

Notes on using R

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Introduction

These notes are a tutorial on using R for empirical projects. I have tried to keep it short while still touching on the topics you need to know about to start using R for your own work. There

is much to know about R, and I encourage interested readers to consult the resources at the end of the notes for even more information.

Getting R (and R Studio)

You can download R from <https://www.r-project.org>. Technically this is all you need, but most people prefer to work in the RStudio editor, which makes it easy for you to run commands, view graphs and data, browse through help files, and more. You can download RStudio for free from <https://posit.co> (they also offer an editor called Positron that works with R and Python, but I don't have any experience with it).

Packages

Packages extend R's functionality. You can install them using `install.packages("packagename")` and load them for use using `library(packagename)` (note the you use quotes when installing but not loading). You can also use a command from a package without loading it using the syntax `packagename::command`.

Base R, Tidyverse, and data.table. The *Tidyverse* is a popular set of packages that provide different ways of working in R, and refine some of R's built-in capabilities. They can be loaded using `library(tidyverse)`. I recommend learning how to do things using R's built-in capabilities ("base R") as well as using the Tidyverse, since they both have their advantages. Another popular package for data manipulation which I don't discuss here is `data.table`, which is capable and fast, and may be of interest to you, particularly if you work with very large datasets.

Interacting with R

You can use R by directly typing commands into the *console* (the command line in the bottom window), but it is much better to create an R script (ending in `.R`), which contains a record of all of your commands. In RStudio, you can run the selected line by clicking the Run button, or by hitting `Command + Enter` on macOS or `Ctrl + Enter` on Windows. You can also run the entire file by selecting `Run all` from the dropdown menu on the Run button.

If you prefer, you can also run an entire R script using the command `source("filename.R")`. Before you do this, you will need to set the working directory to the path of your script file, which you can do using the command `setwd("path/to/file")`. If you prefer to use backslashes on Windows, you will need to *escape* them, so your command will look like `setwd("path\\to\\file")`.

It's a good idea to comment your R scripts with notes about the purpose of your code. You can use the `#` character to start a comment. In RStudio, you can also type **Command + Shift + R** to insert a nice-looking separator line.

You can view the documentation for any command by typing `help(commandname)` (this may take you to a page of search results if there are multiple similarly named commands). For me, the R documentation is hit or miss: sometimes it answers my questions and teaches me about new and useful options/commands, sometimes it's frustratingly terse. You get used to it, and you can always try a web search or LLM.

Types of data

The main data types that you will work with in R are *vectors*, *lists*, *matrices*, and *data frames* (there are also *arrays*, which generalize matrices and include matrices and data frames as a subset, but we won't be working with them).

Vectors

A vector is a list of entries that are all the same *type* (usually numeric or character). You can create a vector using the concatenate function `c()` and assign it a name using the assignment operator `<-1`:

```
a <- c(1, 2, 3)
b <- c(4, 5, 6)
c <- c("a", "b", "c")
```

You can determine an object's type by using the `class` command. You can refer to the elements of a vector using `a[1]`, or for multiple elements, `a[1:2]`. You can add/subtract and perform elementwise multiplication/division on numeric vectors using `+`, `-`, `*`, and `/`. Transformations are *vectorized*, meaning that they will be applied to all elements of a vector:

```
a + b
a - b
a * b
a / b
exp(a)
b^2
```

If you have a vector of categorical data, you can turn it into a *factor* using the `factor` command:

¹If you're feeling rebellious, you can go the other way: `c(1, 2, 3) -> a`.

```
fac.a <- factor(a)
fac.a
```

```
[1] 1 2 3
Levels: 1 2 3
```

You can also work with a vector as though it had been converted to a factor using the `as.factor` function. R provides facilities for working with factor variables (see `help(factor)`), although often you can simply leave them as numeric vectors.

Lists

A list is essentially a vector where not every element is of the same type. Lists can include vectors or other lists as elements. You can create a list using the `list()` command:

```
d <- list(1, 2, "a", "b")
e <- list(a, b, c)
```

You can view the structure of a list (or any object in R) using the `str` function. Referring to elements of a list works a little differently: `e[1]` returns a list that contains the vector `(1, 2, 3)`, while `e[[1]]` returns the vector `(1, 2, 3)` itself. Hence, `class(e[1])` returns `list` while `class(e[[1]])` returns `numeric`.

You can add names to the elements of a list using the `names` function, or when creating the list:

```
names(e) <- c("a", "b", "c")
f <- list(a = a, b = b, c = c)
```

(In the above, `a = a` means that the element named “a” takes the value `a`.) You can refer to the elements of a named list using `e[["a"]]` or `e$a`, both of which mean the same thing.

Matrices

A matrix is a collection of vectors that all have the same type and length. You can create a matrix using the `matrix` function. By default, R creates matrices by filling in the columns (this can be reversed using the `by.row = TRUE` option). All of the following create the same matrix:

```

g <- matrix(c(1, 2, 3, 4), nrow = 2)
g <- matrix(1:4, ncol = 2)
g <- matrix(seq(1, 4, by=1), nrow = 2, ncol = 2)
g

```

```

[,1] [,2]
[1,]    1    3
[2,]    2    4

```

You can create an $n \times k$ matrix of zeros using `matrix(0, n, k)` or an $n \times n$ identity matrix using `diag(n)`.² You can refer to the elements of a matrix using, e.g., `g[1, 2]` or a submatrix using `g[1:2, 2]`. You can name the rows of a matrix using `rownames` and the columns using `colnames`:

```

h <- matrix(1:9, nrow = 3)
rownames(h) <- c("r1", "r2", "r3")
colnames(h) <- c("x1", "x2", "x3")
h

```

```

x1 x2 x3
r1  1  4  7
r2  2  5  8
r3  3  6  9

```

If the rows or columns of a matrix are named, you can subset the matrix using row or column names instead of numbers, as in `h[c("r1", "r2"), c("x1", "x3")]`. You can also index a matrix by another matrix. For example, the following returns a vector containing the third element of the first row of `h`, the second element of the second row, and the first element of the third row:

```

index <- cbind(1:3, c(3, 2, 1))
index

```

```

[,1] [,2]
[1,]    1    3
[2,]    2    2
[3,]    3    1

```

²The `diag` function contains multitudes. If you have a vector `a`, `diag(a)` will create a matrix with the elements of `a` on the diagonal, and zeros elsewhere. If you have a square matrix `A`, `diag(A)` will return a vector consisting of the diagonal elements of `A`. Somehow this all makes sense when you're actually using it.

```
h[index]
```

```
[1] 7 5 3
```

`cbind` stands for column bind, and allows you to append two columns side by side (there is also `rbind`, which operates analogously on rows). Note that `h[index]` returns a vector.

Matrix addition works as you might expect. The `%*%` operator performs matrix multiplication on conformable matrices, while `*` does element by element multiplication. You can invert a matrix using `solve` and transpose it using `t`.

```
i <- matrix(5:8, nrow = 2)
i
```

```
 [,1] [,2]
[1,]    5    7
[2,]    6    8
```

```
g %*% i
```

```
 [,1] [,2]
[1,]   23   31
[2,]   34   46
```

```
g * i
```

```
 [,1] [,2]
[1,]    5   21
[2,]   12   32
```

```
solve(g)
```

```
 [,1] [,2]
[1,]   -2   1.5
[2,]    1  -0.5
```

```
t(g)
```

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

If you multiply a matrix by a vector elementwise (using `*`), or add a matrix and a vector, R will use the “recycling rule” to turn the vector into a conformable matrix. Since R fills matrices columnwise, this means that elementwise multiplication of the matrix `g` by the vector `j` in the following is the same as multiplying `g` by a matrix consisting of two copies of `j`, stacked side-by-side:

```
j <- c(1, 2)
g * j
```

```
[,1] [,2]
[1,]    1    3
[2,]    4    8
```

```
g + j
```

```
[,1] [,2]
[1,]    2    4
[2,]    4    6
```

R will interpret `j` as a row vector if you pre-multiply `g` by `j` (using matrix multiplication `%*%`), and as a column vector if you post-multiply `g` by `j` (i.e., `j %*% g` is equivalent to `matrix(j, nrow=1) %*% g`).

Data frames

A data frame is basically a list of vectors that all have the same length (i.e., a dataset). You can create a data frame using the `data.frame` function:

```
dat <- data.frame(a, b, c)
```

Since a data frame is a list, you can subset it using the same rules as for lists. For example, `dat["a"]` returns a data frame consisting of only the variable `a`, while `dat[["a"]]` and `dat$a` return the variable (vector) `a` itself. You can also subset data frames like matrices, so `dat[2:3, c("a", "c")]` returns a data frame consisting of the second and third observations on the variables `a` and `c`.

You can preview a data frame (or vector) using `str(dat)`, `head(dat)`, or in the Tidyverse, `glimpse(dat)`. In RStudio, you can also view a data frame (or any data object) by typing `View(dat)`, or by double clicking on it in the *Environment* pane.

Programming tools

Functions

You can extend R by writing your own functions. For example, suppose that you are working with the `mtcars` dataset (which comes with R and contains characteristics of different cars), and you want to know how average milage changes with the number of cylinders a car has. While there are built-in functions to handle this kind of thing, you can also write your own:

```
cyl.mean.mpg <- function(cylinders) {  
  mean(mtcars[mtcars$cyl==cylinders, ]$mpg)  
}  
cyl.mean.mpg(4)
```

```
[1] 26.66364
```

```
cyl.mean.mpg(6)
```

```
[1] 19.74286
```

Note that we use the double equals sign “`==`” when checking whether a condition holds (when evaluating a character variable, we put the condition in quotes, as in `char.var == "a"`).

This illustrates the basic use of `function`, but there are a few ways that we can make our function more useful. R functions can accept multiple inputs. And while by default they return the last object created, we can also specify what we want the function to return, which can be a scalar, vector, matrix or list.

Here is a somewhat more complex version of our function:

```
cyl.stats <- function(cylinders, variable) {  
  mean <- mean(mtcars[mtcars$cyl==cylinders, ][[variable]])  
  sd <- sd(mtcars[mtcars$cyl==cylinders, ][[variable]])  
  return(list(mean = mean, sd = sd))  
}  
cyl.stats(4, "mpg")
```

```
$mean  
[1] 26.66364
```

```
$sd  
[1] 4.509828
```

```
cyl.stats(6, "wt")
```

```
$mean  
[1] 3.117143  
  
$sd  
[1] 0.3563455
```

This function takes two arguments, `cylinders` and `variable`, and returns a list containing the mean and standard deviation of `variable` among all cars with the specified number of cylinders.

Loops

Loops allow you to perform actions iteratively. As a simple example, we could use a loop to run our basic `cyl.mean.mpg` function for cars with 4, 6, or 8 cylinders:

```
means <- rep(0, 3)  
values <- c(4, 6, 8)  
for (i in 1:3) {  
  means[i] <- cyl.mean.mpg(values[i])  
}  
means
```

```
[1] 26.66364 19.74286 15.10000
```

A few notes on this: We initially set `means` to a vector of zeros with the correct final length (an alternative would be to build the vector `means` by joining the result of each iteration with the previous one). On bigger jobs, this makes the loop faster because R doesn't have to make a new copy of the vector each time. Also, we use `cyl.mean.mpg(values[i])` to make sure that the first entry in `means` consists of the *value* of the first element of `values`, and so on.

One limitation of the loop above is that we had to specify the values of cylinders. We can avoid this by having R determine the possible values of `cyl` and the length of `means`:

```
values <- sort(unique(mtcars$cyl))  
means <- rep(0, length(values))  
for (i in seq_along(values)) {  
  means[i] <- mean(mtcars[mtcars$cyl==values[i], ]$mpg)  
}  
means
```

```
[1] 26.66364 19.74286 15.10000
```

Here, `sort(unique(mtcars$cyl))` gives us the unique values of `cyl` (in increasing order, as opposed to the order in which they occur, which `unique` provides by default), and `seq_along(cyls)` tells the loop to iterate over those values (this is basically the same as saying `for (i in 1:length(values))`), but won't throw an error if the length is zero). Note that we could have also called our `cyl.mean.mpg` function inside the loop instead of defining the value of `means` "by hand".

Loops can be nested. If we wanted to know how the means of both `mpg` and `wt` change with `cyl`, we could use:

```
values <- sort(unique(mtcars$cyl))
vars <- c("mpg", "wt")
means <- matrix(0, nrow=length(values), ncol=2)
for (i in seq_along(values)) {
  for (j in seq_along(vars)) {
    means[i, j] <- cyl.stats(values[i], vars[j])[[1]]
  }
}
rownames(means) <- paste0("cyl = ", values)
colnames(means) <- vars
means
```

	mpg	wt
cyl = 4	26.66364	2.285727
cyl = 6	19.74286	3.117143
cyl = 8	15.10000	3.999214

As we will see below, loops are useful for modifying a data frame. They can also be used to modify objects that aren't elements of an existing data frame. For example, suppose that we wanted the exponential of three matrices, `x1`, `x2`, and `x3`. One way we could create these matrices using a loop is with the `assign` function:

```
set.seed(12345)
for (i in 1:3) {
  assign(paste0("x", i), matrix(runif(4), ncol = 2))
}
```

Here, we use the `paste0` function to create the strings "`x1`", "`x2`", and "`x3`" (there is also `paste`, which allows you to specify a separator like "`_`"). If we tried to say `paste0("x", 1) <- matrix(...)`, we'd get an error message because R would think we were trying to assign

something to the literal string “x1”. Instead, we use the `assign` function to tell R that we are creating an object whose name is that string. We also use the `runif` function to populate the matrices with pseudo-random draws from a uniform distribution (and seed the random number generator to make sure that we get the same values every time).

We could also use a loop to define transformations of these matrices. To do this, we use the `get` function (which is the inverse of `assign`) to retrieve the objects whose names are given by the strings “x1”, “x2”, and “x3”:

```
for (i in 1:3) {
  assign(paste0("x", i, "_exp"), exp(get(paste0("x", i))))
}
```

Alternatively, we could collect our matrices into a list, then work with the list. The `mget` function allows us to get a list of multiple objects, all at once. Once we have a list, we can iterate through the elements of the list:

```
matrices <- mget(paste0("x", 1:3))
for (i in 1:3) {
  matrices[[i]] <- exp(matrices[[i]])
}
```

The first time I tried this, I accidentally typed `exp(matrices[i])` and got an error, because `matrices[1]` is a list containing `x1` (which can't be exponentiated), while `matrices[[1]]` is the matrix `x1` itself, which can. Now, we could return to the `assign` function to add these transformed matrices as objects in our environments. Alternatively, we can use the `list2env` function, which handles this for us (in this case, we have to tell R to save the matrices in the *global environment*, which is the main workspace where objects are stored):

```
names(matrices) <- paste0("x", 1:3, "_exp")
list2env(matrices, envir = .GlobalEnv)
```

Conditionals and while loops

Instead of looping over a set index, we can use `while` to continue looping while a condition is met. Similarly, we can use `if`, `else if` and `else` to execute commands based on whether a condition is met.

In the following example, we illustrate these commands by using the bisection method to find the positive root of $f(x) = x^2 - 16$.³

³Here is the bisection algorithm. Pick a point L where $f(L) < 0$ and a point H where $f(H) > 0$, then calculate the midpoint M . If $|f(M)| < \epsilon$, stop. Otherwise, if $f(L) * f(M) < 0$, the root is between L and M , so start over, replacing H with M . If $f(L) * f(M) > 0$, the root is between M and H , so start over, replacing L with M .

```

f <- function(x) x^2 - 16
# initial values
eps <- 1
low <- 0
high <- 5
i <- 1 # no. of iterations
while (eps > 10^(-5)) {
  mid <- .5*(low + high)
  eps <- abs(f(mid))
  if (f(low) * f(mid) < 0) {
    high <- mid
  } else {
    low <- mid
  }
  i <- i + 1
}
eps

```

[1] 7.629395e-06

mid

[1] 4.000001

i

[1] 21

Apply functions

Loops are useful programming tools, but sometimes it is more convenient to use one of the “apply” family of functions, which are abstractions of loops known as *functionals* (programming with them is called *functional programming*).

The `apply` function itself applies a function to every row or column of an array (e.g., a matrix or data frame). For example, if we wanted the mean of every column of `mtcars`, we could use

```
apply(mtcars, 2, mean)
```

	mpg	cyl	disp	hp	drat	wt	qsec
20.090625	6.187500	230.721875	146.687500	3.596563	3.217250	17.848750	
vs	am	gear	carb				
0.437500	0.406250	3.687500	2.812500				

Or, if we wanted the sum of every row of the matrix `x1`, we could use

```
apply(x1, 1, sum)
```

```
[1] 1.481886 1.761898
```

The `lapply` function applies a function to every element of a list (I think of this as a general-purpose substitute for loops) and returns the results as a list.⁴ For example, instead of using a loop to get the mean mileage for every value of cylinder, we can use:

```
lapply(unique(mtcars$cyl), cyl.mean.mpg)
```

```
[[1]]  
[1] 19.74286  
  
[[2]]  
[1] 26.66364  
  
[[3]]  
[1] 15.1
```

Note that, here, we don't need to use constructs like `seq_along` – `lapply` automatically sets the first element of the output list to the value of `cyl.mean.mpg` applied to the first element of the input list, and so on.

We can also define functions within the `lapply` statement. Both of the following produce the same result:

```
lapply(unique(mtcars$cyl),  
      function(x) mean(mtcars[mtcars$cyl==x, ]$mpg))  
lapply(unique(mtcars$cyl), \(x) mean(mtcars[mtcars$cyl==x, ]$mpg))
```

⁴The `purrr` package from the Tidyverse has a modified version of `lapply` called `map`, which works essentially the same way, but offers a few additional conveniences (see *R for Data Science* for more).

The `\(x)` syntax is called an *anonymous function*, but it's just a shorthand for the usual function syntax. Also note that for short functions, we can dispense with the braces and just type `function(x) thing_to_do`.

You can also use `lapply` with functions that take multiple arguments by supplying additional arguments after the function:

```
lapply(unique(mtcars$cyl), cyl.stats, "mpg")
```

```
[[1]]  
[[1]]$mean  
[1] 19.74286  
  
[[1]]$sd  
[1] 1.453567
```

```
[[2]]  
[[2]]$mean  
[1] 26.66364  
  
[[2]]$sd  
[1] 4.509828
```

```
[[3]]  
[[3]]$mean  
[1] 15.1  
  
[[3]]$sd  
[1] 2.560048
```

There is a variant of `lapply` called `sapply` that tries to simplify the output as a vector or matrix, if possible (the “s” stands for simplify):

```
sapply(unique(mtcars$cyl), \(\(x) mean(mtcars[mtcars$cyl==x,]$mpg))
```

```
[1] 19.74286 26.66364 15.10000
```

The `tapply` command applies a function to a vector within groups defined by levels of another vector. We could have been using `tapply` instead of our `cyl.mpg.mean` function all along:

```
tapply(mtcars$mpg, mtcars$cyl, mean)
```

```
4       6       8  
26.66364 19.74286 15.10000
```

This is equivalent to using the `split` command to make a list of vectors, each containing the values of `mpg` for a particular level of `cyl`, then using `sapply` on that list:

```
mpg.split <- split(mtcars$mpg, mtcars$cyl)  
sapply(mpg.split, mean)
```

```
4       6       8  
26.66364 19.74286 15.10000
```

We can also use `lapply` instead of loops to work with lists of data objects. Let's use the `rm` command to remove our modified matrices, then recreate them using `lapply`:

```
rm(list=paste0("x", 1:3, "_exp"), matrices)  
matrices <- lapply(paste0("x", 1:3), get)  
matrices_exp <- lapply(matrices, exp)  
names(matrices_exp) <- paste0("x", 1:3, "_exp")  
lapply(1:3, \(i) assign(paste0("x", i, "_exp"),  
                         matrices_exp[[i]],  
                         envir = .GlobalEnv))
```

```
[[1]]  
     [,1]      [,2]  
[1,] 2.056291 2.140378  
[2,] 2.400731 2.425711
```

```
[[2]]  
     [,1]      [,2]  
[1,] 1.578509 1.384163  
[2,] 1.181012 1.664000
```

```
[[3]]  
     [,1]      [,2]  
[1,] 2.070324 1.035139  
[2,] 2.690527 1.164595
```

The above illustrates how `lapply` can be used in conjunction with `get` and `assign` to replicate the functionality provided by the `mget` and `list2env` commands. Note that, when we use `assign` inside a function, we have to specify the environment (as we did when using `list2env`), which we didn't have to do when using a loop.

There are two other functions that are useful for functional programming. Suppose that you wanted to use `cbind` to join a list of matrices or data frames. It's straightforward to do this with a loop (NULL below is a convenient way to define an object which is initially empty):

```
big.matrix <- NULL
for (i in seq_along(matrices)) {
  big.matrix <- cbind(big.matrix, matrices[[i]])
}
big.matrix
```

```
[,1]      [,2]      [,3]      [,4]      [,5]      [,6]
[1,] 0.7209039 0.7609823 0.4564810 0.3250954 0.7277053 0.03453544
[2,] 0.8757732 0.8861246 0.1663718 0.5092243 0.9897369 0.15237349
```

What if you are using `apply` functions instead? `do.call` applies a function with the elements of a list as its arguments. For example, the following is equivalent to the loop above:

```
do.call(cbind, matrices)
```

```
[,1]      [,2]      [,3]      [,4]      [,5]      [,6]
[1,] 0.7209039 0.7609823 0.4564810 0.3250954 0.7277053 0.03453544
[2,] 0.8757732 0.8861246 0.1663718 0.5092243 0.9897369 0.15237349
```

`Reduce` applies a function to the first two elements of a list, then applies the function to the result of the first step and the third element of the list, and so on. In this case, `Reduce(cbind, matrices)` produces the same result as `do.call`. However, `Reduce` also allows you to view the intermediate steps (we could replace `+` below with `sum`; the former shows how we can refer to symbolic mathematical operators):

```
Reduce(`+`, 1:5, accumulate=TRUE)
```

```
[1] 1 3 6 10 15
```

Loops vs. functionals. In R, a vectorized command like `v <- c(1:5)^2` is much faster than the loop `for (i in 1:5) v[i] <- i^2`. However, apply-style functions are really just a more concise way of writing a loop (and sometimes more concise means harder to code), so you should use whatever works best for the task at hand.

Working with data

Reading and writing data

R can read many different types of data files, but I am going to focus on the most common types. One frequently used format that is easy to work with is CSV (comma seperated value). If you have a csv file named `data.csv` saved on your desktop (on macOS), you can read it and store it as a data frame using

```
setwd("~/Desktop")
library(tidyverse)
d <- read_csv("data.csv")
```

By default, the entries in the first row will be interpreted as column headings. The `read_csv` function is part of the `readr` package from the Tidyverse. Alternatively, you can use the built-in function `read.csv`. The advantages of `read_csv` are that it can be faster and simpler to work with. It stores the data as a “Tibble,” which is a modified data frame (see *R for Data Science* for a discussion of the differences), while some packages expect a traditional data frame. If you think you’ve run into this problem, you can convert a Tibble to a data frame using `d <- data.frame(d)`.

You can save a data frame (or matrix) as a csv using the command `write_csv(d, "filename.csv")`. Alternatively, you can use the built-in command `write.csv`.

You can read Excel files using the `read_excel` command from the `readxl` package. The basic syntax is

```
d <- readxl::read_excel("data.xlsx", sheet="Sheet1")
```

The `haven` package from the Tidyverse can import data from several other statistical programs. In particular, you can use `read_dta` to import a Stata file. The `foreign` package also reads data from other statistical software.

Once you’ve imported some data, you might want to clean up the formatting a little. You can rename one or more variables using

```
names(d)[names(d)=="y1"] <- c("why1")
names(d)[names(d) %in% paste0("y", 1:3)] <- paste0("why", 1:3)
```

or in the Tidyverse using

```
d |> rename(why1 = y1)
```

You can also change the case of variables names using either of the following:

```
names(d) <- tolower(names(d))
d |> rename_with(tolower)
```

You can save the image of your entire workspace (all data, functions, etc.) as an RData file using

```
save.image("filename.RData")
```

and load an existing image using

```
load("filename.RData")
```

Aside. Here is the code I used to generate these data:

```
d <- data.frame(replicate(6, rnorm(50)))
names(d) <- c(paste0("x", 1:3), paste0("y", 1:3))
d[c("z1", "z2")] <- as.factor(replicate(2, sample(1:5, 50,
                                                 replace=TRUE)))
write.csv(d, file="data.csv", row.names = FALSE)
```

`replicate` is a member of the `apply` family (it is equivalent to `sapply(1:2, \(\lambda\)(x) sample(1:5, 50, replace=TRUE))`). `rnorm` creates draws from a normal distribution.

Subsetting data

There are several ways to prune a data frame so that it only contains certain variables. For example, if you want to drop the y variables from d, you can use any of the following:

```
d[c("x1", "x2", "x3", "z1", "z2")]
d[ , c("x1", "x2", "x3", "z1", "z2")]
d[c(paste0("x", 1:3), paste0("z", 1:2))]
d[names(d) %in% c(paste0("x", 1:3), paste0("z", 1:2))]
d[startsWith(names(d), "x") | startsWith(names(d), "y")]
d[!startsWith(names(d), "y")]
d[grep("^xz", names(d))]
subset(d, select=c(paste0("x", 1:3), paste0("z", 1:2)))
subset(d, select=-c(y1, y2, y3))
d[-c(4, 5, 6)]
```

Here are a few notes on these commands: `!startsWith(names(d), "y")` specifies the names in `d` that *don't* start with "y" (`!` is the logical negation operator). `grep("^[xz]", names(d))` is a *regular expression* that returns the names of `d` that start with either "x" or "z" (regular expressions are useful for working with strings, but I avoid them because I can never remember the syntax). `select=-c(y1, y2, y3)` returns the subset of `d` excluding the "y" variables, and `d[-c(4, 5, 6)]` returns the subset that excludes columns 4-6.

You can also drop a variable using `d$z1 <- NULL`.

In the Tidyverse, this can be accomplished using the `select` command from the `dplyr` package. All of the following do the same:

```
d |> select(x1, x2, x3, z1, z2)
d |> select(starts_with(c("x", "z")))
d |> select(num_range("x", 1:3), num_range("z", 1:2))
d |> select(!starts_with("y"))
```

This syntax deserves a bit of explanation. "`|>`" is the "pipe." It takes data and "pipes" it into the first argument of the next command (and so only works if the next command takes data as its first argument). The pipe is part of base R, but is most often used with Tidyverse commands. Note that since we didn't use `d <-` to modify the data frame, running these commands will simply print the data frame (actually, it will print a preview, which is one of the ways that Tibbles differ from data frames).

Now suppose that you only wanted to retain observations where `z1` is 4 or 5 and `z2` is 2. Any of the following would work:

```
d[(d$z1==4 | d$z1==5) & d$z2==2,]
d[d$z1 %in% c(4, 5) & d$z2==2, ]
subset(d, d$z1 %in% c(4, 5) & d$z2==2)
```

The syntax `|` means "or". Sometimes it's useful to have a detailed understanding of how this kind of subsetting works. The first command above is equivalent to

```
condition <- (d$z1==4 | d$z1==5) & d$z2==2
d[condition, ]
```

In the above, `condition` is a logical vector (TRUE/FALSE values) indicating whether the condition holds, and `d[condition,]` is the subset of `d` for observations that satisfy the condition (i.e., where `condition == TRUE`).

We can also use `dplyr`'s `filter` command:

```
d |> filter((z1==4 | z1==5) & z2==2)
d |> filter(z1 %in% c(4, 5) & z2==2)
```

Note. You can use the pipe to “chain” commands together (this is more common with Tidyverse commands, but it works in base R, too):

```
d |> select(z1, z2) |> filter(z1 %in% c(4, 5) & z2==2)
d |> subset(select=c(z1, z2)) |> subset((z1==4 | z1==5) & z2==2)
```

We could sort our data according to the variable `z1` using either of

```
d[order(d$z1), ]
d |> arrange(z1)
```

If we wanted to sort according to `z1`, and then the value of `z2` within levels of `z1`, we could use `order(d$z1, d$z2)` or `arrange(z1, z2)`.

Transforming data

The variable `z1` in the data frame `d` takes the values 1-5. Suppose we want to recode 4s and 5s to 1 and all other values to zero. We could do this using

```
d[d$z1==3 | d$z1==2 | d$z1==1, ]$z1 <- 0
d[d$z1==4 | d$z1==5, ]$z1 <- 1
```

If `d` is saved as a data frame instead of a Tibble, this will give us an error if no observations satisfy the condition in brackets, because then we’re trying to replace something that has length zero with something that has positive length (this is another way that Tibbles differ from data frames). We can avoid this by using the following alternative syntax, which always works:

```
d[d$z1==3 | d$z1==2 | d$z1==1, "z1"] <- 0
d[d$z1==4 | d$z1==5, "z1"] <- 1
```

Missing values. R stores missing values as `NA`. Our data don’t have any missing values, but if they did, R would return an error using either of these approaches. How missing values are handled differs across commands, so it’s a good idea to be vigilant about potential NAs. If we have missing values (or to be safe if we’re not sure), we can use the following:

```
d[(d$z1==3 | d$z1==2 | d$z1==1) & !is.na(d$z1), "z1"] <- 0
d[(d$z1==4 | d$z1==5) & !is.na(d$z1), "z1"] <- 1
```

We could simplify this using the `%in%` operator (which also handles NAs correctly):

```
d[d$z1 %in% c(1, 2, 3), "z1"] <- 0
d[d$z1 %in% c(4, 5), "z1"] <- 1
```

We could also do this in a single line using *indicator function* notation, in which `(expression)` evaluates to TRUE if the expression is true and FALSE otherwise. In numeric settings, R will interpret TRUE as 1 and FALSE as 0, and we can use this to force the result to be numeric. Since the indicator function will return zeros for NAs, to be safe we should use:

```
d[!is.na(d$z1),]$z1 <- 1*(d[!is.na(d$z1),]$z1 %in% c(4, 5))
```

If we wanted to do this for both `z1` and `z2`, we could use a loop:

```
vars <- paste0("z", 1:2)
for (i in vars) {
  d[!is.na(d[i]), ][i] <- 1*(d[!is.na(d[i]), ][i] %in% c(4, 5))
}
```

Note here that we need to use `d[[i]] %in% ...` because `%in%` expects a vector, not a data frame.

To avoid the problems with these approaches, it might be easier to use the built-in `replace` function:

```
d$z1 <- replace(d$z1, d$z1 %in% c(1, 2, 3), 0)
d$z1 <- replace(d$z1, d$z1 %in% c(4, 5), 1)
```

which we could automate using a loop or `lapply`:

```
d[paste0("z", 1:2)] <- lapply(d[paste0("z", 1:2)],
                                \x) replace(x, x %in% c(1,2,3), 0)
d[paste0("z", 1:2)] <- lapply(d[paste0("z", 1:2)],
                                \x) replace(x, x %in% c(4,5), 1))
```

The `dplyr` package from the Tidyverse has convenient functions for doing this sort of thing. To recode `z1`, we can use:

```
d <- d |> mutate(z1 = case_when(z1 %in% c(1, 2, 3) ~ 0,
                                    z1 %in% c(4, 5) ~ 1,
                                    .default = z1))
```

`mutate` is dplyr's command for transforming and adding variables. The `case_when` syntax tells R to replace `z1` with 0 when `z1=5`, and to use the existing value of `z1` otherwise.

If we wanted to transform both `z1` and `z2` at the same time, we could use:

```
d <- d |> mutate(across(c(z1, z2),
                           \((x) case_when(x %in% c(1, 2, 3) ~ 0,
                                             x %in% c(4, 5) ~ 1,
                                             .default = x)))
```

In the above, we could replace `c(z1, z2)` with `all_of(vars)`, `num_range("z", 1:2)`, or `starts_with("z")`.

If we wanted to create a new variable `x1_sq` equal to the square of `x1`, we could use

```
d$x1_sq <- d$x1^2
```

If we wanted to do this for all of the `x` variables, we could use either

```
for (i in 1:3) {
  d[paste0("x", i, "_sq")] <- d[paste0("x", i)]^2
}
```

or the vectorized approach

```
d[paste0("x", 1:3, "_sq")] <- d[paste0("x", 1:3)]^2
```

In the Tidyverse, we could use

```
d <- d |> mutate(x1_sq = x1^2)
```

for one variable, or for multiple variables,

```
d <- d |> mutate(x1_sq = x1^2, x2_sq = x2^2)
d <- d |> mutate(across(num_range("x", 1:3),
                        .fns = list(sq = \((x) x^2),
                        .names = "{col}_{fn}")))
```

We could add additional functions to the list if we wanted to add multiple transformations of these variables.

Merging and reshaping data

Imagine that our (pre re-coding) variables `z1` and `z2` represent “state” and “year”. Suppose that we have another dataset that records the values of some different variables for each state and year. How could we merge this dataset to our original dataset `d`?

First, let’s make such a dataset. We want all combinations of `z1` and `z2`, each of which take the values 1-5. We can create all of these combinations using the `expand.grid` function, then make some fake data for each combination:

```
temp <- expand.grid(z1=1:5, z2=1:5)
w1 <- runif(25)
w2 <- runif(25)
d.new <- data.frame(cbind(temp, w1, w2))
```

We can use the `merge` function to combine this new data frame with `d`:

```
d.merged <- merge(d, d.new, by=c("z1", "z2"))
```

There are some options to the `merge` function. For example, you might want the merged data to include values from `d.new` that don’t appear in `d` (which doesn’t occur in our example), see `help(merge)` for details.

You can also do this using the `left_join` command from `dplyr`:

```
d.merged2 <- d |> left_join(d.new, join_by(z1, z2))
```

A left join keeps all rows in the original dataset (we might throw away some rows from the new dataset if they don’t correspond to any values in the original). There are also right joins (keep all rows from the new dataset), full joins (keep all rows from both), and inner joins (only keep rows that appear in both). These correspond to the options in the base-R `merge` function.

Now suppose that the variables `x1-x3` and `y1-y3` represent observations on the same variable for different units in different time periods (i.e., panel data). It is often easier to work with panel data in *long form*, which has one column per variable but multiple rows per unit, one for each time period.

It isn’t too hard to reshape a dataset into long form in base R, but I always found the documentation a little confusing. First, we need to add an “id” variable that identifies each unit (we can do this using the `dim` function, which returns a vector containing the dimensions of an array). Then we can use the `reshape` command:

```

d <- read.csv("data.csv")
d$id <- 1:dim(d)[1]
d.long <- reshape(d, idvar="id",
                   varying=c(paste0("x", 1:3), paste0("y", 1:3)),
                   sep="", direction="long")

```

We could use regular expressions to replace the `varying` command with `varying=grep("^x|^y", names(d))`, which selects all variables that start with “x” or “y” (or we could just type everything manually using `varing=c("x1", "x2", ...)`).

In the Tidyverse, this can be accomplished using the `pivot_longer` command:

```

d.long <- d |> pivot_longer(cols = c(paste0("x", 1:3),
                                         paste0("y", 1:3)),
                               names_to = c(".value", "time"),
                               names_pattern = "([A-Za-z]+)(\\d+)")

```

Here, `cols` determines which columns get reshaped to long form, `names_to` determines what we do with the different parts of variable names like `x1` (we use the first part to name the variable and the second to index the time period), and `names_pattern` determines how we split names like `x1` into parts. In this case, `names_pattern` uses a regular expression that looks for letters followed by numbers. Unfortunately, this is the only syntax that reliably works in all cases (if our variables were named like “`x_1`”, we could replace this with the simpler `names_sep="_"`). We could, however, replace the `cols` statement with the simpler `cols = starts_with(c("x", "y"))` or `cols = c(num_range("x", 1:3), num_range("y", 1:3))`.

To reshape back to “wide” form, we can use

```

d.wide <- reshape(d.long, idvar="id", timevar="time",
                   direction="wide", v.names=c("x", "y"))

```

or, in the Tidyverse,

```

d.wide <- d.long |> pivot_wider(names_from="time",
                                    values_from=c("x", "y"),
                                    names_glue=".value}{time}")

```

Analysis

Summary statistics

R doesn't have a great built-in way to easily get descriptive statistics. You can get some descriptives for a data frame (or a subset of one) using:

```
summary(d)
```

	x1	x2	x3	y1
Min.	-1.94396	-2.0193	-2.395694	-2.15245
1st Qu.	-0.62954	-0.8372	-0.620201	-0.81119
Median	0.07309	-0.2028	0.006369	0.13922
Mean	0.10972	-0.1018	0.018148	-0.02562
3rd Qu.	0.61040	0.4488	0.593582	0.55230
Max.	2.72821	2.1323	1.918224	1.90431
	y2	y3	z1	z2
Min.	-1.8943	-2.7330	1.00	1.00
1st Qu.	-0.4492	-0.8780	2.00	2.00
Median	0.2419	-0.4337	3.00	3.00
Mean	0.1493	-0.2758	2.92	3.06
3rd Qu.	0.7228	0.6060	4.00	4.00
Max.	2.4996	2.0274	5.00	5.00
	id			
Min.	1.00			
1st Qu.	13.25			
Median	25.50			
Mean	25.50			
3rd Qu.	37.75			
Max.	50.00			

but chances are you want to know the sample size, standard deviation, etc. as well.

However, there are several packages that can do this. One easy approach is using `describe` from the `psych` package:

```
psych::describe(d, fast=TRUE)
```

vars	n	mean	sd	median	min	max	range	skew	kurtosis	se	
x1	1	50	0.11	1.02	0.07	-1.94	2.73	4.67	0.52	0.09	0.14
x2	2	50	-0.10	0.96	-0.20	-2.02	2.13	4.15	0.24	-0.44	0.14

```

x3    3 50  0.02  0.91   0.01 -2.40  1.92  4.31 -0.05   0.22 0.13
y1    4 50 -0.03  1.04   0.14 -2.15  1.90  4.06 -0.25   -0.78 0.15
y2    5 50  0.15  0.87   0.24 -1.89  2.50  4.39 -0.08   -0.06 0.12
y3    6 50 -0.28  0.99   -0.43 -2.73  2.03  4.76 -0.22   -0.01 0.14
z1    7 50  2.92  1.35   3.00  1.00  5.00  4.00  0.00   -1.27 0.19
z2    8 50  3.06  1.41   3.00  1.00  5.00  4.00 -0.15   -1.32 0.20
id    9 50 25.50 14.58  25.50  1.00 50.00 49.00  0.00   -1.27 2.06

```

Here, we used the `fast=TRUE` option to get a more limited set of statistics. You can also use `describeBy` to get descriptives within groups.

You can also calculate your own descriptive statistics. For example, to get the sample size, mean and standard deviation, you could use

```
apply(d[paste0("y", 1:3)], 2, \(\text{x}) c(n=length(x), mean=mean(x), sd=sd(x)))
```

```

          y1        y2        y3
n      50.0000000 50.000000 50.000000
mean -0.02562224 0.149286 -0.2757643
sd     1.04306947 0.868742  0.9878856

```

Our sample data don't have any missing values, but if they did, our statistics would also be `NA`. We could use `is.na` to get the sample size, and add the `na.rm=TRUE` option to the `mean` and `sd` functions:

```
apply(d[paste0("y", 1:3)], 2, \(\text{x}) c(n=sum(is.na(x)),
                                         mean=mean(x, na.rm=TRUE),
                                         sd=sd(x, na.rm=TRUE)))
```

You can calculate statistics within groups using `aggregate` (which omits missing values by default). To get the sample size, mean and median of `y1-y3` by `z1` and `z2` you could use:

```
head(aggregate(cbind(y1, y2, y3) ~ z1 + z2, d,
                \(\text{x}) c(n = length(x), mean=mean(x), sd=sd(x))))
```

	<code>z1</code>	<code>z2</code>	<code>y1.n</code>	<code>y1.mean</code>	<code>y1.sd</code>	<code>y2.n</code>	<code>y2.mean</code>	<code>y2.sd</code>
1	2	1	3.00000000	-0.15625681	1.59970107	3.0000000	-0.1648793	1.5722075
2	3	1	2.00000000	-0.02454541	0.20067036	2.0000000	0.2116156	1.0023726
3	4	1	3.00000000	0.08727342	0.64314253	3.0000000	0.1789289	0.7663552
4	5	1	2.00000000	1.02753392	0.14723381	2.0000000	0.9686784	0.3591143
5	1	2	2.00000000	-0.20859931	2.67238968	2.0000000	0.9521382	0.9171654

```

6 2 2 1.00000000 -1.02335712      NA 1.0000000 -0.2331576      NA
      y3.n     y3.mean     y3.sd
1 3.0000000 -0.7569015 0.1534366
2 2.0000000  0.1597525 0.6511911
3 3.0000000 -1.3442514 1.2223017
4 2.0000000 -0.2588588 0.9739089
5 2.0000000 -0.3362057 0.4113071
6 1.0000000 -0.7457210      NA

```

Here, we used `cbind` to apply `aggregate` to multiple variables (this trick often works with commands that take “~” formulas, including the regression command `lm` introduced below). We also used the `head` function to only print the first few rows of output. From here, you could also store the results as a data frame in order to export them to a spreadsheet program for further editing.

The `dplyr` package is particularly convenient for descriptive statistics. To get the sample size, mean and standard deviation, we can use `summarize`:

```
d |> summarize(y1_mean = mean(y1, na.rm=TRUE), y1_sd = sd(y1, na.rm=TRUE),
n=n())
```

```

y1_mean     y1_sd   n
1 -0.02562224 1.043069 50

```

```
d |> summarize(across(num_range("y", 1:3),
list(mean = \((x) mean(x), sd = \((x) sd(x))), n=n()))
```

```

y1_mean     y1_sd   y2_mean     y2_sd     y3_mean     y3_sd   n
1 -0.02562224 1.043069 0.149286 0.868742 -0.2757643 0.9878856 50

```

the sample size `n` gets special treatment here because it applies to the entire dataset, not any specific variable (alternatively, we could use `n=\(x) sum(is.na(x))` as above).

We could do this by `z1` and `z2`:

```
d |> group_by(z1, z2) |> summarize(across(num_range("y", 1:3),
list(mean = \((x) mean(x, na.rm=TRUE),
sd = \((x) sd(x, na.rm=TRUE))), n=n()))
```

```
`summarise()` has grouped output by 'z1'. You can override using the `groups` argument.
```

```
# A tibble: 23 x 9
# Groups:   z1 [5]
  z1    z2 y1_mean  y1_sd y2_mean  y2_sd y3_mean  y3_sd     n
  <int> <int>    <dbl>   <dbl>    <dbl>   <dbl>    <dbl>   <dbl> <int>
1     1     2  -0.209   2.67    0.952   0.917  -0.336   0.411     2
2     1     3  -0.388   NA     -0.155   NA     0.896   NA      1
3     1     4   0.691   1.57    0.162   0.953  -0.160   0.675     4
4     1     5   0.641   0.288   0.193   0.726  -0.346   1.36      3
5     2     1  -0.156   1.60    -0.165   1.57   -0.757   0.153     3
6     2     2  -1.02    NA     -0.233   NA     -0.746   NA      1
7     2     3   0.199   0.798   0.303   0.444  -0.106   1.84      4
8     2     4  -0.995   NA     0.275   NA     0.723   NA      1
9     2     5  -0.716   NA     -1.35    NA     -0.806   NA      1
10    3     1  -0.0245  0.201   0.212   1.00    0.160   0.651     2
# i 13 more rows
```

If you wanted to add these summaries back to your dataset, you could save them as a data frame, then do a merge. Alternatively, you can use base-R's `ave` function:

```
d[paste0("y", 1:3, "_mean")] <- lapply(d[paste0("y", 1:3)],
  \((x) ave(x, list(d$z1, d$z2),
  FUN=mean))
```

(it might be simpler to do this using a loop), or obtain the statistics using dplyr's `mutate`:

```
d <- d |> group_by(z1, z2) |> mutate(across(num_range("y", 1:3),
  list(mean = \((x) mean(x),
  sd = \((x) sd(x)),
  .names = "{col}_{fn}"),
  n=n()))
```

You can create tables for discrete variables using the `table` command. The basic syntax simply produces counts, but it can be used with `prop.table` to get proportions:

```
table(d$z1)
```

```
1 2 3 4 5
10 10 11 12 7
```

```
prop.table(table(d$z1))
```

```
1   2   3   4   5  
0.20 0.20 0.22 0.24 0.14
```

To perform crosstabs, you can use either `table` or the somewhat more convenient `xtabs` command. When we use `prop.table` with a two-dimensional table, we have to specify whether we want row proportions (with a 1) or column proportions (with a 2).

```
prop.table(table(d$z1, d$z2), 1)
```

```
1           2           3           4           5  
1 0.00000000 0.20000000 0.10000000 0.40000000 0.30000000  
2 0.30000000 0.10000000 0.40000000 0.10000000 0.10000000  
3 0.18181818 0.18181818 0.00000000 0.36363636 0.27272727  
4 0.25000000 0.08333333 0.33333333 0.25000000 0.08333333  
5 0.28571429 0.28571429 0.14285714 0.14285714 0.14285714
```

```
prop.table(xtabs(~ z1 + z2, d), 2)
```

```
z2  
z1      1       2       3       4       5  
1 0.00000000 0.25000000 0.10000000 0.30769231 0.33333333  
2 0.30000000 0.12500000 0.40000000 0.07692308 0.11111111  
3 0.20000000 0.25000000 0.00000000 0.30769231 0.33333333  
4 0.30000000 0.12500000 0.40000000 0.23076923 0.11111111  
5 0.20000000 0.25000000 0.10000000 0.07692308 0.11111111
```

We could use, e.g., `round(prop.table(xtabs(~ z1 + z2, d), 2), 2)`, to round the decimals to two places.

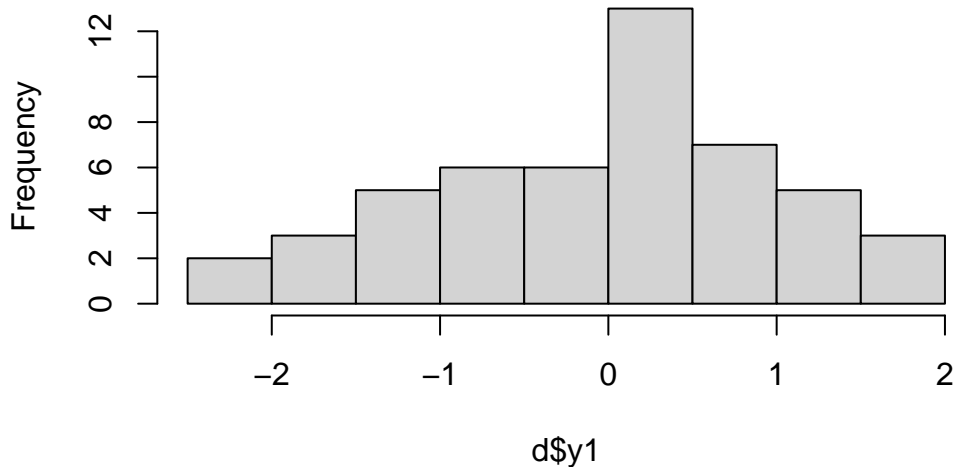
Visualizing data

R's graphing capabilities extend well beyond what can be covered in a short introduction, so we'll only sketch the basics, give a sense of the possibilities, and point in the direction of further resources.

The basics of R's built-in plotting functions are easy. For example, you can obtain histogram using:

```
hist(d$y1)
```

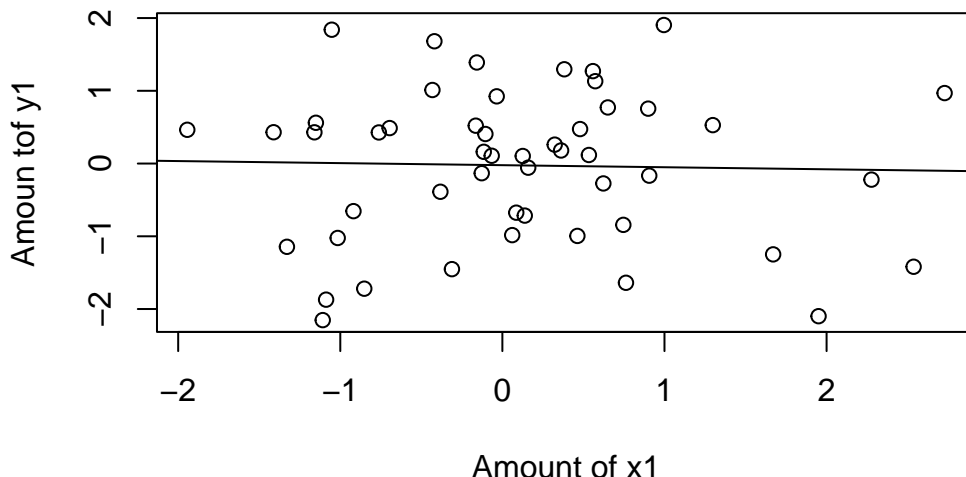
Histogram of d\$y1



Here is a scatter plot, with a title and custom axis labels and a regression line (via the `abline` command):

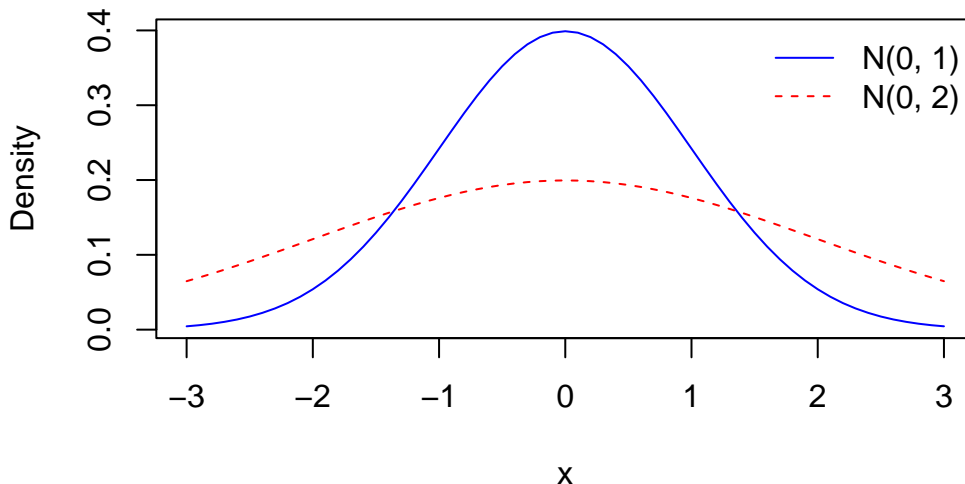
```
plot(d$x1, d$y1, xlab="Amount of x1", ylab="Amount of y1",
      main="Scatter plot of y1 against x1")
abline(lm(y1 ~ x1, d))
```

Scatter plot of y1 against x1



I often use Base-R graphics to create simple line diagrams. Here's how we can plot the densities for two normal distributions with different variances:

```
x <- seq(-3, 3, by=.1)
d1 <- dnorm(x, 0, 1)
d2 <- dnorm(x, 0, 2)
plot(x, d1, type="l", col="blue", ylab="Density")
lines(x, d2, type="l", lty=2, col="red")
legend("topright", bty="n", c("N(0, 1)", "N(0, 2)"),
       col=c("blue", "red"), lty=1:2)
```



Here, `dnorm` returns the normal density (there is also `pnorm` for the distribution function and `qnorm` for the quantiles, useful for computing critical values), `plot` displays the initial curve, `lines` adds the second, `legend` adds the legend, `type="l"` gives a line (rather than underlying points), `col` controls the colors, `lty` controls the line type (solid or dashed), and `bty="n"` stops R from putting a box around the legend.

As you can see, base-R plots have a lot of options. You can read about them by typing `help(plot)`, or just search the web.

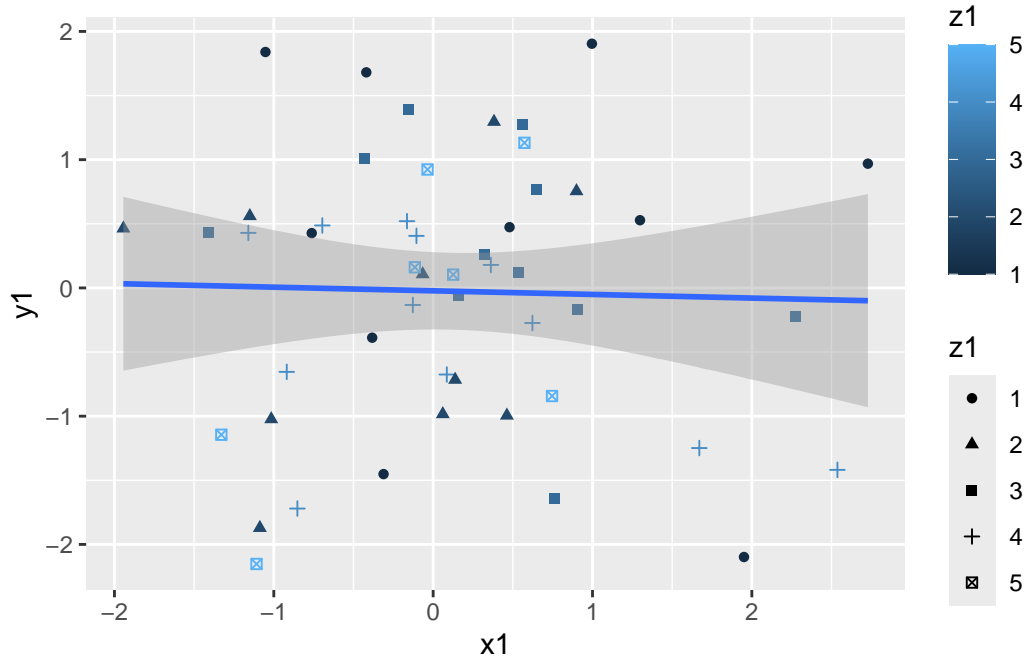
There is another plotting system called `ggplot2` which is part of the Tidyverse, is very versatile, and has much more accessible documentation. Roughly speaking, you specify the data, then the geometric objects, describing the aesthetics of the plot along the way. For example, we could make a basic histogram and scatterplot using, saving the latter as a PDF, using

```
ggplot(data=d, aes(x=x1)) + geom_histogram()
ggplot(d, aes(x=x1, y=y1)) + geom_point()
ggsave("scatter.pdf")
```

Here are some more complicated examples, adapted from *R for Data Science*. The following produces a scatterplot of y_1 against x_1 , with the color and shape of the points determined by the value of z_1 , along with a linear fit and labelled axes (note that this requires that we treat z_1 as a factor variable):

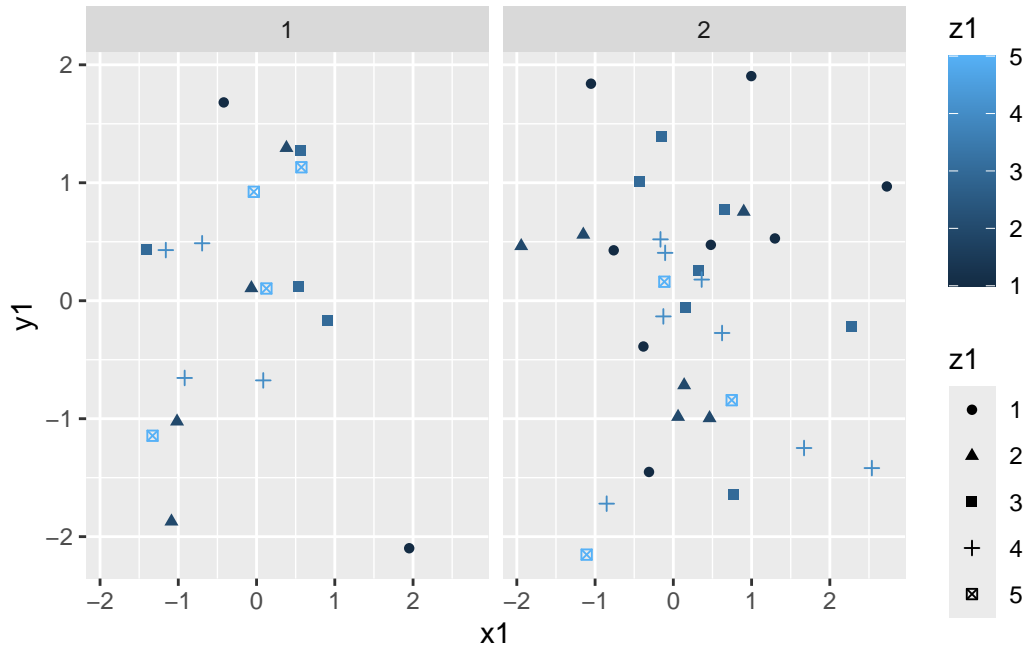
```
ggplot(d, aes(x1, y1)) + geom_point(aes(color=z1, shape=as.factor(z1))) +
  geom_smooth(method="lm") +
  labs(x="x1", y="y1", color="z1", shape="z1")
```

`geom_smooth()` using formula = 'y ~ x'



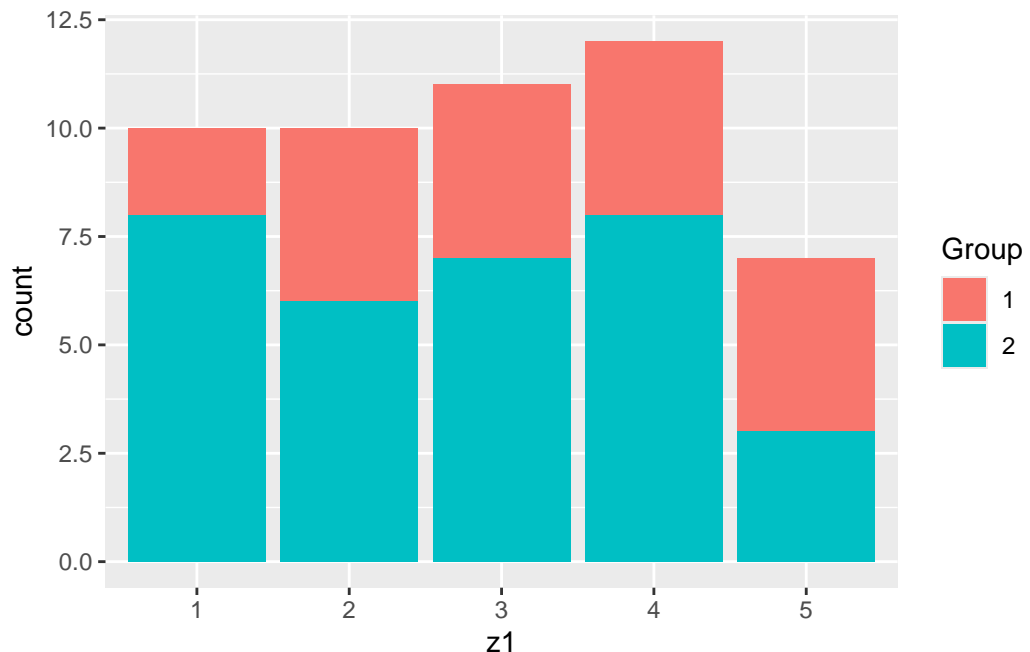
We can also use `facet_wrap` (also see `facet_grid`) to make subplots for different values of a categorical variable:

```
# make a binary group variable based on z2
d <- d |> mutate(gp = factor(1*(z2<3) + 2*(z2>=3)))
ggplot(d, aes(x1, y1)) + geom_point(aes(color=z1, shape=as.factor(z1))) +
  facet_wrap(~ gp) + labs(x="x1", y="y1", color="z1", shape="z1")
```



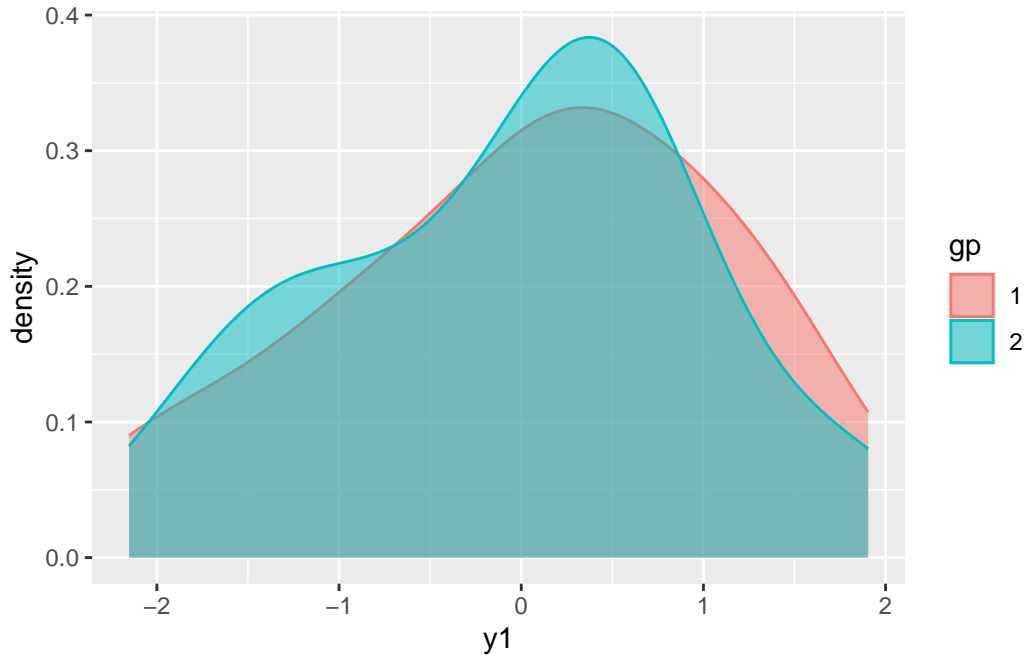
Here is a bar graph of $z2$, with the bars broken down by `group` (note that we use `as.factor` because $z1$ is not saved as a factor variable):

```
ggplot(d, aes(as.factor(z1), fill=gp)) + geom_bar() +
  labs(x="z1", fill="Group")
```



And here are separate kernel densities for y_1 for each value of $group$:

```
ggplot(d, aes(y1, color=gp, fill=gp)) + geom_density(alpha=.5)
```



Regression basics

An introduction like this one is also not the place to go into detail on statistical or econometric estimation commands, but we can highlight some techniques that come up frequently.

In R, estimated models are most usefully stored as objects which can be accessed later. For example, to run an OLS regression and view the results, you can use

```
model1 <- lm(y1 ~ x1 + x2 + x3, d)
summary(model1)
```

Call:
lm(formula = y1 ~ x1 + x2 + x3, data = d)

Residuals:

Min	1Q	Median	3Q	Max
-2.0277	-0.7679	0.1993	0.7355	2.1677

Coefficients:

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	-0.01206	0.14987	-0.080	0.936
x1	-0.06640	0.15017	-0.442	0.660
x2	0.01543	0.15725	0.098	0.922
x3	-0.25943	0.16790	-1.545	0.129

Residual standard error: 1.049 on 46 degrees of freedom

Multiple R-squared: 0.05134, Adjusted R-squared: -0.01053

F-statistic: 0.8298 on 3 and 46 DF, p-value: 0.4843

Since the model has been saved, we can refer back to it and everything saved along with it (you can see everything saved with the model using `str(model1)` or `names(model1)`). For example, we can extract the coefficients using either of the following `model$coefficients` or `coef(model1)`.

We can extract the variance-covariance matrix using

```
vcov(model1)
```

	(Intercept)	x1	x2	x3
(Intercept)	0.0224618504	-0.002226572	0.002111048	-0.0007521723
x1	-0.0022265716	0.022551337	0.003262413	0.0046395898
x2	0.0021110485	0.003262413	0.024728827	0.0026378130
x3	-0.0007521723	0.004639590	0.002637813	0.0281889529

and the residuals and predicted values using `model1$residuals` and `model1$fitted.values`. If we wanted to obtain the predicted values from applying our model to a different dataset, we could use `predict(model1, dataset)`.

R has useful syntax for adding terms to a model. We can use : to get the interaction between two variables, * to get interactions and main effects, I() for transformations of variables, and `as.factor` to include indicators for every level of a categorical variable:

```
lm(y1 ~ x1*x2 + x1:x3 + I(x2^2) + as.factor(z1), d)
```

Call:

```
lm(formula = y1 ~ x1 * x2 + x1:x3 + I(x2^2) + as.factor(z1),
  data = d)
```

```

Coefficients:
(Intercept)          x1          x2      I(x2^2) as.factor(z1)2
               0.38126     -0.09714     0.06716     0.11604     -0.71730
as.factor(z1)3 as.factor(z1)4 as.factor(z1)5      x1:x2      x1:x3
               -0.15187    -0.86856    -0.80041     0.02989     -0.03123

```

Note that when we use `lm` without saving it, R simply prints the coefficients (without standard errors or other statistics).

When we use `as.factor` to include all levels of `z1`, R automatically chooses an omitted category. We can omit the constant by starting the right-hand side of the formula with a zero (`y1 ~ 0 + as.factor(z1) + ...`), in which case R won't omit any category. We can also specify the omitted category by including `z1` in the formula using `relevel(as.factor(z1), ref=2)`. If we are going to be treating `z1` as a factor in many commands, we can permanently convert and set its baseline category using

```

d$z1 <- as.factor(d$z1)
d$z1 <- relevel(d$z1, ref=2)

```

We can use the `linearHypothesis` function from the `car` package to test linear hypotheses, and the `deltaMethod` function to test nonlinear ones:

```

library(car)
linearHypothesis(model1, c("x1 + x2=0"))

```

```

Linear hypothesis test:
x1 + x2 = 0

Model 1: restricted model
Model 2: y1 ~ x1 + x2 + x3

Res.Df   RSS Df Sum of Sq    F Pr(>F)
1      47 50.628
2      46 50.575  1  0.053104 0.0483  0.827

```

```

deltaMethod(model1, c("x1^2 + x2/2"))

```

	Estimate	SE	2.5 %	97.5 %
<code>x1^2 + x2/2</code>	0.012123	0.078401	-0.141540	0.1658

By default, R doesn't report confidence intervals, but you can obtain them using `confint(model1)`.

The sandwich package supports robust and clustered standard errors. The easiest way to report them is using the `coeftest` function from the lmtest package:

```
library(sandwich)
library(lmtest)
coeftest(model1, vcov=vcovHC)
```

t test of coefficients:

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	-0.012058	0.159304	-0.0757	0.9400
x1	-0.066404	0.177596	-0.3739	0.7102
x2	0.015426	0.151925	0.1015	0.9196
x3	-0.259434	0.205761	-1.2609	0.2137

```
coeftest(model1, vcov=vcovCL(model1, cluster = ~ z2))
```

t test of coefficients:

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	-0.012058	0.099796	-0.1208	0.90436
x1	-0.066404	0.076382	-0.8694	0.38916
x2	0.015426	0.171451	0.0900	0.92870
x3	-0.259434	0.139238	-1.8632	0.06882 .

```
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Note that there are several variants on the robust and cluster-robust variance estimators, and you can use the `type` option to specify which one you want (see `help(vcovHC)` and `help(vcovCL)` for some details).

You can estimate binary choice models using the generalized linear models function `glm`. The syntax for a probit is

```
d$y1.bin <- (d$y1 >= 0)
probit <- glm(y1.bin ~ x1 + I(x1^2) + as.factor(z1),
               family=binomial(link=probit), d)
summary(probit)
```

```

Call:
glm(formula = y1.bin ~ x1 + I(x1^2) + as.factor(z1), family = binomial(link = probit),
     data = d)

Coefficients:
              Estimate Std. Error z value Pr(>|z|)
(Intercept) 0.74569   0.47774  1.561   0.119
x1          -0.16422   0.21318 -0.770   0.441
I(x1^2)      -0.08172   0.13168 -0.621   0.535
as.factor(z1)2 -0.73018   0.60978 -1.197   0.231
as.factor(z1)3 -0.24982   0.58816 -0.425   0.671
as.factor(z1)4 -0.87343   0.57885 -1.509   0.131
as.factor(z1)5 -0.54821   0.66249 -0.828   0.408

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 68.593 on 49 degrees of freedom
Residual deviance: 64.627 on 43 degrees of freedom
AIC: 78.627

Number of Fisher Scoring iterations: 4

```

We could get a logit by using `link=logit` instead.

Both the `marginaleffects` and `margins` package can be used to obtain marginal effects. For example, we can obtain average marginal effects using

```

library(marginaleffects)
avg_slopes(probit)

```

Term	Contrast	Estimate	Std. Error	z	Pr(> z)	S	2.5 %	97.5 %
x1	dY/dX	-0.0666	0.074	-0.899	0.368	1.4	-0.212	0.0785
z1	2 - 1	-0.2663	0.210	-1.266	0.205	2.3	-0.679	0.1459
z1	3 - 1	-0.0844	0.197	-0.429	0.668	0.6	-0.470	0.3010
z1	4 - 1	-0.3212	0.196	-1.636	0.102	3.3	-0.706	0.0637
z1	5 - 1	-0.1959	0.233	-0.841	0.400	1.3	-0.653	0.2607

Type: response

Since we used `I()` to include the quadratic term and `as.factor` to include `z1`, `marginalEffects` knows to treat `x1` and its square as a single expression, and to determine the marginal effects of `z1` using discrete differences.

There are several packages that format regression tables and export them to other formats (e.g., LaTeX or HTML), including `texreg`, `stargazer`, and `modelsummary`, all of which allow you to customize the contents of the tables.⁵

For example, we can run regressions for each level of `z1` (as we saw in the *Programming tools* section, there are multiple ways to do this), then summarize the results in a single table, and export that table (the following example uses `texreg`, but the alternative packages work similarly):

```
library(texreg)
models.z1 <- lapply(sort(unique(d$z1)), \((i) lm(y1 ~ x1 + x2 + x3,
                                                 d[d$z1==i, ]))
names(models.z1) <- paste0("z1 = ", sort(unique(d$z1)))
screenreg(models.z1)
```

=====					
	z1 = 1	z1 = 2	z1 = 3	z1 = 4	z1 = 5
(Intercept)	0.57 (0.60)	-0.20 (0.48)	0.45 (0.30)	-0.16 (0.17)	-0.04 (0.27)
x1	-0.30 (0.46)	0.17 (0.47)	-0.46 (0.32)	-0.44 * (0.17)	1.11 (0.36)
x2	-0.03 (0.81)	0.02 (0.61)	-0.12 (0.29)	-0.07 (0.15)	-0.43 (0.35)
x3	-0.35 (0.52)	-0.02 (0.49)	-0.27 (0.35)	-0.59 * (0.20)	-1.68 (0.65)
R^2	0.10	0.02	0.25	0.62	0.83
Adj. R^2	-0.35	-0.46	-0.07	0.48	0.67
Num. obs.	10	10	11	12	7
=====					

*** p < 0.001; ** p < 0.01; * p < 0.05

⁵Although these commands export LaTeX tables, I often find that they need further editing. Instead, I export to HTML (which can be opened in Excel), edit them, then paste them into LyX or use Excel2LaTeX to get LaTeX-formatted tables.

```
htmlreg(models.z1, file="models_z1.html")
```

If we want to use robust or clustered standard errors, we need to do a little more work to add them to the table. Here's how we can display robust standard errors (and significance stars based on them) using stargazer:

```
library(texreg)
ses.z1 <- lapply(models.z1, \(\(x) coeftest(x, vcovHC)[, 2]))
ps.z1 <- lapply(models.z1, \(\(x) coeftest(x, vcovHC)[, 4]))
screenreg(models.z1, override.se = ses.z1, override.pvalues = ps.z1)
```

```
=====
      z1 = 1   z1 = 2   z1 = 3   z1 = 4   z1 = 5
-----
(Intercept)  0.57    -0.20    0.45   -0.16   -0.04
             (0.81)   (0.62)   (0.43)   (0.19)   (0.39)
x1          -0.30    0.17   -0.46   -0.44 **   1.11
             (0.66)   (1.00)   (0.50)   (0.09)   (0.62)
x2          -0.03    0.02   -0.12   -0.07   -0.43
             (1.34)   (1.30)   (0.34)   (0.21)   (0.56)
x3          -0.35   -0.02   -0.27   -0.59   -1.68
             (0.68)   (0.67)   (0.55)   (0.37)   (0.75)
-----
R^2          0.10    0.02    0.25    0.62    0.83
Adj. R^2     -0.35   -0.46   -0.07    0.48    0.67
Num. obs.    10      10      11      12      7
=====
*** p < 0.001; ** p < 0.01; * p < 0.05
```

As another example, we could create a table consisting of different model specifications:

```
formulas <- c("x1", "x1 + x2", "x1 + x2 + x3")
specifications <- lapply(formulas, \(\(x) lm(paste0("y1 ~", x), d))
screenreg(specifications)
```

```
=====
      Model 1   Model 2   Model 3
-----
```

(Intercept)	-0.02	-0.02	-0.01
	(0.15)	(0.15)	(0.15)
x1	-0.03	-0.02	-0.07
	(0.15)	(0.15)	(0.15)
x2		0.04	0.02
		(0.16)	(0.16)
x3			-0.26
			(0.17)
<hr/>			
R^2	0.00	0.00	0.05
Adj. R^2	-0.02	-0.04	-0.01
Num. obs.	50	50	50
<hr/>			
*** p < 0.001; ** p < 0.01; * p < 0.05			

Here are a few additional packages that are useful for regressions and econometrics:

- The plm package estimates panel data models, including fixed and random effects
- The fixest package estimates fixed effects models efficiently, works with multiple dimensions of fixed effects, and has built-in clustering and graphing capabilities
- The ivreg packages handles instrumental variables and two-stage least squares

Resources for more

- [An introduction to R](#) by Venables, Smith, et al. is a classic introduction to R, and also comes bundled with R itself
- [FasteR: Fast Lane to Learning R](#) by Matloff is a detailed tutorial, with a focus on base R
- [R for Data Science](#) by Wickham, Cetinkaya-Rundel and Grolemund is an excellent introduction to using R via the Tidyverse (the [first edition](#) touches on a few useful topics omitted from the second)
- [Advanced R](#) by Wickham delves into some of the more technical details of working with R (also see the [first edition](#))
- [Ggplot2: Elegant Graphics for Data Analysis](#) by Wickham is a deep dive into ggplot2
- [The Art of R Programming](#) by Matloff provides a lot of details on R programming, with a focus on base R