` using method = 'loess'
```



4.2 Saving Plots

When you're writing in R Markdown, the plots go straight into your document without much fuss. Odds are, your dissertation will contain plots but won't be written in R Markdown, which means you'll need to learn how to save them.

It's pretty simple:

- 1. Assign your ggplot() call to a variable.
- 2. Pass that variable to the ggsave() function.

If you want plot types other than PDF, just set a different extension. See ?ggsave for the possibilities.

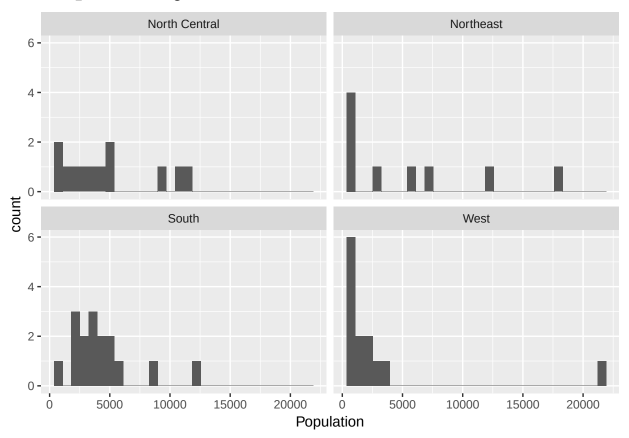
4.3 Faceting

Suppose you want to split the data into subgroups, as defined by some variable in the data (e.g., the region states are in), and make the same plot for each subgroup. ggplot's *faceting* functions, facet_wrap() and facet_grid(), make this easy.

To split up plots according to a single grouping variable, use facet_wrap(). This uses R's formula syntax, defined by the tilde ~, which you'll become well acquainted with once we start running regressions.

```
ggplot(state_data, aes(x = Population)) +
  geom_histogram() +
  facet_wrap(~ Region)
```

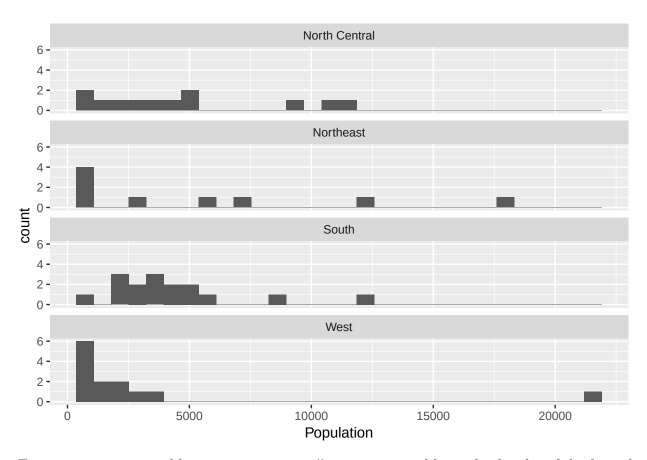
`stat_bin()` using `bins = 30`. Pick better value with `binwidth`.



If you don't like the default arrangement, use the ncol argument.

```
ggplot(state_data, aes(x = Population)) +
  geom_histogram() +
  facet_wrap(~ Region, ncol = 1)
```

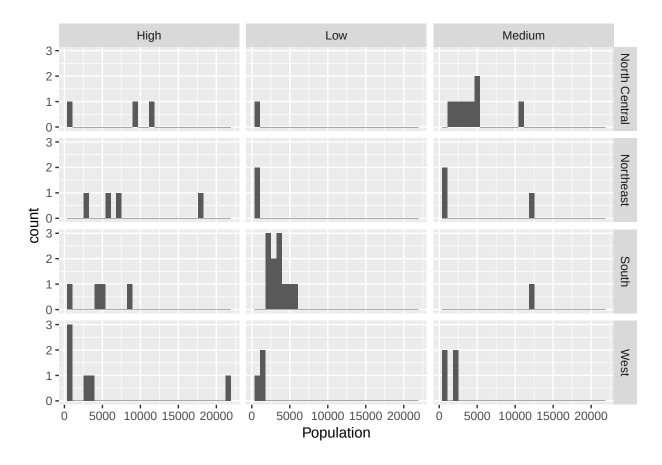
`stat_bin()` using `bins = 30`. Pick better value with `binwidth`.



For two grouping variables, use facet_grid(), putting variables on both sides of the formula.

```
ggplot(state_data, aes(x = Population)) +
  geom_histogram() +
  facet_grid(Region ~ IncomeGroup)
```

`stat_bin()` using `bins = 30`. Pick better value with `binwidth`.

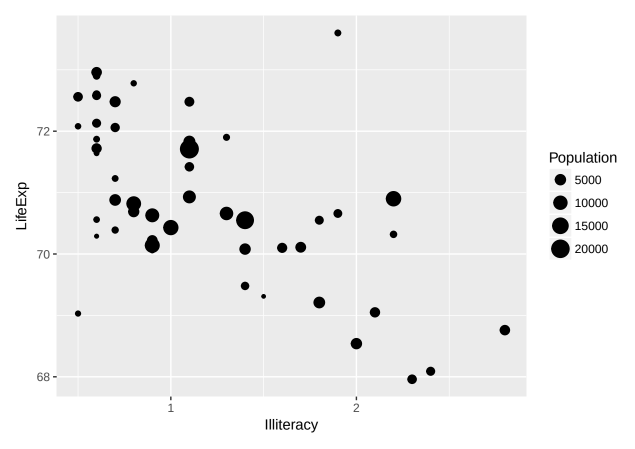


4.4 Aesthetics

Faceting is one way to incorporate information about additional variables into what would otherwise be a plot of just one or two variables. Aesthetics—which alter the appearance of particular plot features depending on the value of a variable—provide another way to do that.

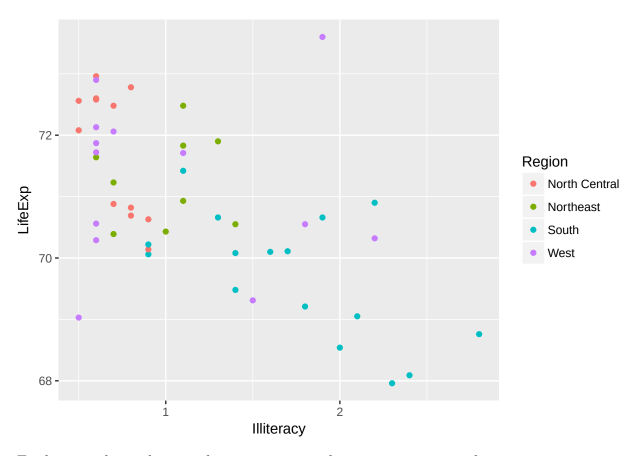
For example, when visualizing the relationship between statewide illiteracy and life expectancy, you might want larger states to get more visual weight. You can set the size aesthetic of the point geometry to vary according to the state's population.

```
ggplot(state_data, aes(x = Illiteracy, y = LifeExp)) +
  geom_point(aes(size = Population))
```



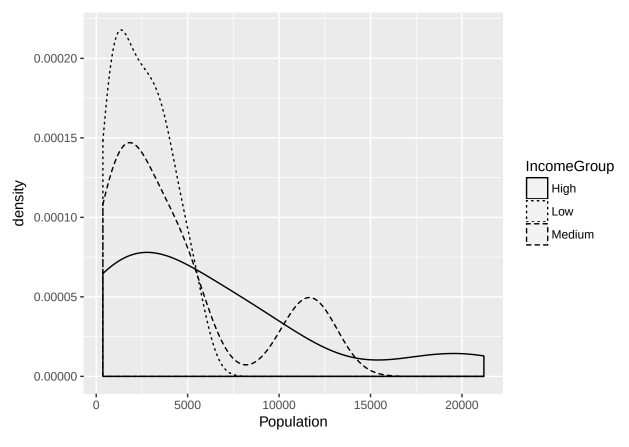
The **ggplot2** documentation lists the available aesthetics for each function. Another popular one is **colour**, which is great for on-screen display but not so much for the printed page. (And terrible for the colorblind!)

```
ggplot(state_data, aes(x = Illiteracy, y = LifeExp)) +
  geom_point(aes(colour = Region))
```



For line graphs or density plots, you can set the linetype to vary by category.

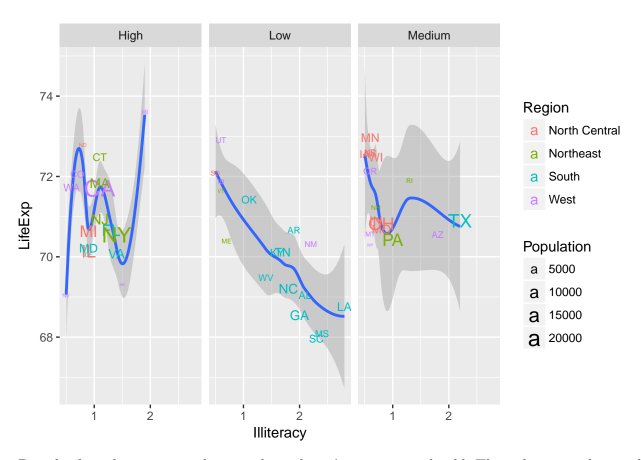
```
ggplot(state_data, aes(x = Population)) +
  geom_density(aes(linetype = IncomeGroup))
```



(I always find these incomprehensible with more than two lines, but maybe that's just me.) You can use multiple aesthetics together, and you can even combine aesthetics with faceting, as in the following example.

```
ggplot(state_data, aes(x = Illiteracy, y = LifeExp)) +
  geom_smooth() +
  geom_text(aes(label = Abbrev, colour = Region, size = Population)) +
  facet_wrap(~ IncomeGroup)
```

`geom_smooth()` using method = 'loess'



But the fact that you can do something doesn't mean you should. That plot is so cluttered that it's hard to extract the relevant information from it. Data visualizations should communicate a clear message to viewers without overwhelming them. To do this well takes practice, patience, and maybe even a bit of taste.

4.5 Appendix: Creating the Example Data

The example data comes from data on U.S. states in 1977 that are included with base R. See ?state.

Chapter 5

Bivariate Regression

The goal of empirical social science is usually to learn about the relationships between variables in the social world. Our goals might be descriptive: were college graduates more likely to vote for Clinton in 2016? Or causal: does receiving more education make a person more liberal on average? Or predictive: what kinds of voters should Democrats target in 2020 to have the best chance of victory?

The linear model is one of the simplest ways to model relationships between variables. Ordinary least squares regression is one of the easiest and (often) best ways to estimate the parameters of the linear model. Consequently, a linear model estimated by OLS is the starting point for many analyses. We will start with the simplest case: regression on a single covariate.

5.1 Probability Refresher

Let Y be a random variable that takes values in the finite set \mathcal{Y} according to the probability mass function $f_Y : \mathcal{Y} \to [0,1]$. The *expected value* (aka *expectation*) of Y is the weighted average of each value in \mathcal{Y} , where the weights are the corresponding probabilities:

$$E[Y] = \sum_{y \in \mathcal{Y}} y \, f_Y(y); \tag{5.1}$$

For a continuous random variable Y on \mathbb{R} with probability density function f_Y , the expected value is the analogous integral:

$$E[Y] = \int y f_Y(y) dy. \tag{5.2}$$

Now suppose (X,Y) is a pair of discrete random variables drawn according to the joint mass

function f_{XY} on $\mathcal{X} \times \mathcal{Y}$, with respective marginal mass functions f_X and f_Y .¹ Recall the formula for conditional probability,

$$\Pr(Y = y \mid X = x) = \frac{\Pr(X = x, Y = y)}{\Pr(X = x)} = \frac{f_{XY}(x, y)}{f_X(x)}.$$
 (5.3)

For each $x \in \mathcal{X}$, we have the conditional mass function

$$f_{Y|X}(y \mid x) = \frac{f_{XY}(x, y)}{f_{X}(x)}$$
 (5.4)

and corresponding conditional expectation

$$E[Y|X = x] = \sum_{y \in \mathcal{Y}} y \, f_{Y|X}(y \mid x). \tag{5.5}$$

For continuous random variables, the conditional expectation is

$$E[Y|X = x] = \int y \, f_{Y|X}(y|x) \, dy, \tag{5.6}$$

where $f_{Y|X}$ is the conditional density function.

The *variance* of a random variable Y is

$$V[Y] = E[(Y - E[Y])^{2}]. (5.7)$$

Given a sample Y_1, \ldots, Y_N of observations of Y, we usually estimate V[Y] with the sample variance

$$S_Y^2 = \frac{1}{N-1} \sum_n (Y_n - \bar{Y})^2, \tag{5.8}$$

where \bar{Y} is the sample mean and \sum_{n} denotes summation from n=1 to N.

Similarly (in fact a generalization of the above), the *covariance* between random variables X and Y is

$$Cov[X, Y] = E[(X - E[X])(Y - E[Y])],$$

which we estimate with the sample covariance

$$S_{XY} = \frac{1}{N-1} \sum_{n} (X_n - \bar{X})(Y_n - \bar{Y}). \tag{5.9}$$

The marginal mass function, if you don't recall, is $f_X(x) = \sum_{y \in \mathcal{Y}} f_{XY}(x,y)$. In the continuous case, the marginal density function is $f_X(x) = \int f_{XY}(x,y) \, dy$.

A fun fact about the sample covariance is that

$$S_{XY} = \frac{1}{N-1} \sum_{n} (X_n - \bar{X})(Y_n - \bar{Y})$$
 (5.10)

$$= \frac{1}{N-1} \left[\sum_{n} X_n (Y_n - \bar{Y}) + \sum_{n} \bar{X} (Y_n - \bar{Y}) \right]$$
 (5.11)

$$= \frac{1}{N-1} \left[\sum_{n} X_n (Y_n - \bar{Y}) + \bar{X} \sum_{n} (Y_n - \bar{Y}) \right]$$
 (5.12)

$$= \frac{1}{N-1} \sum_{n} X_n (Y_n - \bar{Y}). \tag{5.13}$$

If we had split up the second term instead of the first, we would see that

$$S_{XY} = \frac{1}{N-1} \sum_{n} Y_n (X_n - \bar{X})$$

as well.

Since the (sample) variance is a special case of the (sample) covariance, by the same token we have

$$S_Y^2 = \frac{1}{N-1} \sum_n Y_n (Y_n - \bar{Y}). \tag{5.14}$$

5.2 The Linear Model

Suppose we observe a sequence of N draws from f_{XY} , denoted $(X_1, Y_1), (X_2, Y_2), \ldots, (X_N, Y_N)$, or $\{(X_n, Y_n)\}_{n=1}^N$ for short. What can we learn about the relationship between X and Y from this sample of data?

If we were really ambitious, we could try to estimate the shape of the full joint distribution, f_{XY} . The joint distribution encodes everything there is to know about the relationship between the two variables, so it would be pretty useful to know. But except in the most trivial cases, it would be infeasible to estimate f_{XY} precisely. If X or Y can take on more than a few values, estimating the joint distribution would require an amount of data that we're unlikely to have.²

The first way we simplify our estimation task is to set our sights lower. Let Y be the response or the dependent variable—i.e., the thing we want to explain. We call X the covariate or the independent variable. Instead of estimating the full joint distribution, we're just going to try to learn the conditional expectation, E[Y | X]. In other words, for each potential value

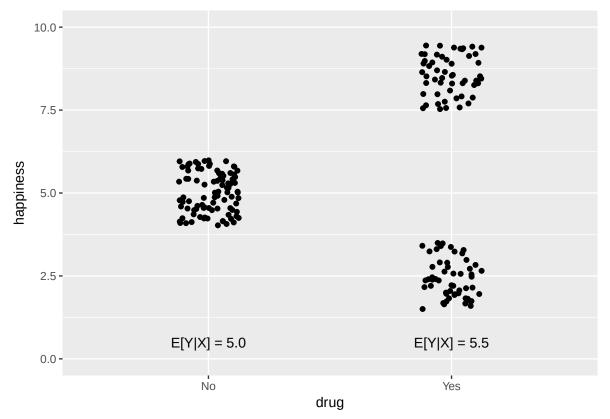
²This problem only gets worse as we move from bivariate into multivariate analysis, a phenomenon called the *curse of dimensionality*.

of the covariate, what is the expected value of the response? This will allow us to answer questions like whether greater values of X are associated with greater values of Y.

Two important things about the estimation of conditional expectations before we go any further.

- 1. Statements about conditional expectations are not causal. If Y is rain and X is umbrella sales, we know E[Y|X] increases with X, but that doesn't mean umbrella sales make it rain.
 - We will spend some time in the latter part of the course on how to move from conditional expectations to causality. Then, in Stat III, you will learn about causal inference in excruciating detail.
- 2. The conditional expectation doesn't give you everything you'd want to know about the relationship between variables.

As a hypothetical example, suppose I told you that taking a particular drug made people happier on average. In other words, $E[\text{Happiness} \mid \text{Drug}] > E[\text{Happiness} \mid \text{No Drug}]$. Sounds great! Then imagine the dose-response graph looked like this:



The fact that expected happiness rises by half a point doesn't quite tell the whole story. In spite of these caveats, conditional expectation is a really useful tool for summarizing the relationship between variables.

If X takes on sufficiently few values (and we have enough data), we don't need to model the

conditional expectation function. We can just directly estimate E[Y|X=x] for each $x \in \mathcal{X}$. The graph above, where there are just two values of X, is one example.

But if X is continuous, or even if it is discrete with many values, estimating E[Y|X] for each distinct value is infeasible. In this case, we need to model the relationship. The very simplest choice—and thus the default for social scientists—is to model the conditional expectation of Y as a linear function of X:

$$E[Y \mid X] = \alpha + \beta X. \tag{5.15}$$

In this formulation, α and β are the parameters to be estimated from sample data. We call α and β "coefficients," with α the "intercept" and β the "slope." Regardless of how many different values X might take on, we only need to estimate two parameters of the linear model.

Exercise your judgment before using a linear model. Ask yourself, is a linear conditional expectation function at least minimally plausible? Not perfect—just a reasonable approximation. If X is years of education and Y is annual income, the answer is probably yes (depending on the population!). But if X is hour of the day (0-24) and Y is the amount of traffic on I-65, probably not.

To obtain the linear conditional expectation, we usually assume the following model of the response variable:

$$Y_n = \alpha + \beta X_n + \epsilon_n, \tag{5.16}$$

where ϵ_n is "white noise" error with the property

$$E[\epsilon_n \mid X_1, \dots X_N] = 0. \tag{5.17}$$

You can think of ϵ_n as the summation of everything besides the covariate X_n that affects the response Y_n . The assumption that $E[\epsilon_n | X_1, \dots, X_N] = 0$ implies that these external factors are uncorrelated with the covariate. This is not a trivial technical condition that you can ignore—it is a substantive statement about the variables in your model. It requires justification, and it is difficult to justify.

For now we will proceed assuming that our data satisfy the above conditions. Later in the course, we will talk about how to proceed when $E[\epsilon_n | X_1, \dots, X_N] \neq 0$, and you will learn much more about such strategies in Stat III.

5.3 Least Squares

To estimate the parameters of the linear model, we will rely on a mathematically convenient method called *least squares*. We will see that this method not only is convenient, but also

has nice statistical properties.

Given a parameter estimate $(\hat{\alpha}, \hat{\beta})$, define the *residual* of the *n*'th observation as the difference between the true and predicted values:

$$e_n(\hat{\alpha}, \hat{\beta}) = Y_n - \hat{\alpha} - \hat{\beta}X_n. \tag{5.18}$$

The residual is directional. The residual is positive when the regression line falls below the observation, and vice versa when it is negative.

We would like the regression line to lie close to the data—i.e., for the residuals to be small in magnitude. "Close" can mean many things, so we need to be a bit more specific to derive an estimator. The usual one, *ordinary least squares*, is chosen to minimize the sum of squared errors,

$$SSE(\hat{\alpha}, \hat{\beta}) = \sum_{n} e_n(\hat{\alpha}, \hat{\beta})^2.$$

(Throughout the rest of this chapter, I write \sum_n as shorthand for $\sum_{n=1}^N$.) When we focus on squared error, we penalize a positive residual the same as a negative residual of the same size. Moreover, we penalize one big residual proportionally more than a few small ones.

It is important to keep the linear model and ordinary least squares distinct in your mind. The linear model is a model of the data. Ordinary least squares is one estimator—one among many—of the parameters of the linear model. Assuming a linear model does not commit you to estimate it with OLS if you think another estimator is more appropriate. And using OLS does not necessarily commit you to the linear model, as we will discuss when we get to multiple regression.

To derive the OLS estimator, we will derive the conditions for minimization of the sum of squared errors. The SSE is a quadratic and therefore continuously differentiable function of the estimands, $\hat{\alpha}$ and $\hat{\beta}$. You will remember from calculus that, at any extreme point of a continuous function, all its partial derivatives equal zero. To derive necessary conditions for minimization,³ we can take the derivatives of the SSE and set them to equal zero.

The derivative with respect to the intercept is

$$\frac{\partial \operatorname{SSE}(\hat{\alpha}, \hat{\beta})}{\partial \hat{\alpha}} = -2\sum_{n} (Y_n - \hat{\alpha} - \hat{\beta}X_n).$$

Setting this to equal zero gives

$$\hat{\alpha} = \frac{1}{N} \sum_{n} (Y_n - \hat{\beta} X_n) = \bar{Y} - \hat{\beta} \bar{X}.$$

This gives us one important property of OLS: the regression line estimated by OLS always passes through (\bar{X}, \bar{Y}) .

³In fact, since the SSE function is strictly convex, these conditions are sufficient for global minimization.

The derivative with respect to the slope is

$$\frac{\partial \operatorname{SSE}(\hat{\alpha}, \hat{\beta})}{\partial \hat{\beta}} = -2 \sum_{n} X_n (Y_n - \hat{\alpha} - \hat{\beta} X_n).$$

Setting this equal to zero and substituting in the expression for $\hat{\alpha}$ we derived above gives

$$\sum_{n} X_n(Y_n - \bar{Y}) = \hat{\beta} \sum_{n} X_n(X_n - \bar{X}).$$

As long as the sample variance of X is non-zero (i.e., X is not a constant), we can divide to solve for $\hat{\beta}$:

$$\hat{\beta} = \frac{\sum_{n} X_{n} (Y_{n} - \bar{Y})}{\sum_{n} X_{n} (X_{n} - \bar{X})} = \frac{S_{XY}}{S_{X}^{2}}.$$

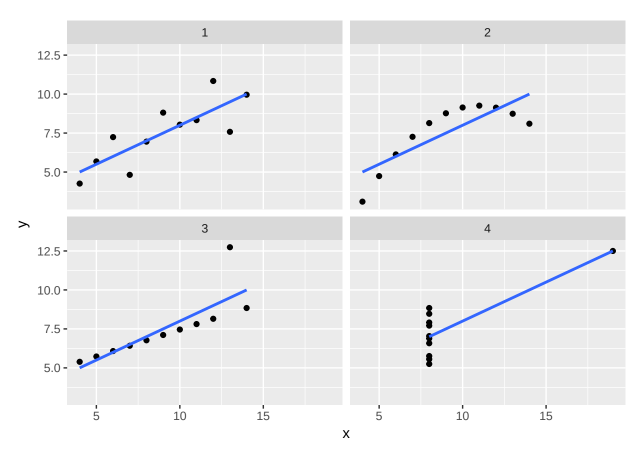
Combining these two results, we have the OLS estimators of the intercept and slope of the bivariate linear model. We write them as functions of $(X_1, \ldots, X_N, Y_1, \ldots, Y_N)$, or (X, Y) for short,⁴ to emphasize that an estimator is a statistic, which in turn is a function of sample data. We place the "OLS" subscript on them to emphasize that there are many estimators of these parameters, of which OLS is just one (good!) choice.

$$\hat{\alpha}_{\text{OLS}}(X,Y) = \bar{Y} - \frac{S_{XY}}{S_X^2} \bar{X},$$

$$\hat{\beta}_{\text{OLS}}(X,Y) = \frac{S_{XY}}{S_Y^2}.$$

Regression is a convenient way to summarize the relationship between variables, but it is a complement to—not a substitute for—graphical analysis. The statistician Francis Anscombe found that OLS yields nearly identical regression lines for all four of the datasets in the following graph:

 $^{^4}$ This is a bit of an abuse of notation, since previously I used X and Y to refer to the random variables and now I'm using them to refer to vectors of sample data. Sorry.



Unless your data all lie along a line, the regression line estimated by OLS will not predict the data perfectly. Let the *residual sum of squares* be the squared error left over by OLS,

$$RSS = SSE(\hat{\alpha}_{OLS}, \hat{\beta}_{OLS}),$$

and let the $total\ sum\ of\ squares$ be the squared error that would result from a horizontal regression line through the mean of Y,

$$TSS = SSE(\bar{Y}, 0).$$

The \mathbb{R}^2 statistic is the proportion of "variance explained" by X, calculated as

$$R^2 = 1 - \frac{\text{RSS}}{\text{TSS}}.$$

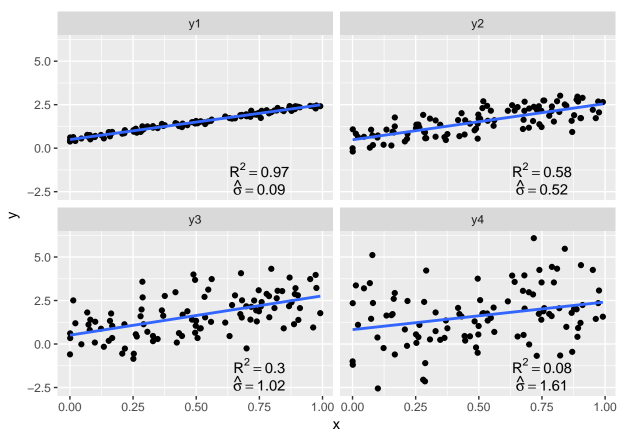
If the regression line is flat, in which case $\hat{\beta}_{OLS} = 0$ and RSS = TSS, we have $R^2 = 0$. Conversely, if the regression line fits perfectly, in which case RSS = 0, we have $R^2 = 1$.

A statistic that is often more useful than R^2 is the *residual variance*. The residual variance is (almost) the sample variance of the regression residuals, calculated as

$$\hat{\sigma}^2 = \frac{1}{N-2} \sum_n e_n(\hat{\alpha}_{\text{OLS}}, \hat{\beta}_{\text{OLS}})^2 = \frac{\text{RSS}}{N-2}$$

Since bivariate regression uses two degrees of freedom (one for the intercept, one for the slope), we divide by N-2 instead of the usual N-1. The most useful quantity is $\hat{\sigma}$,

the square root of the residual variance. $\hat{\sigma}$ is measured in the same units as Y, and it is a measure of the spread of points around the regression line. If the residuals are roughly normally distributed, then we would expect roughly 95% of the data to lie within $\pm 2\hat{\sigma}$ of the regression line.



5.4 Properties

We didn't use any fancy statistical theory to derive the OLS estimator. We just found the intercept and slope that minimize the sum of squared residuals. As it turns out, though, OLS indeed has some very nice statistical properties as an estimator of the linear model.

The first desirable property of OLS is that it is *unbiased*. Recall that an estimator $\hat{\theta}$ of the parameter θ is unbiased if $E[\hat{\theta}] = \theta$. This doesn't mean the estimator always gives us the right answer, just that on average it is not systematically biased upward or downward. In other words, if we could take many many samples and apply the estimator to each of them, the average would equal the true parameter.

We will begin by showing that the OLS estimator of the slope is unbiased; i.e., that $E[\hat{\beta}_{OLS}(X,Y)] = \beta$. At first, we'll take the conditional expectation of the slope estimator,

treating the covariates (X_1, \ldots, X_N) as fixed.

$$E[\hat{\beta}_{OLS}(X,Y) \mid X] = E\left[\frac{S_{XY}}{S_X^2} \mid X\right]$$

$$= E\left[\frac{\sum_n Y_n(X_n - \bar{X})}{\sum_n X_n(X_n - \bar{X})} \mid X\right]$$

$$= \frac{\sum_n E[Y_n \mid X](X_n - \bar{X})}{\sum_n X_n(X_n - \bar{X})}$$

$$= \frac{\sum_n (\alpha + \beta X_n)(X_n - \bar{X})}{\sum_n X_n(X_n - \bar{X})}$$

$$= \frac{\alpha \sum_n (X_n - \bar{X}) + \beta \sum_n X_n(X_n - \bar{X})}{\sum_n X_n(X_n - \bar{X})}$$

$$= \frac{\beta \sum_n X_n(X_n - \bar{X})}{\sum_n X_n(X_n - \bar{X})}$$

$$= \beta.$$

It then follows from the law of iterated expectation⁵ that

$$E[\hat{\beta}_{OLS}(X,Y)] = \beta.$$

Then, for the intercept, we have

$$E[\hat{\alpha}_{OLS}(X,Y) \mid X] = E[\bar{Y} - \hat{\beta}_{OLS}(X,Y)\bar{X} \mid X]$$

$$= E[\bar{Y} \mid X] - E[\hat{\beta}_{OLS}(X,Y) \mid X]\bar{X}$$

$$= E\left[\frac{1}{N}\sum_{n}Y_{n} \mid X\right] - \beta\bar{X}$$

$$= E\left[\frac{1}{N}\sum_{n}(\alpha + \beta X_{n} + \epsilon_{n}) \mid X\right] - \beta\bar{X}$$

$$= \frac{1}{N}\sum_{n}E[\alpha + \beta X_{n} + \epsilon_{n} \mid X] - \beta\bar{X}$$

$$= \frac{1}{N}\sum_{n}\alpha + \frac{\beta}{N}\sum_{n}X_{n} + \frac{1}{N}\sum_{n}E[\epsilon_{n} \mid X] - \beta\bar{X}$$

$$= \alpha + \beta\bar{X} - \beta\bar{X}$$

$$= \alpha.$$

As with the slope, this conditional expectation gives us the unconditional expectation we want:

$$E[\hat{\alpha}_{\text{OLS}}(X,Y)] = \alpha.$$

To sum up: as long as the crucial condition $E[\epsilon_n | X_1, \dots, X_N] = 0$ holds, then OLS is an unbiased estimator of the parameters of the linear model.

⁵For random variables A and B, $E[f(A,B)] = E_A[E_B[f(A,B) \mid A]] = E_B[E_A[f(A,B) \mid B]].$

Another important property of OLS is that it is *consistent*. Informally, this means that in sufficiently large samples, the OLS estimates $(\hat{\alpha}_{OLS}, \hat{\beta}_{OLS})$ are very likely to be close to the true parameter values (α, β) . Another way to think of consistency is that, as $N \to \infty$, the bias and variance of the OLS estimator both go to zero.⁶

Of course the bias "goes to" zero, since OLS is unbiased. The real trick to proving consistency is to show that the variance goes to zero. If you wanted to do that for the slope estimate, you'd derive an expression for

$$V[\hat{\beta}_{\text{OLS}}] = E[(\hat{\beta}_{\text{OLS}} - E[\hat{\beta}_{\text{OLS}}])^2] = E[(\hat{\beta}_{\text{OLS}} - \beta)^2]$$

and show that

$$\lim_{N \to \infty} V[\hat{\beta}_{\text{OLS}}] = 0.$$

This takes more algebra than we have time for, so I leave it as an exercise for the reader.

5.5 Appendix: Regression in R

We will be using the **tidyverse** package as always, the **car** package for the **Prestige** data, and the **broom** package for its convenient post-analysis functions.

```
library("tidyverse")
library("car")
library("broom")
```

Let's take a look at Prestige, which records basic information (including perceived prestige) for a variety of occupations.

head(Prestige)

```
##
                        education income women prestige census type
                            13.11
                                                           1113 prof
## gov.administrators
                                   12351 11.16
                                                    68.8
## general.managers
                            12.26
                                   25879 4.02
                                                           1130 prof
                                                    69.1
## accountants
                            12.77
                                    9271 15.70
                                                    63.4
                                                           1171 prof
## purchasing.officers
                                                           1175 prof
                            11.42
                                    8865
                                          9.11
                                                    56.8
## chemists
                            14.62
                                    8403 11.68
                                                    73.5
                                                           2111 prof
## physicists
                            15.64
                                   11030 5.13
                                                    77.6
                                                           2113 prof
```

Suppose we want to run a regression of prestige on education. We will use the lm() function, which stands for *linear model*. This will employ the "formula" syntax that you previously saw when faceting in ggplot. The basic syntax of a formula is response ~ covariate, where response and covariate are the names of the variables in question. In this case, with prestige (note that the variable is lowercase, while the dataset is capitalized) as the response and education as the covariate:

⁶What I am describing here is *mean square consistency*, which is stronger than the broadest definitions of consistency in statistical theory.

(Intercept)

-10.73

##

```
lm(prestige ~ education, data = Prestige)

##
## Call:
## lm(formula = prestige ~ education, data = Prestige)
##
## Coefficients:
```

education

5.36

You'll notice that didn't give us very much. If you've previously used statistical programs like Stata, you might expect a ton of output at this point. It's all there in R too, but R has a different philosophy about models. R sees the fitted model as an object in its own right—like a data frame, a function, or anything else you load or create in R. Therefore, to analyze regression results in R, you will typically save the regression results to a variable.

Like any other variable, you'll want to give your regression results meaningful names. I typically call them fit_ to indicate a fitted model, followed by some memorable description.

```
fit_educ <- lm(prestige ~ education, data = Prestige)</pre>
```

When you do this, the output doesn't get printed. To see the default output, just run the variable name, just like you would to see the content of a data frame:

```
fit_educ
```

```
##
## Call:
## lm(formula = prestige ~ education, data = Prestige)
##
## Coefficients:
## (Intercept) education
## -10.73 5.36
```

For a more detailed readout, use the summary() method:

```
summary(fit_educ)
```

```
##
## Call:
## lm(formula = prestige ~ education, data = Prestige)
##
## Residuals:
##
       Min
                     Median
                                  3Q
                                         Max
                 1Q
                      0.661
##
  -26.040
            -6.523
                               6.743
                                      18.164
##
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
##
```

This prints out a whole boatload of information, including inferential statistics that we're going to wait until later in the course to discuss how to interpret:

- The model you ran
- Basic statistics about the distribution of the residuals
- For each coefficient:
 - Parameter estimate
 - Standard error estimate
 - Test statistic for a hypothesis test of equality with zero
 - p-value associated with the test statistic
- $\hat{\sigma}$ (called the "residual standard error", a term seemingly unique to R)
- R^2 and an "adjusted" variant that accounts for the number of variables in the model
- F statistic, degrees of freedom, and associated p-value for a hypothesis test that every coefficient besides the intercept equals zero

Strangely, summary() doesn't give you the sample size. For that you must use nobs():

```
nobs(fit_educ)
```

```
## [1] 102
```

You can use a fitted model object to make predictions for new data. For example, let's make a basic data frame of education levels.

```
my_data <- data_frame(education = 8:16)
my_data</pre>
```

```
## # A tibble: 9 × 1
##
     education
##
          <int>
## 1
              8
## 2
              9
## 3
             10
## 4
             11
## 5
             12
## 6
             13
## 7
             14
## 8
             15
## 9
             16
```

To calculate the predicted level of prestige for each education level, use predict():

```
predict(fit_educ, newdata = my_data)
## 1 2 3 4 5 6 7 8 9
## 32.155 37.516 42.877 48.238 53.599 58.959 64.320 69.681 75.042
```

When using predict(), it is crucial that the newdata have the same column names as in the data used to fit the model.

You can also extract a confidence interval for each prediction:

```
predict(fit educ,
        newdata = my_data,
        interval = "confidence",
        level = 0.95)
##
        fit
               lwr
                      upr
## 1 32.155 29.615 34.695
## 2 37.516 35.393 39.639
## 3 42.877 41.024 44.730
## 4 48.238 46.441 50.034
## 5 53.599 51.627 55.571
## 6 58.959 56.632 61.287
## 7 64.320 61.525 67.116
## 8 69.681 66.353 73.010
## 9 75.042 71.142 78.942
```

One of the problems with summary() and predict() is that they return inconveniently shaped output. The output of summary() is particularly hard to deal with. The **broom** package provides three utilities to help get model output into shape. The first is tidy(), which makes a tidy data frame out of the regression coefficients and the associated inferential statistics:

```
tidy(fit_educ)
```

```
## term estimate std.error statistic p.value
## 1 (Intercept) -10.7320 3.67709 -2.9186 4.3434e-03
## 2 education 5.3609 0.33199 16.1478 1.2863e-29
```

The second is glance(), which provides a one-row data frame containing overall model characteristics (e.g., R^2 and $\hat{\sigma}$):

```
glance(fit educ)
     r.squared adj.r.squared sigma statistic
##
                                                  p.value df logLik
                                                                        AIC
                     0.72003 9.1033
                                        260.75 1.2863e-29
                                                                -369 744.01
## 1
        0.7228
                                                            2
##
        BIC deviance df.residual
## 1 751.88
                8287
                              100
```

The third is augment(), which "augments" the original data—or new data you supply, as in

7

8

9

14

15

16

64.320 1.40897 69.681 1.67763

75.042 1.96574

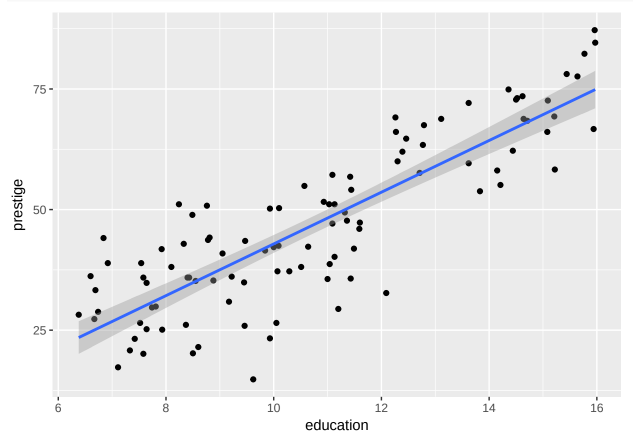
predict()—with information from the model, such as predicted values.

```
# Lots of output, so only printing first 10 rows
head(augment(fit educ), 10)
##
                .rownames prestige education .fitted .se.fit
                                                               .resid
                                                                          .hat
## 1
       gov.administrators
                              68.8
                                              59.549 1.19689
                                                              9.2509 0.017287
                                       13.11
## 2
        general.managers
                              69.1
                                              54.992 1.03332 14.1076 0.012885
                                       12.26
## 3
              accountants
                              63.4
                                       12.77
                                              57.726 1.12584
                                                              5.6736 0.015295
## 4
     purchasing.officers
                              56.8
                                       11.42 50.489 0.92936 6.3108 0.010422
## 5
                 chemists
                              73.5
                                       14.62
                                              67.644 1.57269
                                                              5.8559 0.029846
## 6
               physicists
                              77.6
                                       15.64 73.112 1.86034
                                                              4.4879 0.041763
## 7
                              72.6
               biologists
                                       15.09
                                              70.164 1.70291
                                                              2.4363 0.034993
               architects
                              78.1
                                       15.44 72.040 1.80254 6.0600 0.039208
## 8
## 9
          civil.engineers
                              73.1
                                       14.52 67.108 1.54561 5.9920 0.028827
## 10
        mining.engineers
                              68.8
                                       14.64 67.751 1.57814 1.0487 0.030053
##
                .cooksd .std.resid
      .sigma
## 1
      9.1010 0.00924267
                           1.02511
## 2 9.0372 0.01587881
                           1.55981
## 3 9.1311 0.00306360
                           0.62807
## 4 9.1269 0.00255745
                           0.69688
      9.1296 0.00656112
                           0.65310
## 6 9.1375 0.00552702
                           0.50362
## 7
      9.1458 0.00134578
                           0.27244
## 8 9.1280 0.00941104
                           0.67914
## 9 9.1287 0.00662109
                           0.66793
## 10 9.1485 0.00021198
                           0.11697
augment(fit educ,
        newdata = my data)
##
     education .fitted .se.fit
## 1
                32.155 1.28013
## 2
             9
               37.516 1.07023
## 3
            10
               42.877 0.93407
## 4
               48.238 0.90555
            11
## 5
            12
               53.599 0.99397
## 6
            13
               58.959 1.17319
```

Notice that you get back more information for the data used to fit the model than for newly supplied data. The most important is .fitted, the predicted value. See ?augment.lm for what all the various output represents.

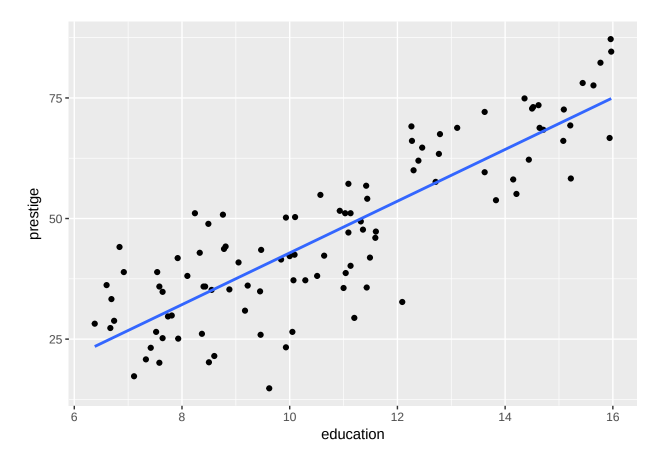
One last note on plotting regression lines with ggplot. Use geom smooth(method = "lm").

```
ggplot(Prestige, aes(x = education, y = prestige)) +
  geom_point() +
  geom_smooth(method = "lm")
```



To get rid of the confidence interval:

```
ggplot(Prestige, aes(x = education, y = prestige)) +
  geom_point() +
  geom_smooth(method = "lm", se = FALSE)
```



Chapter 6

Matrix Algebra: A Crash Course

Some material in this chapter is adapted from notes Hye Young You wrote for the math boot camp for the political science PhD program at Vanderbilt.

Matrix algebra is an essential tool for understanding multivariate statistics. You are probably already familiar with matrices, at least informally. The data representations we have worked with so far—each row an observation, each column a variable—are formatted like matrices.

An introductory treatment of matrix algebra is a semester-long college course. We don't have that long, or even half that long. This chapter gives you the *bare minimum* you need to understand to get up and running with the matrix algebra we need for OLS with multiple covariates. If you want to use advanced statistical methods in your research and haven't previously taken a matrix algebra or linear algebra course, I recommend taking some time this summer to catch up. For example, MIT has its undergraduate linear algebra course available online, including video lectures.

6.1 Vector Operations

A *vector* is an ordered array. To denote a vector v of k elements, we write $\mathbf{v} = (v_1, v_2, \dots, v_k)$, or sometimes

$$\mathbf{v} = \begin{pmatrix} v_1 \\ v_2 \\ \vdots \\ v_k \end{pmatrix}.$$

Notice the convention of using a lowercase bold letter to denote a vector. We will usually be dealing with vectors of real numbers. To denote the fact that \mathbf{v} is a vector of k real numbers, we write $\mathbf{v} \in \mathbb{R}^k$.

A vector can be multiplied by a scalar $c \in \mathbb{R}$, producing what you would expect:

$$c\mathbf{v} = \begin{pmatrix} cv_1 \\ cv_2 \\ \vdots \\ cv_k \end{pmatrix}$$

You can also add and subtract two vectors of the same length.¹

$$\mathbf{u} + \mathbf{v} = \begin{pmatrix} u_1 + v_1 \\ u_2 + v_2 \\ \vdots \\ u_k + v_k \end{pmatrix},$$

$$\mathbf{u} - \mathbf{v} = \begin{pmatrix} u_1 - v_1 \\ u_2 - v_2 \\ \vdots \\ u_k - v_k \end{pmatrix}.$$

A special vector is the *zero vector*, which contains—you guessed it—all zeroes. We write $\mathbf{0}_k$ to denote the zero vector of length k. When the length of the zero vector is clear from the context, we may just write $\mathbf{0}$.

The last important vector operation is the *dot product*. The dot product of \mathbf{u} and \mathbf{v} , written $\mathbf{u} \cdot \mathbf{v}$, is the sum of the products of the entries:

$$\mathbf{u} \cdot \mathbf{v} = u_1 v_1 + u_2 v_2 + \dots + u_k v_k = \sum_{m=1}^k u_m v_m.$$

An important concept for regression analysis is the linear independence of a collection of vectors. Let $\mathbf{v}_1, \dots, \mathbf{v}_J$ be a collection of J vectors, each of length k. We call \mathbf{u} a linear combination of $\mathbf{v}_1, \dots, \mathbf{v}_J$ if there exist real numbers c_1, \dots, c_J such that

$$\mathbf{u} = c_1 \mathbf{v}_1 + \dots + c_J \mathbf{v}_J = \sum_{j=1}^J c_j \mathbf{v}_j.$$

A collection of vectors is *linearly independent* if the only solution to

$$c_1\mathbf{v}_1 + \dots + c_J\mathbf{v}_J = \mathbf{0}$$

is $c_1 = 0, \dots, c_J = 0$. Otherwise, we call the vectors *linearly dependent*. Some fun facts about linear independence:

 $^{^{1}}$ R will let you add and subtract vectors of different lengths, via a technique called "recycling". For example c(1, 0) + c(1, 2, 3, 4) will produce c(2, 2, 4, 4). This is kosher in R, but not in mathematical derivations.

- If any vector in $\mathbf{v}_1, \dots, \mathbf{v}_J$ is a linear combination of the others, then these vectors are linearly dependent.
- A collection of J vectors of length k cannot be linearly independent if J > k. In other words, given vectors of length k, the most that can be linearly independent of each other is k.
- If any $\mathbf{v}_i = \mathbf{0}$, then $\mathbf{v}_1, \dots, \mathbf{v}_J$ are linearly dependent. (Why?)

Examples:

$$\mathbf{v}_{1} = \begin{pmatrix} 1\\1\\1 \end{pmatrix}, \mathbf{v}_{2} = \begin{pmatrix} 0\\1\\0 \end{pmatrix}, \mathbf{v}_{3} = \begin{pmatrix} 0\\0\\1 \end{pmatrix};$$

$$\mathbf{v}_{1} = \begin{pmatrix} 1\\0\\0 \end{pmatrix}, \mathbf{v}_{2} = \begin{pmatrix} 14\\12\\0 \end{pmatrix}, \mathbf{v}_{3} = \begin{pmatrix} 0\\-1\\0 \end{pmatrix};$$

$$\mathbf{v}_{1} = \begin{pmatrix} 1\\2\\3 \end{pmatrix}, \mathbf{v}_{2} = \begin{pmatrix} 1\\4\\9 \end{pmatrix}, \mathbf{v}_{3} = \begin{pmatrix} 1\\8\\27 \end{pmatrix}.$$

6.2 Matrix Operations

A matrix is a two-dimensional array of numbers, with entries in rows and columns. We call a matrix with n rows and m columns an $n \times m$ matrix. For example, the following is a 2×3 matrix:

$$\mathbf{A} = \begin{bmatrix} 99 & 73 & 2 \\ 13 & 40 & 41 \end{bmatrix}$$

Notice the convention of using an uppercase bold letter to denote a matrix. Given a matrix \mathbf{A} , we usually write a_{ij} to denote the entry in the *i*'th row and *j*'th column. In the above example, we have $a_{13} = 2$.

You can think of a vector $\mathbf{v} \in \mathbb{R}^k$ as a $1 \times k$ row matrix or as a $k \times 1$ column matrix. Throughout this book, I will treat vectors as column matrices unless otherwise noted.

Like vectors, matrices can be multipled by a scalar $c \in \mathbb{R}$.

$$c\mathbf{A} = \begin{bmatrix} ca_{11} & ca_{12} & \cdots & ca_{1m} \\ ca_{21} & ca_{22} & \cdots & ca_{2m} \\ \vdots & \vdots & \ddots & \vdots \\ ca_{n1} & ca_{n2} & \cdots & ca_{nm} \end{bmatrix}$$

Matrices of the same dimension (i.e., both with the same number of rows n and columns m)

can be added ...

$$\mathbf{A} + \mathbf{B} = \begin{bmatrix} a_{11} + b_{11} & a_{12} + b_{12} & \cdots & a_{1m} + b_{1m} \\ a_{21} + b_{21} & a_{22} + b_{22} & \cdots & a_{2m} + b_{2m} \\ \vdots & \vdots & \ddots & \vdots \\ a_{n1} + b_{n1} & a_{n2} + b_{n2} & \cdots & a_{nm} + b_{nm} \end{bmatrix}$$

 \dots and subtracted \dots

$$\mathbf{A} - \mathbf{B} = \begin{bmatrix} a_{11} - b_{11} & a_{12} - b_{12} & \cdots & a_{1m} - b_{1m} \\ a_{21} - b_{21} & a_{22} - b_{22} & \cdots & a_{2m} - b_{2m} \\ \vdots & \vdots & \ddots & \vdots \\ a_{n1} - b_{n1} & a_{n2} - b_{n2} & \cdots & a_{nm} - b_{nm} \end{bmatrix}$$

Sometimes you will want to "rotate" an $n \times m$ matrix into an $m \times n$ one, so that the first row becomes the first column, the second row becomes the second column, and so on. This is called the *transpose*. I write the transpose of \mathbf{A} as \mathbf{A}^{\top} , though you will often also see it written \mathbf{A}' . For example:

$$\mathbf{A} = \begin{bmatrix} 99 & 73 & 2 \\ 13 & 40 & 41 \end{bmatrix} \qquad \Leftrightarrow \qquad \mathbf{A}^{\top} = \begin{bmatrix} 99 & 13 \\ 73 & 40 \\ 2 & 41 \end{bmatrix}$$

Some of the most commonly invoked properties of the transpose are:

$$(\mathbf{A}^{\top})^{\top} = \mathbf{A},$$

$$(c\mathbf{A})^{\top} = c\mathbf{A}^{\top},$$

$$(\mathbf{A} + \mathbf{B})^{\top} = \mathbf{A}^{\top} + \mathbf{B}^{\top},$$

$$(\mathbf{A} - \mathbf{B})^{\top} = \mathbf{A}^{\top} - \mathbf{B}^{\top}.$$

A matrix is *square* if it has the same number of rows as columns, i.e., it is $n \times n$. Every matrix is special, but some kinds of square matrix are *especially* special.

• A symmetric matrix is equal to its transpose: $\mathbf{A} = \mathbf{A}^{\top}$. Example:

$$\begin{bmatrix} 1 & 10 & 100 \\ 10 & 2 & 0.1 \\ 100 & 0.1 & 3 \end{bmatrix}$$

• A diagonal matrix contains zeroes everywhere except along the main diagonal: if $i \neq j$, then $a_{ij} = 0$. A diagonal matrix is symmetric by definition. Example:

$$\begin{bmatrix} 1 & 0 & 0 \\ 0 & 2 & 0 \\ 0 & 0 & 3 \end{bmatrix}$$

• The $n \times n$ identity matrix, written \mathbf{I}_n (or just \mathbf{I} when the size is clear from context), is the $n \times n$ diagonal matrix where each diagonal entry is 1. Example:

$$\begin{bmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{bmatrix}$$

And last we come to matrix multiplication. Whereas matrix addition and subtraction are pretty intuitive, matrix multiplication is not. Let \mathbf{A} be an $n \times m$ matrix and \mathbf{B} be an $m \times p$ matrix. (Notice that the number of columns of \mathbf{A} must match the number of rows of \mathbf{B} .) Then $\mathbf{C} = \mathbf{A}\mathbf{B}$ is an $n \times p$ matrix whose ij'th element is the dot product of the i'th row of \mathbf{A} and the j'th column of \mathbf{B} :

$$c_{ij} = a_{i1}b_{1j} + a_{i2}b_{2j} + \cdots + a_{im}b_{mj}$$
.

Some examples might make this more clear.

$$\mathbf{A} = \begin{bmatrix} 2 & 10 \\ 0 & 1 \\ -1 & 5 \end{bmatrix}, \mathbf{B} = \begin{bmatrix} 1 & 4 \\ -1 & 10 \end{bmatrix}$$

$$\mathbf{AB} = \begin{bmatrix} 2 \cdot 1 + 10 \cdot (-1) & 2 \cdot 4 + 10 \cdot 10 \\ 0 \cdot 1 + 1 \cdot (-1) & 0 \cdot 4 + 1 \cdot 10 \\ (-1) \cdot 1 + 5 \cdot (-1) & (-1) \cdot 4 + 5 \cdot 10 \end{bmatrix} = \begin{bmatrix} -8 & 108 \\ -1 & 10 \\ -6 & 46 \end{bmatrix}$$

And here's one that you'll start seeing a lot of soon.

$$\mathbf{A} = \begin{bmatrix} 1 & x_{11} & x_{12} \\ 1 & x_{21} & x_{22} \\ \vdots & & \\ 1 & x_{N1} & x_{N2} \end{bmatrix}, \mathbf{B} = \begin{bmatrix} \beta_0 \\ \beta_1 \\ \beta_2 \end{bmatrix}$$
$$\mathbf{AB} = \begin{bmatrix} \beta_0 + \beta_1 x_{11} + \beta_2 x_{12} \\ \beta_0 + \beta_1 x_{21} + \beta_2 x_{22} \\ \vdots \\ \beta_0 + \beta_1 x_{N1} + \beta_2 x_{N2} \end{bmatrix}$$

Some important properties of matrix multiplication:

- Matrix multiplication is associative: (AB)C = A(BC).
- Matrix multiplication is distributive: A(B + C) = AB + AC.
- For any $n \times m$ matrix \mathbf{A} , we have $\mathbf{AI}_m = \mathbf{I}_n \mathbf{A} = \mathbf{A}$. In this way, the identity matrix is kind of like the matrix equivalent of the number one. (More on this when we get to matrix inversion.)
- Matrix multiplication is *not* commutative. In other words, $\mathbf{AB} \neq \mathbf{BA}$ except in very special cases (e.g., one of them is the identity matrix).

This is obvious when we're dealing with non-square matrices. Let **A** be $n \times m$ and **B** be $m \times p$, so that **AB** exists. Then **BA** doesn't even exist unless n = p. Even then, if $n \neq m$, then **AB** is $n \times n$ and **BA** is $m \times m$, so they can't possibly be the same.

For an example that $AB \neq BA$ even for square matrices:

$$\mathbf{A} = \begin{bmatrix} 1 & 0 \\ 2 & 0 \end{bmatrix}, \mathbf{B} = \begin{bmatrix} 1 & 0 \\ 0 & 0 \end{bmatrix},$$
$$\mathbf{AB} = \begin{bmatrix} 1 & 0 \\ 2 & 0 \end{bmatrix}, \mathbf{BA} = \begin{bmatrix} 1 & 0 \\ 0 & 0 \end{bmatrix}.$$

• The transpose of the product is the product of the transposes ... but the other way around: $(\mathbf{A}\mathbf{B})^{\top} = \mathbf{B}^{\top}\mathbf{A}^{\top}$.

This is intuitive, if you think about it. Suppose **A** is $n \times m$ and **B** is $m \times p$. Then **AB** is $n \times p$, so $(\mathbf{AB})^{\top}$ should be $p \times n$. Therefore, \mathbf{B}^{\top} must come first.

6.3 Matrix Inversion

We've covered matrix addition, subtraction, and multiplication. What about division?

Let's think about division of real numbers for a second. We know that any division problem can be rewritten as a multiplication problem,

$$\frac{a}{b} = a \times b^{-1},$$

where b^{-1} is the unique real number such that

$$b \times b^{-1} = 1$$
.

Similarly, in matrix algebra, we say that the $n \times n$ matrix \mathbf{C} is an *inverse* of the $n \times n$ matrix \mathbf{A} if $\mathbf{AC} = \mathbf{CA} = \mathbf{I}_n$.

Some basic properties of matrix inverses:

- If C is an inverse of A, then A is an inverse of C. This is immediate from the definition.
- If C and D are both inverses of A, then C = D. Proof: If C and D are inverses of A, then we have

$$\begin{aligned} \mathbf{AC} &= \mathbf{I} \Leftrightarrow \mathbf{D(AC)} = \mathbf{DI} \\ &\Leftrightarrow (\mathbf{DA)C} = \mathbf{D} \\ &\Leftrightarrow \mathbf{IC} = \mathbf{D} \\ &\Leftrightarrow \mathbf{C} = \mathbf{D}. \end{aligned}$$

As a consequence of this property, we write the inverse of A, when it exists, as A^{-1} .

- The inverse of the inverse of **A** is **A**: $(\mathbf{A}^{-1})^{-1} = \mathbf{A}$.
- If the inverse of **A** exists, then the inverse of its transpose is the transpose of the inverse: $(\mathbf{A}^{\top})^{-1} = (\mathbf{A}^{-1})^{\top}$.
- Matrix inversion inverts scalar multiplication: if $c \neq 0$, then $(c\mathbf{A})^{-1} = (1/c)\mathbf{A}^{-1}$.
- The identity matrix is its own inverse: $\mathbf{I}_n^{-1} = \mathbf{I}_n$.

Some matrices are not *invertible*; i.e., their inverse does not exist. As a simple example, think of

$$\mathbf{A} = \begin{bmatrix} 0 & 0 \\ 0 & 0 \end{bmatrix}.$$

It's easy to see that, for any 2×2 matrix **B**, we have

$$\mathbf{AB} = \begin{bmatrix} 0 & 0 \\ 0 & 0 \end{bmatrix} \neq \mathbf{I}_2.$$

Therefore, A does not have an inverse.

Remember that matrix inversion is kind of like division for scalar numbers. In that light, the previous example is a generalization of the principle that you can't divide by zero. But matrices full of zeroes are not the only ones that aren't invertible. For instance, it may not be obvious at first glance, but the following matrix is not invertible:

$$\mathbf{A} = \begin{bmatrix} 1 & 2 \\ 2 & 4 \end{bmatrix}.$$

We know that because of the following theorem: A matrix is invertible if and only if its columns are linearly independent. In the above example, the second column is 2 times the first column, so the columns are not linearly independent, so the matrix is not invertible.

Consider the general 2×2 matrix

$$\mathbf{A} = \begin{bmatrix} a & b \\ c & d \end{bmatrix}.$$

We have a simple criterion for linear independence here. In particular, the columns of **A** are linearly independent if and only if $ad \neq bc$, or $ad - bc \neq 0$. We call this the *determinant* of the matrix, since it determines whether the matrix is invertible.² Moreover, if $ad - bc \neq 0$ we have

$$\mathbf{A}^{-1} = \frac{1}{ad - bc} \begin{bmatrix} d & -b \\ -c & a \end{bmatrix}.$$

I'll leave it to you to convince yourself that's true. For now, let's try a couple of examples.

 $^{^2}$ On the determinants of 3×3 and larger matrices, see your friendly local linear algebra textbook. Calculating the determinant becomes exponentially more complicated with the size of the matrix.

$$\mathbf{A} = \begin{bmatrix} 1 & 1 \\ 0 & 1 \end{bmatrix}, \mathbf{A}^{-1} = \begin{bmatrix} 1 & -1 \\ 0 & 1 \end{bmatrix},$$

$$\mathbf{A} = \begin{bmatrix} 4 & 6 \\ 2 & 4 \end{bmatrix}, \mathbf{A}^{-1} = \begin{bmatrix} 1 & -1.5 \\ -0.5 & 1 \end{bmatrix},$$

$$\mathbf{A} = \begin{bmatrix} 10 & 25 \\ 4 & 10 \end{bmatrix}, \mathbf{A}^{-1} \text{ does not exist.}$$

6.4 Solving Linear Systems

You may remember from high school being asked to solve for x_1 and x_2 in systems of equations like the following one:

$$2x_1 + x_2 = 10,$$

$$2x_1 - x_2 = -10.$$

Matrix algebra lets us write this whole system as a single equation, $\mathbf{A}\mathbf{x} = \mathbf{b}$, where

$$\mathbf{A} = \begin{bmatrix} 2 & 1 \\ 2 & -1 \end{bmatrix},$$

$$\mathbf{x} = \begin{bmatrix} x_1 \\ x_2 \end{bmatrix},$$

$$\mathbf{b} = \begin{bmatrix} 10 \\ -10 \end{bmatrix}.$$

This suggests a natural way to solve for \mathbf{x} :

$$\mathbf{x} = \mathbf{A}^{-1}\mathbf{b}.$$

In fact, the linear system of equations $\mathbf{A}\mathbf{x} = \mathbf{b}$ has a unique solution if and only if \mathbf{A} is invertible. Otherwise, it has either zero solutions or infinitely many solutions.

Example with zero solutions:

$$x_1 + x_2 = 1,$$

$$2x_1 + 2x_2 = 10.$$

Example with infinitely many solutions:

$$x_1 + x_2 = 1,$$

$$2x_1 + 2x_2 = 2.$$

6.5 Appendix: Matrices in R

We use the matrix() command to create matrices.

```
A \leftarrow matrix(c(2, 1, 3, 4),
             nrow = 2,
             ncol = 2)
Α
##
         [,1] [,2]
## [1,]
            2
                  3
## [2,]
                  4
            1
Notice that it fills "down" by column. To fill "across" instead, use the byrow argument:
B \leftarrow matrix(c(2, 1, 3, 4),
             nrow = 2,
             ncol = 2,
             byrow = 2)
В
         [,1] [,2]
##
## [1,]
            2
                  1
## [2,]
            3
There are a few utilities for checking the dimension of a matrix.
nrow(A)
## [1] 2
ncol(A)
## [1] 2
dim(A)
## [1] 2 2
To extract the i'th row, j'th column, or ij'th element, use square brackets.
A[1, ] # 1st row
## [1] 2 3
A[, 2] # 2nd column
## [1] 3 4
A[2, 1] # entry in 2nd row, 1st column
```

[1] 1

Notice that when you extract a row or column, R turns it into a vector—the result has only a single dimension. If you dislike this behavior (i.e., you want an extracted column to be a 1-column matrix), use the drop = FALSE option in the square brackets.

A[, 2, drop = FALSE]

```
## [,1]
## [1,] 3
## [2,] 4
```

Adding and subtracting matrices works as you'd expect.

A + B

A - B

As does scalar multiplication.

5 * A

```
## [,1] [,2]
## [1,] 10 15
## [2,] 5 20
```

-1 * B

```
## [,1] [,2]
## [1,] -2 -1
## [2,] -3 -4
```

However, the * operator performs element-by-element multiplication, not matrix multiplication.

A * B

```
## [,1] [,2]
## [1,] 4 3
## [2,] 3 16
```

To perform matrix multiplication, use the %*% operator.

A %*% B

```
## [,1] [,2]
## [1,] 13 14
## [2,] 14 17
```

To invert a matrix or solve a linear system, use the solve() function.

```
# Invert A
solve(A)

## [,1] [,2]
## [1,] 0.8 -0.6
## [2,] -0.2 0.4

# Solve for x in Ax = (3, 2)
solve(A, c(3, 2))

## [1] 1.2 0.2
```

Here is a not-so-fun fact about matrix inversion in R: it's not entirely exact. To see this, let's invert a matrix with some decimal elements.

```
X <- matrix(c(1.123, 2.345, 3.456, 4.567), 2, 2)
Y <- solve(X)
Y

## [,1] [,2]
## [1,] -1.53483  1.16145
## [2,] 0.78808 -0.37741</pre>
```

Now let's see what we get when we multiply X and Y.

```
X %*% Y
```

```
## [,1] [,2]
## [1,] 1.00e+00 1.4798e-16
## [2,] 8.21e-17 1.0000e+00
```

Let's check that our result is *numerically* equal to what we expected. By numerically equal, I mean, loosely speaking, that any differences are less than the amount of error you would expect due to floating point error. First we'll use diag() to generate a 2×2 identity matrix, then we'll compare numerical equality using all.equal().

```
I <- diag(2)
all.equal(X %*% Y, I)</pre>
```

```
## [1] TRUE
```

Whereas the traditional == operator is stricter, checking for exact equality.

```
X %*% Y == I
```

```
## [,1] [,2]
```

```
## [1,] TRUE FALSE
## [2,] FALSE FALSE
```

Moral of the story: when comparing decimal numbers, use all.equal() rather than ==. When all.equal() is not TRUE, it returns a message indicating how far apart the numbers are. This is annoying if you want to use all.equal() in, say, an if/else statement. To get around that, we have the isTRUE() function.

```
all.equal(1.0, 1.5)
```

[1] "Mean relative difference: 0.5"

```
isTRUE(all.equal(1.0, 1.5))
```

```
## [1] FALSE
```

One last thing. If solve() throws an error that says "reciprocal condition number..." or "system is exactly singular", that means you tried to invert a matrix that is not invertible.

```
Z <- matrix(c(1, 1, 2, 2), 2, 2)
solve(Z)</pre>
```

Error in solve.default(Z): Lapack routine dgesv: system is exactly singular: U[2,2] = 0
Sad!

Bibliography

Bowers, J. (2011). Six Steps to a Better Relationship with Your Future Self. *The Political Methodologist*, 18(2):2–8.

Leek, J. (2015). The Elements of Data Analytic Style. Leanpub.

Wickham, H. (2014). Tidy Data. Journal of Statistical Software, 59(10).

Wilson, G., Aruliah, D. A., Brown, C. T., Hong, N. P. C., Davis, M., Guy, R. T., Haddock, S. H. D., Huff, K. D., Mitchell, I. M., Plumbley, M. D., Waugh, B., White, E. P., and Wilson, P. (2014). Best Practices for Scientific Computing. *PLOS Biology*, 12(1):e1001745.