Exploratory Data Analysis with R

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Contents

X.	hat	you will learn	7
* *		sou win learn	7
		ics	8
	_	nnologies	8
	reci	inologies	0
н	ow to	o use this book	13
	Text	t, instructions, and explanations	13
		ode and output	14
\mathbf{G}	et ur	o and running	15
	-	erent ways to run RStudio	15
		alling R and RStudio locally	16
		uick look at RStudio	16
	_	king at the Console in RStudio	19
	,,,,,	ming at the common in increase	10
Ι	Int	roduction to R	21
1	Δ α	uick introduction to R	23
_	1.1	Using R as a big calculator	23
	1.2	Problematic calculations	26
	1.3	Storing and reusing results	$\frac{20}{26}$
	$1.3 \\ 1.4$	How does assignment work?	30
	1.5	Global environment	32
	1.6	Naming rules and conventions	$\frac{32}{32}$
	1.0	Naming rules and conventions	32
2	Usi	ng functions	35
	2.1	Introduction	35
	2.2	Functions and arguments	35
	2.3	Evaluating arguments and returning results	37
	2.4	Specifying function arguments	39
	2.5	Combining functions	40
	2.6	Functions do not have 'side effects'	41

3	Vec	tors	43
	3.1	Introduction	43
	3.2	Atomic vectors	43
	3.3	Numeric vectors	44
	3.4	Constructing numeric vectors	46
	3.5	Named vectors	46
	3.6	Vectorised operations	47
	3.7	Other kinds of atomic vectors	48
4	Dat	a frames	5 3
	4.1	Introduction	53
	4.2	Data frames	54
	4.3	Exploring data frames	56
	4.4	Extracting and adding a single variable	57
5	Pac	kages	61
	5.1	The R package system	61
	5.2	Task views	62
	5.3	Using packages	62
	5.4	Package data	65
	5.5	The tidyverse ecosystem of packages	66
II	D	ata Wrangling	67
6	Get		69
	6.1	Introduction	69
	6.2	Tidy data	CC
			69
	6.3	Penguins! $+=$	71
	$6.3 \\ 6.4$	Penguins! +=	
			71
	6.4	Um tibbles?	71 72
7	6.4 6.5 6.6	Um tibbles?	71 72 73
7	6.4 6.5 6.6	Um tibbles? Missing values. Introducing dplyr rking with variables Introduction.	71 72 73 74
7	6.4 6.5 6.6 Wo s	Um tibbles?	71 72 73 74 7 9
7	6.4 6.5 6.6 Wo 7.1	Um tibbles? Missing values. Introducing dplyr rking with variables Introduction.	71 72 73 74 79 79
	6.4 6.5 6.6 Wo 7.1 7.2 7.3	Um tibbles? Missing values Introducing dplyr rking with variables Introduction Subset variables with select Creating variables with mutate	71 72 73 74 79 79 80
	6.4 6.5 6.6 Wo 7.1 7.2 7.3	Um tibbles? Missing values Introducing dplyr rking with variables Introduction Subset variables with select Creating variables with mutate	71 72 73 74 79 79 80 85
	6.4 6.5 6.6 Wo 7.1 7.2 7.3	Um tibbles? Missing values. Introducing dplyr rking with variables Introduction. Subset variables with select Creating variables with mutate rking with observations	71 72 73 74 79 79 80 85 89
	6.4 6.5 6.6 Wo 7.1 7.2 7.3 Wo 8.1	Um tibbles? Missing values	71 72 73 74 79 79 80 85 89 89
	6.4 6.5 6.6 Wo : 7.1 7.2 7.3 Wo : 8.1 8.2	Um tibbles? Missing values Introducing dplyr rking with variables Introduction Subset variables with select Creating variables with mutate rking with observations Introduction Relational and logical operators	71 72 73 74 79 79 80 85 89 89
	6.4 6.5 6.6 Wo 7.1 7.2 7.3 Wo 8.1 8.2 8.3 8.4	Um tibbles? Missing values Introducing dplyr rking with variables Introduction Subset variables with select Creating variables with mutate rking with observations Introduction Relational and logical operators Subset observations with filter Reordering observations with arrange	71 72 73 74 79 79 80 85 89 89 90 93
8	6.4 6.5 6.6 Wo 7.1 7.2 7.3 Wo 8.1 8.2 8.3 8.4	Um tibbles? Missing values Introducing dplyr rking with variables Introduction Subset variables with select Creating variables with mutate rking with observations Introduction Relational and logical operators Subset observations with filter Reordering observations with arrange	71 72 73 74 79 79 80 85 89 89 90

CONTENTS	F
CONTENIOS	5
CONTENIO	· ·

9.3 Removing grouping information	. 105
10 Building pipelines	107
10.1 Why do we need 'pipes'?	. 107
10.2 Using pipes (%>%)	
1012 COM Proce (W N)	. 100
11 Helper functions	111
11.1 Introduction	. 111
11.2 Working with select	
11.3 Working with mutate and transmute	. 113
11.4 Working with filter	. 114
11.5 Working with summarise	. 115
III Exporing Data	117
•	
12 Exploratory data analysis	119
12.1 Introduction	
12.2 Statistical variables and data	
12.3 Populations and samples	
12.4 Types of EDA	
12.5 A primer of descriptive statistics	. 124
13 Introduction to ggplot2	131
13.1 The anatomy of ggplot2	. 132
13.2 A quick introduction to ggplot2	
13.3 A standard way of using ggplot2	
13.4 Increasing the information density	
14 Customising plots	149
14.1 Geom properties	. 150
14.2 Plot scales	. 153
14.3 Labels	. 157
14.4 Themes	. 158
14.5 Advice for making plots	. 162
15 Exploring one variable	163
15.1 Exploring numerical variables	
15.2 Exploring categorical variables	
10.2 Exploring categorical variables	. 110
16 Exploring associations	181
16.1 Associations between numeric variables	. 181
16.2 Associations between categorical variables	. 185
16.3 Categorical-numerical associations	. 190
16.4 Multivariate associations	. 194
17 Doing more with ggplot2	197

6	CONTENTS
---	----------

	17.2 17.3	Comparing descriptive statistics	05 06
A	A.1 A.2 A.3 A.4	ing help27Introduction2Browsing the help system2Searching for help files2Navigating help files2Vignettes2	11 11 12 13
В	Mar	aging projects, scripts and data files	17

What you will learn

This book provides a self-contained introduction to how to use R for exploratory data analysis. Think of it as a resource to be referred to when needed. There is no need to memorise everything in this book. Instead, aim to understand the key concepts and familiarise yourself with the content, so that you know where to look for information when you need it. The details will get easier with practise.

Aims

This book has three related aims:

- 1. Introduce the R ecosystem. R is widely used by biologists and environmental scientists to manipulate and clean data, produce high quality figures, and carry out statistical analyses. We will teach you some basic R programming so that you are in a position to address these needs in future if you need to. You don't have to become an expert programmer to have a successful career in biology but knowing a little bit of programming has almost become a prerequisite for doing research in the 21st century.
- 2. Demonstrate how to use R to carry out data manipulation and visualisation. Designing good experiments, collecting data, and analysis are hard, and these activities often take a great deal time and money. If you want to effectively communicate your hard-won results, it is difficult to beat a good figure or diagram; conversely, if you want to be ignored, put everything into a boring table. R is really good at producing figures, so even if you end up just using it as a platform for visualising data, your time hasn't been wasted.
- 3. Provides a foundation for learning statistics later on. If you want to be a biologist, particularly one involved in research, there is really no way to avoid using statistics. You might be able to dodge it by becoming a theoretician, but if that really is your primary interest you should probably being studying for a mathematics degree. For the rest of us who collect and analyse data knowing about statistics is essential: it allows us to

distinguish between real patterns (the "signal") and chance variation (the "noise").

Topics

The topics we will cover are divided into three 'blocks':

The Getting Started with R block introduces the R language and the RStudio environment. The aim is to quickly run through what you need to know to start using R productively. This includes some basic terminology, how to use R packages, and how to access help. We are not trying to turn you into an expert programmer—though you may be surprised to discover that you do enjoy it. However, by the end of this block you will know enough about R to begin learning the practical material that follows.

The **Data Wrangling** block aims to show you how to manipulate data with R. If you regularly work with data a large amount of time will inevitably be spent getting data into the format you need. The informal name for this is 'data wrangling'. This topic that is often not taught to beginners, which is a shame because mastering the art of data wrangling saves time in the long run. We'll learn how to get data into and out of R, makes subsets of important variables, create new variables, summarise your data, and so on.

The **Exploratory Data Analysis** block is all about using R to help you understand and describe your data. The first step in any analysis after you have managed to wrangle the data into shape should involve some kind of visualisation or numerical summary. You will learn how to do this using one of the best plotting systems in R: **ggplot2**. We will review the different kinds of 'variables' you might have to analyse, discuss the different ways you can describe them, and then learn how to explore relationships between variables.

Technologies

What is R?

The answer to this question very much depends on who you ask. We could go on and on about the various features that R possesses. R is a functional programming language, it supports object orientation, etc etc... but these kinds of explanations are only helpful to someone who already knows about computer languages. Here's what you need to know... When a typical R user talks about "R" they are often referring to two things at once, the GNU R language and the ecosystem that exists around the language:

• R is all about **data analysis**. We can carry out any standard statistical analysis in R, as well as access a huge array of more sophisticated tools with impressive names like "structural equation model", "random forests" and "penalized regression". These days, when statisticians and computer

scientists develop a new analysis tool, they often implement it in R first. This means a competent R user can always access the latest, cutting edge analysis tools. R also has the best graphics and plotting facilities of any platform. With sufficient expertise, we can make pretty much any type of figure we need (e.g. scatter plots, phylogenetic trees, spatial maps, or even volcanoes). In short, R is a very productive environment for doing data analysis.

• Because R is such a good environment for data analysis, a very large **community of users** has grown up around it. The size of this community has increased steadily since R was created, but this growth has really increased up in the last 5-10 years or so. In the early 2000s there were very few books about R and the main way to access help online was through the widely-feared R mailing lists. Now, there are probably hundreds of books about different aspects of R, online tutorials written by enthusiasts, and many websites that exist solely to help people learn R. The resulting ecosystem is vast, and though it can be difficult to navigate at times, when we run into an R-related problem the chances are that the answer is already written down somewhere¹.

R is not just about data analysis. R is a fully-fledged programming language, meaning that once you become proficient with it you can do things such as construct numerical simulation models, solve equations, query websites, send emails or carry out many other tasks we don't have time to write down. We won't do any of this year or next but it is worth noting that R can do much more than just analyse data if we need it to.

What is RStudio?

R is essentially just a computer program that sits there and waits for instructions in the form of text. Those instructions can be typed in by a user or they can be sent to it from another program. R also runs in a variety of different environments. The job of RStudio is to provide an environment that makes R a more pleasant and productive tool.



R and RStudio are not the same thing.

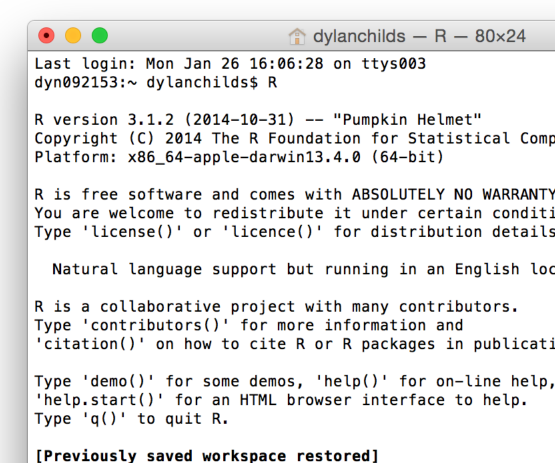
RStudio is a different program from R—it is installed separately and occupies its own place in the Programs menu (Windows PC) or Applications folder (Mac). We can run R without RStudio if we need to, but we cannot run RStudio without R. Remember that!

One way to get a sense of why RStudio is a Very Good Thing is to look at what running R without it is like. The simplest way to run it on a Linux or

¹The other big change is that R is finally starting to become part of the commercial landscape—learning how to use it can only improve your job prospects.

Unix-based machine (like a Mac) is to use something called the Terminal. It's well beyond the scope of this book to get into what this is, but in a nutshell, the Terminal provides a low-level, text-based way to interact with a computer. Here is what R looks like running inside a Terminal on a Mac:

>



We can run R in much the same way on Windows using the "Command Prompt" if we need to. The key thing you need to take away from that screenshot is that running R like this is very "bare bones". We typed the letter "R" in the Terminal and hit Enter to start R. It printed a little information as it started up and then presented us with "the prompt" (>), waiting for input. This is where we type or paste in instructions telling R what to do. There is no other way to interact with it when we run R like this – no menus or buttons, just a lonely prompt waiting for instructions.

So what is RStudio? In one sense RStudio is just another Graphical User Interface for R which improves on the "bare bones" experience. However, it is a GUI on steroids. It is more accurate to describe it as an Integrated Development Environment (IDE). There is no all-encompassing definition of an IDE, but they all exist to make programmer's lives easier by integrating various useful tools into a single piece of software. From the perspective of this book, there are four key features that we care about:

- The R interpreter—the thing that was running in the Terminal above—runs inside RStudio. It's accessed via a window labelled Console. This is where we type in instructions we want to execute when we are working directly with R. The Console also shows us any output that R prints in response to these instructions. So if we just want the "bare bones" experience, we can still have it.
- RStudio provides facilities for working with R programs using something called a Source Code Editor. An R program (also called a "script")" is just is a collection of instructions in the R language that have been saved to a text file. Nothing more! However, it is much easier to work with a script using a proper Source Code Editor than an ordinary text editor like Notepad.
- An good IDE like RStudio also gives you a visual, point-and-click means
 of accessing various language-specific features. This is a bit difficult to
 explain until we have have actually used some of these, but trust us, being
 able to do things like manage packages, set working directories, or inspect
 objects we've made simplifies day-to-day use of R. This especially true for
 new users.
- RStudio is cross-platform—it will run on a Windows PC, a Linux PC or a Mac. In terms of the appearance and the functionality it provides, RStudio is exactly the same on each of these platforms. If we learn to work with R via RStudio on a Windows PC, it's no problem migrating to a Mac or Linux PC later on if we need to. This is a big advantage for those of us who work on multiple platforms.

We'll only scratch the surface of what RStudio can do. The reason for introducing a powerful tool like RStudio is because one day you may need to access sophisticated features like debugging facilities, package build tools, and repository management. RStudio makes it easy to use these advanced tools.

How to use this book

We have adopted a number of formatting conventions in this book to distinguish between normal text, R code, file names, and so on. You need to be aware to make best use of the book.

Text, instructions, and explanations

Normal text—instructions, explanations and so on—are written in the same type as this document. We will tend to use bold for emphasis and italics to highlight specific technical terms when they are first introduced. In addition:

- This typeface is used to distinguish R code within a sentence of text: e.g. "We use the mutate function to change or add new variables."
- A sequence of selections from an RStudio menu is indicated as follows: e.g. File New File R Script
- File names referred to in general text are given in upper case in the normal typeface: e.g. MYFILE.CSV.

At various points in the text you will come across text in different coloured boxes. These are designed to highlight stand-alone exercises or little pieces of supplementary information that might otherwise break the flow. There are three different kinds of boxes:



Action!

This is an **action** box. We use these when we want you to do something. Do not ignore these boxes.



Information!

This is an **information** box. These aim to offer a discussion of why something works the way it does.



Warning!

This is a **warning** box. These usually highlight a common 'gotcha' that might trip up new users.

R code and output

We try to illustrate ideas using snippets of real R code where possible. It's a good idea to run these when working through a topic. The best way to learn something is to use it. Of course, in order to do that we need to know what we're looking at... Stand alone snippets will be formatted like this:

```
tmp <- 1
print(tmp)</pre>
```

[1] 1

At this point it does not matter what the above actually means. You just need to understand how the formatting of R code in this book works. The lines that start with ## show us what R prints to the screen after it evaluates an instruction and does whatever was asked of it, that is, they show the output. The lines that **do not** start with ## show us the instructions, that is, they show us the input. So remember, the absence of ## shows us what we are asking R to do, otherwise we are looking at something R prints in response to these instructions.

Get up and running

Different ways to run RStudio

We can run RStudio in a variety of different ways—

- 1. Most people use the version of RStudio called **RStudio Desktop**, either in its free-to-use guise (Open Source Edition) or the commercial version (RStudio Desktop Pro). The desktop version of RStudio are stand-alone applications that run locally on a computer and have to be installed like any other piece of software. This is generally easy, but you can run into problems if you have an old computer or a Chromebook.
- 2. The second way to use RStudio is by accessing a version called **RStudio Server** through a web browser. RStudio Server is usually administered by professional IT people. Life is easy if you belong to an organisation that has set up RStudio Server, because all you need to get going is a user account, a semi-modern web browser and an internet connection.
- 3. Finally, the company that makes RStudio also runs a commercial cloud-based solution called RStudio Cloud. This allows anyone web browser and an internet connection to use R and RStudio. Although there is a free version, this is fairly limited meaning you end up paying a monthly fee to do 'real work'. However, RStudio Cloud can be a good backup option when all else fails.



Do you need to install R and RStudio?

If you're lucky enough to have access to RStudio Server or an RStudio Cloud account you don't need to install R and RStudio on your own computer. Just access those cloud service through a decent web browser. That said, it can be useful to have a local copy on your own computer, e.g. because you don't have a reliable internet connection. Obviously, if you can't access those cloud services you'll have to install R and RStudio to use them!

Installing R and RStudio locally

It does not need to cost a penny to use R and RStudio. The source code for R is open source, meaning anyone with the time, energy and expertise is free to download it and alter it as they please. Open source does not necessarily mean free, as in it costs £0 to use, but luckily R is free in this sense. On the other hand, RStudio is developed and maintained by a for-profit company (called... RStudio). Luckily, because they make their money selling professional software and services, the open source desktop version of RStudio is also free to use. This section will show you how to download and install R and RStudio.

Installing R

In order to install R you need to download the appropriate installer from the Comprehensive R Archive Network (CRAN). We are going to use the "base distribution" as this contains everything you need to use R under normal circumstances. There is a single installer for Windows. On a Mac, it's important to match the installer to the version of OS X. In either case, R uses a the standard install mechanism that should be familiar to anyone who has installed an application on their machine. There is no need to change the default settings—doing so will probably lead to problems later on.

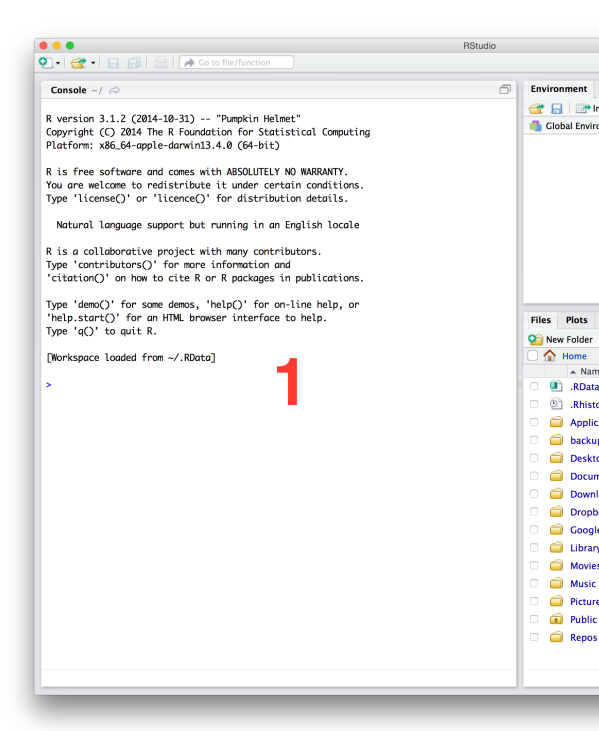
After installing R it should be visible in the Programs menu on a Windows computer or in the Applications folder on a Mac. In fact, that thing labelled 'R' is very simple a Graphical User Interface (GUI) for R. When we launch the R GUI we're presented with something called the Console, which is where we can interact directly with R by typing things at the so-called prompt, >, and a few buttons and menus for managing common tasks. We will not study the GUIs in any detail because we recommend using RStudio, but it's important to be aware they exist so that you don't accidentally use them instead of RStudio.

Installing RStudio

RStudio can be downloaded from the RStudio download page. The one to go for is the Open Source Edition of RStudio Desktop, **not** the commercial version of RStudio Desktop called RStudio Desktop Pro. RStudio installs like any other piece of software—just run the installer and follow the instructions. There's no need to configure after after installation.

A quick look at RStudio

Once installed RStudio runs like any other stand-alone application via the Programs menu or the Applications folder on a Windows PC or Mac, respectively (though it will only work properly if R is also installed). Here is how RStudio appears the first time it runs on a Mac:



There are three panes inside a single window, which we have labelled with red numbers. Each of these has a well-defined purpose. Let's take a quick look at these:

- 1. The large window on the left is the Console. This is basically where R lives inside RStudio. The Console lets you know what R is doing and provides a mechanism to interact with R by typing instructions. All this happens at the prompt, >. You will be working in the Console in the next chapter so we won't say any more about this here.
- 2. The window at the top right contains two or more tabs. One of these, labelled **Environment**, allows us to see all the R 'objects' we can currently access. Another, labelled **History**, allows us to see a list of instructions we've previously sent to R. The buttons in this tab allow us to reuse or save these instructions.
- 3. The window at the bottom right contains five tabs. The first, labelled **Files**, gives us a way to interact with the files and folders. The next tab, labelled **Plots**, is where any figures we produce are displayed. This tab also allows you to save your figures to file. The **Packages** tab is where we view, install and update packages used to extend the functionality of R. The **Help** tab is where you can access and display various different help pages. Finally, **Viewer** is an embedded web browser.



My RStudio looks different!

Don't be alarmed if RStudio looks different on your computer. RStudio saves its state between different sessions, so if you've have already messed about with it you will see these changes when you restart it. For example, there is a fourth pane that is often be visible in RStudio—the source code Editor we mentioned above.

Working at the Console in RStudio

R was designed to be used interactively—it is what is known as an **interpreted language**, which we can interact with via something called a Command Line Interface (CLI). This is just a fancy way of saying that we can type instructions to "do something" into the Console and those instructions will then be interpreted when we hit the Enter key. If our R instructions do not contain any errors, R will then do something like read in some data, perform a calculation, make a figure, and so on. What actually happens obviously depends on what we ask it to do.

Let's briefly see what all this means by doing something very simple with R. Type 1 + 3 at the Console and hit the Enter key:

1+3

[1] 4

The first line above just reminds us what we typed into the Console. The line after that beginning with ## shows us what R printed to the Console after reading and evaluating our instructions.

What just happened? We can ignore the [1] bit for now (the meaning of this will become clear later in the course). What are we left with – the number 4. The instruction we gave R was in effect "evaluate the expression $1\,+\,3$ ". R read this in, decided it was a valid R expression, evaluated the expression, and then printed the result to the Console for us. Unsurprisingly, the expression $1\,+\,3$ is a request to add the numbers 1 and 3, and so R prints the number 4 to the Console.

OK... that was not very exciting. In the next chapter we will start learning to use R to carry out more useful calculations. The important take-away from this is that this sequence of events—reading instructions, evaluating those instructions and printing their output (if there is any output)—happens every time we type or paste something into the Console and hit Enter.



What does that word 'expression' mean?

Why do we keep using that word *expression*? Here is what the Wikipedia page says:

An expression in a programming language is a combination of explicit values, constants, variables, operators, and functions that are interpreted according to the particular rules of precedence and of association for a particular programming language, which computes and then produces another value.

That probably doesn't make much sense! In simple terms, an R expression is a small set of instructions that tell R to do something. That's it. We could write 'instructions' instead of 'expressions' throughout this book but we may as well use the correct word.

$\begin{array}{c} {\rm Part} \; {\rm I} \\ \\ {\rm Introduction} \; {\rm to} \; {\rm R} \end{array}$

Chapter 1

A quick introduction to R

1.1 Using R as a big calculator

1.1.1 Basic arithmetic

The Get up and running chapter showed that R could handle familiar arithmetic operations: division, multiplication, addition and subtraction. If we want to add or subtract two numbers, we place the + or - symbol in between two numbers and hit Enter. R will read the arithmetic expression, evaluate it, and print the result to the Console. This works as you'd expect:

 $Addition\!-\!$

3 + 2

[1] 5

Subtraction-

5 - 1

[1] 4

Multiplication and division are no different. However, we can't use x or \div symbols for these operations. Instead, use * and / to multiply and divide:

Multiplication-

7 * 2

[1] 14

Division-

3 / 2

[1] 1.5

We can also exponentiate numbers, i.e. raise one number to the power of another. Use the ^ operator to do this:

Exponentiation-

4^2

[1] 16

This raises 4 to the power of 2 (i.e. we squared it). In general, we can raise a number x to the power of y using x^y . Neither x nor y need to be whole numbers either.

A

Operators?

What does 'operator' mean? An operator is simply a symbol (or sequence of symbols) that does something specific with one or more inputs. For example, operators like /, *, + and - carry out arithmetic calculations with pairs of numbers. Operators are one of the basic building blocks of a programming language like R.

1.1.2 Combining arithmetic operations

We can also combine arithmetic operations. Assume we want to subtract 6 from 2^3 . The expression to perform this calculation is:

$2^3 - 6$

[1] 2

Simple enough, but what if we had wanted to carry out a slightly longer calculation that required the last answer to then be divided by 2? This is the **wrong** way to do this:

2^3 - 6 / 2

[1] 5

The answer we expect here is 1. So what happened? R evaluated 6/2 first and then subtracted this answer from 2^3 .

If that's obvious, great. If not, it's time to learn about **order of precedence**. R uses a standard set of rules to decide the order in which calculations feed into

one another to unambiguously evaluate any expression. It uses the same order as every other computer language, which thankfully is the same one we all learn in mathematics classes at school. The order of precedence is:

- 1. exponents and roots (also, 'powers' or 'orders')
- 2. division and then multiplication
- 3. additional and then subtraction



BODMAS and friends

If you find it difficult to remember the standard order of precedence there are a load of mnemonics that can to help.

We need to control the order of evaluation to arrive at the answer we were looking for in the above example. Do this by grouping together calculations inside parentheses, i.e. 'round brackets' (and). Here's the expression we should have used:

$$(2^3 - 6) / 2$$

[1] 1

We can use more than one pair of parentheses to control the order of evaluation in more complex calculations. The order of evaluation then happens 'inside-out'. For example, if we want to find the cube root of 2 (i.e. $2^{1/3}$) rather than 2^3 in that last calculation we would instead write:

$$(2^{(1/3)} - 6) / 2$$

[1] -2.370039

The parentheses around the 1/3 are needed to ensure this is evaluated prior to being used as the exponent.



Working efficiently at the Console

Working at the Console soon gets tedious if we have to retype similar things over and over again. There is no need to do this, though. Place the cursor at the prompt and hit the up arrow. What happens? This brings back the last expression sent to R's interpreter. Hit the up arrow again to see the last-but-one expression, and so on. We go back down the list using the down arrow. Once we're at the line we need, we use the left and right arrows to move around the expression and the delete key to remove the parts we want to change. Once an expression has been edited we can hit Enter to send it to R again. Try it.

1.2 Problematic calculations

Now is a good time to highlight how R handles certain kinds of awkward numerical calculations. One of these involves division of a non-zero by 0. Mathematically, division of a finite number by 0 equals A Very Large Number: infinity. Some programming languages will respond to an attempt to do this with an error. R is a bit more forgiving:

1 / 0

[1] Inf

R has a special built-in value that allows it to handle this kind of result. This is Inf, which stands for 'infinity'.

The other special kind of value we sometimes run into is generated by calculations that don't have a well-defined numerical result. For example, look what happens when we try to divide 0 by 0:

0 / 0

[1] NaN

The NaN in this result stands for 'Not a Number'. R produces NaN because 0/0 is not defined mathematically: it produces something that is Not a Number.

The reason we are pointing out Inf and NaN is not that we expect to use them. It's important to know what they represent because they often arise due to a mistake somewhere in our code. It's hard to track down such mistakes if we don't know how Inf and NaN arise.



R as a fancy calculator

What we've seen so far is that we can interact with R via the so-called REPL: the read-evaluate-print loop. R takes user input (e.g. 1 / 0), evaluates it (1 / 0 = Inf), prints the results (## [1] Inf), and then waits for the next input (e.g. 0 / 0). This facility is handy because it means we can use R interactively, working through a set of calculations line-by-line.

1.3 Storing and reusing results

We've not yet tried to do anything remotely complicated or interesting beyond using parentheses to construct longer calculations. This approach is acceptable when a calculation is straightforward, but it quickly becomes unwieldy for dealing with anything more complicated.

The best way to see what we mean is by working through a simple example—solving a quadratic equation. You probably remember these from school. A quadratic equation looks like this:

$$a + bx + cx^2 = 0$$

If we know the values of a, b and c then we can solve this equation to find the values of x that satisfy this equation. Here's the well-known formula for these solutions:

$$x = \frac{-b \pm \sqrt{b^2 - 4ac}}{2a}$$

We can use R to calculate these solutions for us. Say that we want to find the solutions to the quadratic equation when a=1, b=6 and c=5. We have to turn the above equation into a pair of R expressions:

Solution 1-

$$(-6 + (6^2 - 4 * 1 * 5)^(1/2)) / (2 * 1)$$

[1] -1

Solution 2-

$$(-6 - (6^2 - 4 * 1 * 5)^(1/2)) / (2 * 1)$$

[1] -5

The output tells us that the two values of x that satisfy this particular quadratic equation are -1 and -5.

But what should we do if we now need to solve a different quadratic equation? Working at the Console, we could bring up the expressions we typed (using the up arrow) and change the numbers to match the new values of a, b and c. However, editing expressions like this is tedious, and more importantly, it's error-prone because we have to make sure we substitute the new numbers into precisely the right positions.

A partial solution to this problem is to store the values of a, b and c in some way so that we only have to change them one. We'll see why this is useful in a moment.

First, we need to learn how to store results in R. The key to this is to use the **assignment operator**, written as an arrow pointing to the left, <-. Sticking with the current example, we need to store the numbers 1, 6 and 5. We do this by typing out three expressions, one after the another, each time hitting enter to get R to evaluate it:

```
a <- 1
b <- 6
c <- 5
```

The exact sequence <- defines the assignment operator. R won't recognise it as assignment if we try to include a space between the < and - symbols.

Notice that R didn't print anything to screen. So what actually happened? We asked R to first evaluate the expression on the right hand side of each <- (just a number in this case) and then **assigns the result** of that evaluation instead of printing it. Each result has a name associated with it, which appears on the left hand-side of the <-.

The net result of all this is that we have stored the numbers 1, 6 and 5 somewhere in R and associated them with the letters a, b and c, respectively. We can check whether this assignment business has worked by looking at the **Environment** tab in the top right RStudio window. There should be three 'names' listed in that tab now (a, b and c) along with the associated numbers 1, 6 and 5.

What does this mean in practical terms? Look at what happens if we now type the letter **a** into the Console and hit Enter:

```
a
```

[1] 1

It looks the same as if we had typed the number 1 directly into the Console. We stored the output from three separate R expressions, associating each a name so that we can access it again¹. Whenever we use the assignment operator <- we are telling R to keep whatever kind of value results from the calculation on the right-hand side of <-, giving it the name on the left-hand side so that we can access it later.

Why is this useful? Let's imagine we want to do more than one thing with our three numbers. If we want to know their sum or their product we can now use:

Sum-

```
a + b + c
```

[1] 12

Product-

 $^{^{1}}$ Technically, this is called **binding** the name to a value. You don't need to remember this.

```
a * b * c
```

[1] 30

So... once we've stored a result and associated it with a name we can reuse it whenever needed. Returning to our example, we can now calculate the solutions to the quadratic equation by typing these two expressions into the Console:

Solution 1-

```
(-b + (b^2 - 4 * a * c)^(1/2)) / (2 * a)
```

[1] -1

Solution 2-

```
(-b - (b^2 - 4 * a * c)^(1/2)) / (2 * a)
```

[1] -5

Imagine we'd now like to find the solutions to a different quadratic equation where a=1, b=5 and c=5. We only changed the value of b here. To find the new solutions we have to do two things. First we change the value of the number associated with b...

```
b <- 5
```

...then we bring up those lines that calculate the solutions to the quadratic equation and run them, one after the other:

```
(-b + (b^2 - 4 * a * c)^(1/2)) / (2 * a)
```

[1] -1.381966

```
(-b - (b^2 - 4 * a * c)^(1/2)) / (2 * a)
```

```
## [1] -3.618034
```

We don't have to retype those expressions. We can use the up arrow to bring each one back to the prompt and hit Enter. This is much simpler than editing the expressions.

More importantly, we are beginning to see the benefits of using something like R—we can break down complex calculations into a series of steps, storing and reusing intermediate results as required.



RStudio shortcut

We use the assignment operator <- all the time when working with R. Because it's inefficient to type the < and - characters repeatedly, RStudio has a built-in shortcut for typing the assignment operator.

The shortcut is 'Alt +-'. Try it now. Move the cursor to the Console, hold down the Alt key ('Option' on a Mac), and press the - sign key. RStudio will auto-magically add insert <-. If you only learn one RStudio shortcut, learn this one! It will save you a lot of time in the long run.

1.4 How does assignment work?

When we use the assignment operator <- to associate names and values, we refer to this as creating or modifying a variable. This is much less tedious than using words like 'associate', 'value', and 'name' all the time. Why is it called a variable? What happens when we run these lines:

Create myvar and print out its value-

```
myvar <- 1
myvar
```

[1] 1

Modify myvar and and print out its new value -

```
myvar <- 7
myvar
```

[1] 7

The first time we used <- with myvar on the left-hand side, we **created** a variable myvar associated with the value 1. We then printed out the value associated with myvar. The second line myvar <- 7 modified the value of myvar to be 7 (and printed this out again). This is why we refer to myvar as a variable: we can change its value as we please.

What happened to the old value associated with myvar? In short, it is gone, kaput, lost... forever. The moment we assign a new value to myvar the old one is destroyed and can no longer be accessed. Remember this.

Keep in mind that the expression on the right-hand side of <- can be any kind of calculation and the variable can have any (valid) name we like. For example, if we want to perform the calculation $(1 + 2^3) / (2 + 7)$ and associate the result with the word <code>answer</code>, we would do this:

```
answer <- (1 + 2^3) / (2 + 7)
answer
```

[1] 1

Any expression can be used on the right-hand side of the assignment operator as along as it generates an output of some kind. For example, we can create new variables from others:

```
newvar <- 2 * answer
```

What happened here? Start at the right-hand side of <-. The expression on this side contained the variable answer so R went to see if answer actually exists. It does, so it then substituted the value associated with answer into the calculation and assigned the resulting value of 2 to newvar.

Finally, look at what happens if we copy a variable using the assignment operator:

```
myvar <- 7
mycopy <- myvar</pre>
```

We now have a pair of variables, myvar and mycopy, associated with the number 7. Each of these is associated with a **different copy** of this number. If we change the value associated with one of these variables it does not change the value of the other, as this shows:

```
myvar <- 10

myvar

## [1] 10

mycopy</pre>
```

```
## [1] 7
```

R always behaves like this unless we work hard to alter this behaviour. Remember that—every time we assign one variable to another, we actually make a completely new, independent copy of its associated value. That probably doesn't seem like an obvious or important point, but trust us, it is. It will be critical to remember this behaviour when we start learning how to manipulate data sets.

1.5 Global environment

Whenever we associate a name with a value we create a copy of both these things somewhere in the computer's memory. In R the "somewhere" is called an environment. We aren't going to get into a discussion of R's many different kinds of environments—that's an advanced topic well beyond the scope of this book. The one environment we do need to be aware of is the **Global Environment**.

Whenever we perform an assignment in the Console the variable we create is placed into the Global Environment. The set of variables currently in existence are listed in the **Environment** tab in RStudio. Take a look. There are two columns in the **Environment** tab: the first shows the names of the variables, the second summarises their values.



The Global Environment is temporary

By default, R will try to save everything in the Global Environment when we close it down and restore everything when we start the next R session. It does this by writing a copy of the Global Environment to disk. In theory, this means we can close down R, reopen it, and pick things up from where we left off. **Don't rely on this behaviour!** It just increases the risk of making a mistake.

1.6 Naming rules and conventions

We don't have to use a single letter to name things in R. We could use the words tom, dick and harry in place of a, b and c. It might be confusing to use them, but tom, dick and harry are all legal names as far as to R is concerned:

- A legal name in R is any sequence of letters, numbers, ., or _, but the sequence of characters we use must begin with a letter. Both upper and lower case letters are allowed. For example, num_1, num.1, num.1, num.1, num.1 are all legal names, but lnum and _num.1 are not because they begin with 1 and _.
- R is case sensitive—it treats upper and lower case letters as different characters. This means R treats num and Num as distinct names. Forgetting about case sensitivity is a good way to create errors when using R. Try to remember that.



Don't begin a name with.

We are allowed to begin a name with a ., but this usually is A Bad Idea. Why? Because variable names that begin with . are hidden from view in the Global Environment—the value it refers to exists but it's invisible.

This behaviour exists to allow R to create invisible variables that control how it behaves. This is useful, but it isn't really meant to be used by the average user.

Chapter 2

Using functions

2.1 Introduction

Functions are an essential building block of any programming language. The job of a function is to carry out a calculation or computation that would typically require many lines code to do 'from scratch'. Functions allow us to reuse common computations while offering some control over the precise details of what happens. To use R effectively—even if our needs are very simple—we need to understand how to use functions. This chapter aims to explain what functions are for, how to use them, and how to avoid mistakes when doing so, without getting lost in the detail of how they work.

2.2 Functions and arguments

Functions allow us to reuse a calculation. The best way to see what we mean by this is to see one in action. The round function rounds numbers to a significant number of digits (no surprises there). To use it, we could type this into the Console and hit Enter:

```
round(x = 3.141593, digits = 2)
```

We have suppressed the output so that we can unpack things a bit first. We rely on the same basic construct every time we use a function. This starts with the name of the function as the prefix. In the example above, the function name is round. After the function name, we need a pair of opening and closing parentheses. This combination of name and parentheses alerts R to fact that we are using a function.

What about the bits inside the parentheses? These are called the **arguments** of the function. That's a horrible name, but it is the one that everyone uses, so

we have to get used to it. Depending on how it was defined, a function can take zero, one, or more arguments. We will discuss this idea in more detail later in this section. In simple terms, the arguments control the behaviour of a function.

We used the round function with two arguments. We supplied each one as a name-value pair separated by a comma. When working with arguments, name-value pairs occur either side of the equals sign (=), with the argument name on the left-hand side and its value on the right-hand side. The name serves to identify which argument we are working with, and the value is the thing that controls what that argument does.

We call the process of associating argument names and values 'setting the arguments' of the function (or 'supplying the arguments'). Notice the similarity between supplying function arguments and the assignment operation discussed in the last topic. The difference is that name-value pairs are associated with the = symbol when involved in arguments.



Use = to assign arguments

Do not use the assignment operator <- inside the parentheses when working with functions. This is a "trust us" situation—you will end up in all kinds of difficulty if you do this!

The arguments control the behaviour of a function. Our job as users is to set the values of these to get the behaviour we want. However, The function determines arguments we are allowed to use, i.e. we are not free to choose whatever name we like¹.

```
round(x = 3.141593, digits = 2)
```

[1] 3.14

The round function rounds one or more numeric inputs and rounds these to a particular number of significant digits. The argument that specifies the number(s) to round is x; the second argument, digits, specifies the number of decimal places we require. Based on the supplied values of these arguments, 3.141593 and 2, respectively, the round function spits out a value of 3.14, which is then printed to the Console.

What if we had wanted to the answer to 3 significant digits? We would set the digits argument to 3:

```
round(x = 3.141593, digits = 3)
```

¹We say "typically", because R is a very flexible language and so there are certain exceptions to this simple rule of thumb. For now it is simpler to think of the names as constrained by the particular function we're using. Let's return to the example to see how all this works:

```
## [1] 3.142
```

This illustrates what we mean when we say arguments control the behaviour of the function—digits sets the number of significant digits calculated by round.

2.3 Evaluating arguments and returning results

Whenever R evaluates a function, we refer to this action as 'calling the function'. In our simple example, we called the round function with arguments \mathbf{x} and digits. That said, we often just say 'use the function' because that is more natural to most users.

Several things happen when we call functions: first they **evaluate** their arguments, then they perform some action, and finally (optionally) **return** a value to us. Let's work through what all that means...

What do we mean by the word 'evaluate'? When we call a function, what typically happens is:

- 1. the R expression on the right-hand side of an = is evaluated,
- 2. the result is associated with the corresponding argument name, and
- 3. the function does its calculations using the resulting name-value pairs.

To see how the evaluation step works, take a look at a new example using round:

```
round(x = 2.3 + 1.4, digits = 0)
```

```
## [1] 4
```

What happened above is that R evaluated 2.3 + 1.4, resulting in the number 3.7, which was then associated with the argument x. We set digits to 0 this time so that round returns a whole number, 4.

The important thing to realise is that the expression(s) on the right-hand side of the = can be anything we like. This third example essentially equivalent to the last one:

```
y <- 2.3 + 1.4
round(x = y, digits = 0)
```

```
## [1] 4
```

This time we created a new variable called y and supplied this as the value of the x argument. When we use the round function like this, the R interpreter spots that something on the right-hand side of an = is a variable and associates the value of this variable with x argument. As long as we have defined the numeric variable y at some point we can use it as the value of an argument.

At the beginning of this section, we said that a function may optionally **return** a value to us when it completes its task. That word 'return' refers to the process by which a function outputs a value. If we use a function at the Console the returned value is printed out. However, we can use this value in other ways. For example, there is nothing to stop us combining function calls with the arithmetic operations:

```
2 * round(x = 2.64, digits = 0)
```

```
## [1] 6
```

Here the R interpreter evaluates the function call and then multiplies the value it returns by 2. If we want to reuse this value, we have to assign the result of function call, for example:

```
roundnum \leftarrow 2 * round(x = 2.64, digits = 0)
```

Using a function with <- is no different from the examples using multiple arithmetic operations in the last topic. The R interpreter starts on the right-hand side of the <-, evaluates the function call there, and only then assigns the value to roundnum.

A

Argument names vs variable names

Keeping in mind what we've just learned, take a careful look at this example:

```
x <- 0
round(x = 3.7, digits = x)
```

[1] 4

What is going on here? The key to understanding this is to realise that the symbol x is used in two different ways here. When it appears on the left-hand side of the = it represents an argument name. When it appears

left-hand side of the = it represents an argument name. When it appears on the right-hand side, it is treated as a variable name, which must have a value associated with it for the above to be valid. That is a confusing way to use the <code>round</code> function, but it is perfectly valid.

The message here is that what matters is where things appear relative to the =, not the symbols used to represent them.

2.4 Specifying function arguments

So far, we have been concentrating on functions that carry out mathematical calculations with numbers. Functions can do all kinds of things. For example, some functions are designed to extract information about other functions. Take a look at the args function:

```
args(name = round)
```

```
## function (x, digits = 0)
## NULL
```

args prints a summary of the main arguments of a function. What can we learn from the summary of the arguments of round? Notice that the first one, x, is shown without an associated value, whereas the digits part of the summary is printed as digits = 0.

The significance of this is that digits has a default value (0 in this case). This means that we can leave out digits when using the round function:

```
round(x = 3.7)
```

```
## [1] 4
```

This is the same result as we would get using round(x = 3.7, digits = 0). This 'default value' behaviour is useful because it allows us keep our R code concise. Some functions take a large number of arguments, many of which are defined with sensible defaults. Unless we need to change these default arguments, we can ignore them when we call such functions.

Notice that the x argument of round does not have a default, which means we have to supply a value. This is sensible, as the whole purpose of round is to round any number we give it.

There is another way to simplify our use of functions. Take a look at this example:

```
round(3.72, digits = 1)
```

```
## [1] 3.7
```

What does this demonstrate? We do not have to specify argument names. In the absence of a name R uses the position of the supplied argument to work out which name to associate it with. In this example we left out the name of the argument at position 1. This is where x belongs, so we end up rounding 3.71 to 1 decimal place.

R is even more flexible than this. We don't necessarily have to use the full name of an argument, because R can use partial matching on argument

names:

```
round(3.72, dig = 1)
```

```
## [1] 3.7
```

This also works because R can unambiguously match the argument we named dig to digits.



Be careful with your arguments

Here is some advice. Do not rely on partial matching of function names. It just leads to confusion and the odd error. If you use it a lot, you end up forgetting the true name of arguments, and if you abbreviate too much, you create name matching conflicts. For example, if a function has arguments arg1 and arg2 and you use the partial name a for an argument, there is no way to know which argument you meant. We are pointing out partial matching so that you are aware of the behaviour. It is not worth the hassle of getting it wrong to save on a little typing, so do not use it.

What about position matching? This can also cause problems if we're not paying attention. For example, if you forget the order of the arguments to a function and then place your arguments in the wrong place, you will either generate an error or produce a nonsensical result. It is nice not to have to type out the name = value construct all the time though, so our advice is to rely on positional matching only for the first argument. This is a common convention in R that makes sense because it is often obvious what kind of information the first argument should carry.

2.5 Combining functions

Using R to do 'real work' usually involves linked steps, each facilitated by a different function. There is more than one way to achieve this. Here's a simple example that uses an approach we already know about—storing intermediate results:

```
y <- sqrt(10)
round(y, digits = 1)</pre>
```

```
## [1] 3.2
```

These two lines calculate the square root of the number 10 and assigned the result to y, then round this to one decimal place and print the result. We linked the two calculations by assigning a name to the first result and then used this as the input to a function in the second step.

Here is another way to replicate the same calculation:

```
round(sqrt(10), digits = 1)
```

[1] 3.2

The technical name for this is function composition or function nesting: the sqrt function is 'nested inside' the round function. The way we have to read these constructs is from the inside out. The sqrt(10) expression is inside the round function, so this is evaluated first. The result of sqrt(10) is then associated with the first argument of the round function, and only then does the round function do its job.

There aren't any new ideas here. We have already seen that R evaluates whatever is on the right-hand side of the = symbol first before associating the resulting value with the appropriate argument name.

We can extend this nesting idea to do more complicated calculations, i.e. there's nothing to stop us using multiple levels of nesting either. Take a look at this example:

```
round(sqrt(abs(-10)), digits = 1)
```

[1] 3.2

The abs function takes the absolute value of a number, i.e. removes the - sign if it is there. Remember, read nested calls from the inside out:

- 1. we find the absolute value of -10,
- 2. we calculate the square root of the resulting number (+10), and
- 3. then we rounded this to a whole number.

Nested function calls can be useful because they make R code less verbose (we write less), but this comes at a high cost of reduced readability. No reasonable person would say that round(sqrt(abs(-10)), digits = 1) is easy to read! For this reason, we aim to keep function nesting to a minimum. We will occasionally have to use the nesting construct, so it is important to understand how it works even if we don't like it.

The good news is that we'll see a much-easier-to-read method for applying a series of functions in the Data Wrangling block.

2.6 Functions do not have 'side effects'

We'll finish this chapter with an idea every R user needs to understand to avoid confusion. It relates to how functions modify their arguments, or more accurately, how they **do not** modify their arguments. Take a look at this example:

```
y <- 3.7
round(y, digits = 0)
## [1] 4
y</pre>
```

```
## [1] 3.7
```

We created a variable y with the value 3.7, rounded this to a whole number with round, printed out the result, and then printed the value of y. Notice that the value of y has not changed after using it as an argument to round.

This is important. R functions do not typically alter the values of their arguments (we say 'typically' because there are ways to alter this behaviour if we really want to). This behaviour is captured by the phrase 'functions do not have side effects'.

If we had intended to round the value of y so that we can use this new value later on, we have to assign the result of function evaluation, like this:

```
y <- 3.7
y <- round(y, digits = 0)
```

The reason for pointing out this behaviour is because new R users sometimes assume a function will change its arguments. R functions do not typically do this. If we want to make use of changes, rather than print them to the Console, we need to assign the result a name, either by creating a new variable or overwriting the old one. Remember—functions do not have side effects! Forgetting this creates all kinds of headaches.

Chapter 3

Vectors

3.1 Introduction

This chapter has three goals. First, we want to learn how to work with a vector, one of the basic structures used to represent data in R-land. Second, we'll learn how to work with vectors by using **numeric** vectors to perform simple calculations. Finally, we'll introduce a couple of different kinds of vectors—**character** vectors and **logical** vectors. This material provides a foundation for working with real data sets in the next block.



Data structures?

The term 'data structure' is used to describe conventions or rules for organising and storing data on a computer. Computer languages use many different kinds of data structures. Fortunately, we only need to learn about a couple of relatively simple ones to use R for data analysis: 'vectors' and 'data frames'. This chapter will consider vectors. The next chapter will look at collections of vectors (a.k.a. data frames).

3.2 Atomic vectors

We'll start with a definition, even though it probably won't make much sense yet: a **vector** is a 1-dimensional data structure for storing a set of values, each accessible by its position in the vector. The simplest kind of vectors in R are called **atomic vectors**¹.

There are different kinds of atomic vector, but their defining, common feature

 $^{^{1}\}mathrm{The}$ other common vector is called a "list". Lists are very useful but we won't cover them in this book.

is that it can only contain data of one 'type'. They might contain all integers (e.g. 2, 4, 6, ...) or all characters (e.g. "A", "B", "C"), but they can't mix and match integers and characters (e.g. "A", 2, "C", 5).

The word 'atomic' in the name refers to the fact that an atomic vector can't be broken down into anything simpler—they are the simplest kind of data structure R knows. Even when working with a single number we're actually dealing with an atomic vector. Here's the very first expression we evaluated in the Introduction to R chapter:

```
1 + 1
```

[1] 2

Look at the output. What is that [1] at the beginning? It's actually a clue that the output resulting from 1+1 is an atomic vector. We can verify this with the is.atomic functions. First, make a variable called x with the result of the 1+1 calculation:

```
x <- 1 + 1
x
```

[1] 2

Then use is.atomic to check whether x really is an atomic vector:

```
is.atomic(x)
```

[1] TRUE

Atomic vectors really are the simplest kind of data structure in R. Unlike many other languages, there is simply no way to represent just a number. Instead, a single number is always stored as a vector of length one².

3.3 Numeric vectors

A lot of work in R involves **numeric vectors**. After all, data analysis is all about numbers. Here's one simple way to construct a numeric vector (and print it out):

```
numvec <- numeric(length = 50)
numvec</pre>
```

²The same is true for things like sets of characters ("dog", "cat", "fish", ...) and logical values (TRUE or FALSE) discussed in the next two chapters.

What happened? We made a numeric vector with 50 **elements**, each of which is the number 0. The word 'element' is used to refer to the values that reside in a vector.

When we create a vector but don't assign it to a name using <- R just prints it for us. Notice what happens when a vector is printed to the screen. Since a length-50 vector can't fit on one line, it was printed over two. At the beginning of each line there is a [X]: the number X gives the position of the elements printed at the beginning of each line.

If we need to check that we really have made a numeric vector, we can use the is.numeric³ function to do this:

is.numeric(numvec)

[1] TRUE

This confirms numvec is numeric by returning TRUE (a value of FALSE would mean that numvec is some other kind of object).

Keep in mind that R won't always print the exact values of the elements of a vector. For example, when R prints a numeric vector, it only prints the elements to 7 significant figures by default. We can see this by printing the built in constant pi to the Console:

рi

[1] 3.141593

The actual value stored in pi is much more precise than this. We can see this by printing pi again using the print function:

```
print(pi, digits = 16)
```

[1] 3.141592653589793



Different kinds of numbers

Roughly speaking, R stores numbers in two different ways depending of whether they are whole numbers ("integers") or numbers containing decimal points ("doubles" – don't ask). We're not going to worry about this difference. Most of the time, the distinction is invisible to users, so it is easier to think in terms of numeric vectors. We can mix and match integers and doubles in R without having to worry about how it is storing the numbers.

³This may not look like the most useful function in the world, but sometimes we need functions like is.numeric to understand what R is doing or root out mistakes in our scripts.

3.4 Constructing numeric vectors

We just saw to make a numeric vector of zeros using the numeric function. This is arguably not a particularly useful skill because we usually need to work vectors of particular values (not just 0). A useful function for creating custom vectors is the c function. Take a look at this example:

```
c(1.1, 2.3, 4.0, 5.7)
```

```
## [1] 1.1 2.3 4.0 5.7
```

The 'c' in the function name stands for 'combine'. The c function takes a variable number of arguments, each of which must be a vector of some kind, and combines these into a single vector. We supplied the c function with four arguments, each of which was a single number (i.e. a length-one vector). The c function combines these to generate a vector of length 4. Simple.

Now look at this example:

```
vec1 <- c(1.1, 2.3)
vec2 <- c(4.0, 5.7, 3.6)
c(vec1, vec2)</pre>
```

```
## [1] 1.1 2.3 4.0 5.7 3.6
```

This shows that we can use the c function to combine two or more vectors of any length. We combined a length-2 vector with a length-3 vector to produce a new length-5 vector.



The c function is odd

Notice that we did not have to name the arguments in those two examples—there were no = involved. The c function is an example of one of those flexible functions that breaks the simple rules of thumb for using arguments that we set out earlier: it can take a variable number of arguments that do not have predefined names. This behaviour is necessary for c to be of any use: to be useful it needs to be flexible enough to take any combination of arguments.

3.5 Named vectors

What happens if use named arguments with c? Take a look at this:

```
namedv \leftarrow c(a = 1, b = 2, c = 3)
namedv
```

```
## a b c ## 1 2 3
```

What happened? R used the argument names to set the names of each element in the vector. The resulting vector is still a 1-dimensional data structure. When printed to the Console, each element's value is printed along with its associated name above it. We can extract the names from a named vector using the names function:

```
names (namedv)
```

```
## [1] "a" "b" "c"
```

Being able to name the elements of a vector is useful because it makes it easier to identify and extract the bits we need.

3.6 Vectorised operations

All the simple arithmetic operators (e.g. + and -) and many mathematical functions are **vectorised** in R. When we use a vectorised function it operates on vectors on an element-by-element basis. We'll make a couple of simple vectors to illustrate what we mean:

```
x <- c(1, 2, 3, 4, 5, 6, 7, 8, 9, 10)
x
```

```
## [1] 1 2 3 4 5 6 7 8 9 10
```

```
y <- c(0.1, 0.2, 0.3, 0.4, 0.5, 0.6, 0.7, 0.8, 0.9, 1.0)
y
```

```
## [1] 0.1 0.2 0.3 0.4 0.5 0.6 0.7 0.8 0.9 1.0
```

This constructed two length-10 numeric vectors, called \mathbf{x} and \mathbf{y} , where \mathbf{x} is a sequence from 1 to 10 and \mathbf{y} is a sequence from 0.1 to 1.0. \mathbf{x} and \mathbf{y} are the same length. Now look at what happens when we add these using +:

```
x + y
```

```
## [1] 1.1 2.2 3.3 4.4 5.5 6.6 7.7 8.8 9.9 11.0
```

When R evaluates the expression x + y it does this by adding the first element of x to the first element of y, the second element of x to the second element of y, and so on, working through all ten elements of x and y. That's what is meant by a **vectorised** operation.

Vectorisation is implemented in all the standard mathematical functions. For

example, the **round** function rounds each element of a numeric vector to the nearest integer by default:

round(y)

[1] 0 0 0 0 0 1 1 1 1 1

The same behaviour is seen with other mathematical functions like sin, cos, exp, and log—they apply the relevant function to each element of a numeric vector.

It is important to realise that not all functions are vectorised. For example, the sum function takes a vector of numbers and adds them up:

sum(y)

[1] 5.5

Although sum obviously works on a numeric vector it is not 'vectorised' in the sense that it works element-by-element to return an output vector of the same length as its main argument. It just returns a single number—the sum total of the elements of its input.



Vectorisation is not the norm

R's vectorised behaviour may seem like the obvious thing to do, but most computer languages do not work like this. In other languages, we typically have to write a much more complicated expression to do something so simple. This is one reason R is such a good data analysis language: vectorisation allows us to express repetitious calculations in a simple, intuitive way. This behaviour can save a lot of time.

3.7 Other kinds of atomic vectors

The data we collect and analyse are often in the form of numbers. It comes as no surprise, therefore, that we work with numeric vectors a lot in R. Nonetheless, we also need to use other kinds of vectors, either to represent different types of data, or to help us manipulate our data. This section introduces two new types of atomic vector to help us do this: character vectors and logical vectors.

3.7.1 Character vectors

Each element of a **character vectors** is what is known as a "character string" (or "string" if we are feeling lazy). That term "character string" refers to a sequence of characters, such as "Treatment 1", "University of Sheffield", "Pop-

ulation Density". A character vector is an atomic vector that stores an ordered collection of one or more character strings.

If we want to construct a character vector in R, we have to place double (") or single (') quotation marks around the characters. For example, we can print the name "Dylan" to the Console like this:

```
"Dylan"
```

```
## [1] "Dylan"
```

Notice the [1]. This shows that what we just printed is an atomic vector of some kind. We know it's a character vector because the output is printed with double quotes around the value. We often need to make simple character vectors containing only one value—for example, to set the values of arguments to a function.

The quotation marks are not optional—they tell R we want to treat whatever is inside them as a literal value. The quoting is important. If we try to do the same thing as above without the quotes, we end up with an error:

```
Dylan
```

```
## Error in eval(expr, envir, enclos): object 'Dylan' not found
```

What happened? When the interpreter sees the word Dylan without quotes it assumes that this must be the name of a variable, so it goes in search of it in the global environment. We haven't made a variable called Dylan, so there is no way to evaluate the expression and R spits out an error to let us know there's a problem.

Character vectors are typically constructed to represent data of some kind. The c function is one starting point for this kind of thing:

```
# make a length-3 character vector
my_name <- c("Dylan", "Zachary", "Childs")
my_name</pre>
```

```
## [1] "Dylan" "Zachary" "Childs"
```

Here we made a length-3 character vector, with elements corresponding to a first name, middle name, and last name. If we want to extract one or more elements from a character vector by their position

Take note, this is **not** equivalent to the above :

```
my_name <- c("Dylan Zachary Childs")
my_name</pre>
```

[1] "Dylan Zachary Childs"

This length-1 character vector's only element is a single character string containing the first, middle and last name separated by spaces. We didn't even need to use the the c function here because we were only ever working with a length-1 character vector. i.e. we could have typed "Dylan Zachary Childs" and we would have ended up with exactly the same text printed at the Console.

3.7.2 Logical vectors

The elements of **logical vectors** only take two values: TRUE or FALSE. Don't let the simplicity of logical vectors fool you. They're very useful. As with other kinds of atomic vectors, the **c** function can be used to construct a logical vector:

```
l_vec <- c(TRUE, FALSE)
l_vec</pre>
```

[1] TRUE FALSE

So why are logical vectors useful? Their allow us to represent the results of questions such as, "is x greater than y" or "is x equal to y". The results of such comparisons may then be used to carry out various kinds of subsetting operations.

Before we can look at how to use logical vectors to evaluate comparisons, we need to introduce **relational operators**. These sound fancy, but they are very simple: we use relational operators to evaluate the relative value of vector elements. Six are available in R:

- x < y: is x less than y?
- x > y: is x greater than y?
- $x \le y$: is x less than or equal to y?
- $x \ge y$: is x greater than or equal to y?
- x == y: is x = y?
- x != y: is x not equal to y?

The easiest way to understand how these work is by example. We need a couple of numeric variables first:

```
x <- c(11, 12, 13, 14, 15, 16, 17, 18, 19, 20)
y <- c(3, 6, 9, 12, 15, 18, 21, 24, 27, 30)
x
```

```
## [1] 11 12 13 14 15 16 17 18 19 20
```

у

```
## [1] 3 6 9 12 15 18 21 24 27 30
```

Now, if we need to evaluate and represent a question like, "is x greater than than y", we can use either < or >:

x > y

[1] TRUE TRUE TRUE TRUE FALSE FALSE FALSE FALSE FALSE

The x > y expression produces a logical vector, with TRUE values associated with elements in x are less than y, and FALSE otherwise. In this example, x is less than y until we reach the value of 15 in each sequence. Notice too that relational operators are vectorised: they work on an element by element basis.

What does the == operator do? It compares the elements of two vectors to determine if they are exactly equal:

x == y

[1] FALSE FALSE FALSE FALSE TRUE FALSE FALSE FALSE FALSE

The output of this comparison is true only for one element, the number 15, which is at the 5^{th} position in both x and y. The != operator is essentially the opposite of ==. It identifies cases where two elements are not exactly equal. We could step through each of the different relational operators, but hopefully, they are self-explanatory at this point (if not, experiment with them).

A

= and == are not the same

If we want to test for equivalence between the elements of two vectors we must use double equals (==), not single equals (=). Forgetting to use == instead of = is a very common source of mistakes. The = symbol already has a use in R—assigning name-value pairs—so it can't also be used to compare vectors because this would lead to ambiguity in our R scripts. Using = when you meant to use == is a very common mistake. If you make it, this will lead to all kinds of difficult-to-comprehend problems with your scripts. Try to remember the difference!

Chapter 4

Data frames

4.1 Introduction

The quick introduction to R chapter introduced the word 'variable' as a short-hand for a named object. For example, we can make a variable called num_vec that refers to a simple numeric vector using:

```
num_vec <- c(1.1, 2.3, 4.0, 5.7)
num_vec</pre>
```

```
## [1] 1.1 2.3 4.0 5.7
```

When a computer scientist talks about variables they're referring to these sorts of name-value associations.

However, the word 'variable' has a second, more abstract meaning in the world of statistics: it refers to anything we can control or measure. For example, data from an experiment will involve variables whose values describe the experimental conditions (e.g. low temperature vs. high temperature) and the quantities we chose to measure (e.g. enzyme activity). We refer to these kinds of variables as 'statistical variables'.

We'll discuss these statistical variables later on. We're pointing out the dual meaning of the word 'variable' now because we need to work with both interpretations. The duality can be confusing at times, but both meanings are in widespread use, so we just have to get used to them. We try to minimise confusion by using the phrase "statistical variable" when referring to data, rather than R objects.

We're introducing these ideas now because we're going to consider a new type of data object in this chapter—the **data frame**. Real-world data analysis involves collections of related statistical variables. How should we keep a large collection

of variables organised? We could work with them individually, but this tends to be error prone. Instead, we need a way to keep related variables together. This is the problem that **data frames** are designed to manage.

4.2 Data frames

Data frames are one of R's features that mark it out as particularly good for data analysis. We can think of a data frame as a table-like object with rows and columns. A data frame collects together different statistical variables, storing each of them as a separate column. Related observations are all found in the same row.

We'll think about the columns first.

Each column of a data frame is a vector of some kind. These are usually simple atomic vectors containing numbers or character strings, though it is possible to include more complicated vectors. The critical constraint that a data frame applies is that each vector must have the same length. This is what gives a data frame it table-like structure.

The simplest way to get a feel for data frames is to make one. We can make one 'by hand' using some artificial data describing a made-up experiment. Imagine we had conducted a small experiment to examine biomass and community diversity in six field plots. Three plots were subjected to fertiliser enrichment. The other three plots act as experimental controls.

We could store the data describing this experiment in three vectors:

- trt (short for "treatment") shows which experimental manipulation was used in a given plot.
- bms (short for "biomass") shows the total biomass measured at the end of the experiment.
- div (short for "diversity") shows the number of species present at the end of the experiment.

Here's some R code to generate these three vectors (it doesn't matter what the actual values are, they're made up):

```
trt <- rep(c("Control", "Fertilser"), each = 3)
bms <- c(284, 328, 291, 956, 954, 685)
div <- c(8, 12, 11, 8, 4, 5)</pre>
trt
```

```
## [1] "Control" "Control" "Fertilser" "Fertilser" "Fertilser"
```

```
bms
## [1] 284 328 291 956 954 685
div
```

```
## [1] 8 12 11 8 4 5
```

Notice that the information about different observations are linked by their positions in these vectors. For example, the third control plot had a biomass of '291' and a species diversity '11'.

We use the data.frame function to construct a data frame from one or more vectors, i.e. to build a data frame from the three vectors we just created:

```
experim.data <- data.frame(trt, bms, div)
experim.data</pre>
```

```
## trt bms div
## 1 Control 284 8
## 2 Control 328 12
## 3 Control 291 11
## 4 Fertilser 956 8
## 5 Fertilser 954 4
## 6 Fertilser 685 5
```

Notice what happens when we print the data frame: it is displayed as though it has rows and columns. That's what we meant when we said a data frame is a table-like structure.

The data.frame function takes a variable number of arguments. We used the trt, bms and div vectors, resulting in a data frame with three columns. Each of these vectors has 6 elements, so the resulting data frame has 6 rows. The names of the vectors were used to name its columns. The rows do not have names, but they are numbered to reflect their position.

The words trt, bms and div are not very informative. If we prefer to work with more meaningful column names—which is always a good idea—then we can name the data.frame arguments:

```
experim.data <- data.frame(Treatment = trt, Biomass = bms, Diversity = div)
experim.data</pre>
```

```
## Treatment Biomass Diversity
## 1 Control 284 8
## 2 Control 328 12
## 3 Control 291 11
```

```
## 4 Fertilser 956 8
## 5 Fertilser 954 4
## 6 Fertilser 685 5
```

The new data frame contains the same data as the previous one but now the column names correspond to the human-readable words we chose.



Don't bother with row names

We can also name the rows of a data frame using the row.names argument of the data.frame function. We won't bother to show an example of this, though. Why? We can't easily work with the information in row names which means there's not much point adding it. If we need to include row-specific information in a data frame it's best to include an additional variable, i.e. an extra column.

4.3 Exploring data frames

The first things to do when presented with a new data set is to explore its structure to understand what we're dealing with. There are plenty of options for doing this when the data are stored in a data frame. For example, the head and tail functions extract the first and last few rows of a data set:

```
head(experim.data, n = 3)
##
     Treatment Biomass Diversity
## 1
       Control
                    284
## 2
                    328
                                12
       Control
## 3
       Control
                    291
                                11
tail(experim.data, n = 3)
     Treatment Biomass Diversity
## 4 Fertilser
                    956
## 5 Fertilser
                    954
                                 4
## 6 Fertilser
                    685
                                 5
```

Notice that the n argument controls the number of rows printed. The View function can be used to open up the whole data set in a table- or spreadsheet-like like view:

```
View(experim.data)
```

Exactly what happens when we use View depends on how we're interacting with R. When we run it in RStudio a new tab opens up with the data shown inside

it.



View only displays the data

The View function is only designed to display a data frame as a table of rows and columns. We can't change the data in any way with the View function. We can reorder the way the data are presented, but keep in mind that this won't alter the underlying data.

There are quite a few different R functions that will extract information about a data frame. The nrow and ncol functions return the number of rows and columns, respectively:

```
nrow(experim.data)
```

[1] 6

ncol(experim.data)

[1] 3

The names function can be used to extract the column names from a data frame:

colnames(experim.data)

```
## [1] "Treatment" "Biomass" "Diversity"
```

The experim.data data frame has three columns, so names returns a character vector of length three, where each element corresponds to a column name.

4.4 Extracting and adding a single variable

Remember, each column of a data frame can be thought of as a variable. Data frames would not be much use if we could not extract and modify the variables they contain. In this section, we will briefly review how to extract a variable. We'll examine ways to manipulate the data within a data frame in later chapters.

One way of extracting a variable from a data frame uses a double square brackets construct, [[. For example, we extract the Biomass variable from our example data frame with the double square brackets like this:

```
experim.data[["Biomass"]]
```

```
## [1] 284 328 291 956 954 685
```

This prints whatever is in the Biomass column to the Console. What kind of object is this? It's a numeric vector:

```
is.numeric(experim.data[["Biomass"]])
```

[1] TRUE

See? A data frame is a collection of vectors. Notice that all we did was print the resulting vector to the Console. If we want to do something with this numeric vector we need to assign the result:

```
bmass <- experim.data$Biomass
bmass^2</pre>
```

[1] 80656 107584 84681 913936 910116 469225

Here, we extracted the Biomass variable, assigned it to bmass, and then squared this.

Notice that we used "Biomass" instead of Biomass inside the double square brackets, i.e. we quoted the name of the variable. This is because we want R to treat the word "Biomass" as a literal value. This little detail is important! If we don't quote the name, R will assume that Biomass is the name of an object and go in search of it in the global environment. Since we haven't created something called Biomass, leaving out the quotes would generate an error:

```
experim.data[[Biomass]]
```

```
## Error in (function(x, i, exact) if (is.matrix(i)) as.matrix(x)[[i]] else .subset2(x
```

The error message is telling us that R can't find a variable called Biomass.

The second method for extracting a variable uses the \$ operator. For example, to extract the Biomass column from experim.data, we use:

```
experim.data$Biomass
```

```
## [1] 284 328 291 956 954 685
```

We use the \$ operator by placing the name of the data frame we want to work with on the left-hand side and the name of the column (i.e. the variable) we want to extract on the right-hand side. Notice that we didn't have to put quotes around the variable name when using the \$ operator. We can do this if we want to—i.e. experim.data\$"Biomass" also works—but \$ doesn't require it



Why is there more than one way to extract variables?

There's no simple way to answer this question without getting into the details of how R represents data frames. The simple answer is that \$ and [[are not strictly equivalent, even though they appear to do much the same thing. The \$ method is a bit easier to read, and people tend to prefer it for interactive data analysis tasks, whereas the [[construct tends to be used when we need a bit more flexibility for programming.

Chapter 5

Packages

5.1 The R package system

The R package system is the most important single factor driving increased adoption of R. Packages are used to extend the basic capabilities of R. In his book about R packages Hadley Wickam says,

Packages are the fundamental units of reproducible R code. They include reusable R functions, the documentation that describes how to use them, and sample data.

An R package is a collection of folders and files in a standard, well-defined format that bundles together computer code, data, and documentation in a way that is easy to use and share with other users. The computer code might all be R code, but it can also include code written in other languages. Packages provide an R-friendly interface to use this "foreign" code without needing to understand how it works.

The base R distribution it comes with quite a few pre-installed packages. These base R packages represent a tiny subset of all available R packages. The majority of these are hosted on a worldwide network of web servers collectively know as CRAN: the Comprehensive R Archive Network, pronounced either "see-ran" or "kran".

CRAN is a fairly spartan web site, so it's easy to navigate. The landing page has about a dozen links on the right-hand side. Under the *Software* section there is a link called Packages. Near the top of that packages page there is a link called Table of available packages, sorted by name that points to a very long list of all the packages on CRAN. The column on the left shows each package name, followed by a brief description of what the package does on the right. There are 1000s of packages listed there.

5.2 Task views

The huge list of packages on the available packages is pretty overwhelming. A more user-friendly view of many R packages is found on the Task Views page (the link is on the left hand side, under the section labelled CRAN). A Task View is basically a curated guide to the packages and functions that are useful for certain disciplines. The Task Views page shows a list of these discipline-specific topics, along with a brief description. For example—

- The Environmentrics Task View contains information about using R to analyse ecological and environmental data.
- The Clinical Trials Task View contains information about using R to design, monitor, and analyse data from clinical trials.
- The Medical Image Analysis Task View contains information about packages for working with commercial medical image data.
- The Pharmacokinetic Task View contains information about packages for working with pharmacokinetic (PK) data.

Task views are often a good place to start looking for a new package to support a particular analysis in future projects.

5.3 Using packages

Two things need to happen to make use of a package. First, we need to copy the folders and files that make up the package to an appropriate location on our computer. This process is called **installing** the package. Second, we need to **load and attach** the package for use in an R session. The word "session" refers to the time between when we start up R and close it down again.

It's worth unpacking these two ideas because packages are a very frequent source of confusion for new users:

- If we don't have a copy of a package's folders and files our computer, we can't use it. The process of making this copy is called **installing** the package. It's possible to manually install packages by going to the CRAN website, downloading the package, and then using various tools to install it. We don't recommend using this approach though, because it's inefficient and error-prone. Instead, use built-in R functions to grab the package from CRAN and install it one step.
- Once we have a copy of the package on our computer, it will remain there for us to use. We don't need to re-install a package we want to use every time we start a new R session. It is worth saying that again, there is no need to install a package every time we start up R / RStudio. The only exception to this rule is that a major update to R will sometimes require a complete re-install of the packages. Such major updates are infrequent.

• Installing a package does nothing more than place a copy of the relevant files on our hard drive. If we want to use the functions or the data that comes with a package we need to make them available in our current R session. Unlike package installation, this **load and attach** process as it's known has to be repeated every time we restart R. If we forget to load up the package we can't use it.

5.3.1 Viewing installed packages

We sometimes need to check whether a package is currently installed. RStudio provides a simple, intuitive way to see which packages are installed on our computer. The **Packages** tab in the bottom right pane shows the name of every installed package, a brief description and a version number.

There are also a few R functions that can be used to check whether a package is currently installed. For example, the find.package function can do this:

```
find.package("MASS")
```

[1] "/Library/Frameworks/R.framework/Versions/4.0/Resources/library/MASS"

The find.package function either prints a "file path" showing us where the package is located, as above, or return an error if the package can't be found. Alternatively, a function called installed.packages will return a data frame containing a lot of information about the installed packages.

5.3.2 Installing packages

R packages can be installed from several different sources. For example, they can be installed from a local file on a computer, from the CRAN repository, or from an other kind of online repository called Github. Although various alternatives to CRAN are becoming more popular, we're only going to worry about installing packages that live on CRAN.

To install a package from an online repository like CRAN we have to down-load the package files, uncompress them (like we would a ZIP file), and move them to the correct location. All of this can be done using a single function: install.packages. For example, to install a package called **fortunes** we use:

```
install.packages("fortunes")
```

The quotes are necessary by the way. If everything is working—we have an active internet connection, the package name is valid, and so on—R will briefly pause while it communicates with the CRAN servers, we should see some red text reporting back what's happening, and then we're returned to the prompt. The red text is just letting us know what R is up to. As long as this text does not include the word "error", there is usually no need to worry about it.

There are a couple of things to keep in mind. First, package names are case sensitive. For example, **fortunes** is not the same as **Fortunes**. Quite often package installations fail because we used the wrong case somewhere in the package name. The other aspect of packages we need to know about is related to **dependencies**: some packages rely on other packages in order to work properly. By default <code>install.packages</code> will install these dependencies, so we don't usually have to worry too much about them. Just don't be surprised if the <code>install.packages</code> function installs more than one package when only one was requested.

RStudio provides a way of interacting with install.packages via point-and-click. The **Packages** tab has an "Install" button at the top right. Clicking on this brings up a small window with three main fields: "Install from", "Packages", and "Install to Library". We only need to work with the "Packages" field—the other two can be left at their defaults. When we start typing in the first few letters of a package name (e.g. **dplyr**) RStudio provides a list of available packages that match this. After we select the one we want and click the "Install" button, RStudio invokes install.packages for us.

5.3.3 Loading and attaching packages

Once we've installed a package or two, we'll probably want to use them. Two things have to happen to access a package's facilities: the package has to be loaded into memory, and then it has to attached to something called a search path so that R can find it.

It is beyond the scope of this book to get in to "how" and "why" of these events. Fortunately, there's no need to worry about these details, as both loading and attaching can be done in a single step with a function called library. The library function works as we might expect it to. If we want to start using the fortunes package—which was just installed above—all we need is:

```
library("fortunes")
```

Nothing much happens if everything is working as it should. R just returns us to the prompt without printing anything to the Console. The difference is that now we can use the functions that **fortunes** provides. As it turns out, there is only one function, called fortune:

```
##
## Friends don't let friends use Excel for statistics!
## -- Jonathan D. Cryer (about problems with using Microsoft Excel for statistics)
## JSM 2001, Atlanta (August 2001)
```

The **fortunes** package is either very useful or utterly pointless, depending on one's perspective. It dispenses quotes from various R experts delivered to the venerable R mailing list.



5.3.4 Don't use RStudio for loading packages!

As usual, if we don't like working in the Console RStudio can help us out. There is a small button next to each package listed in the **Packages** tab. Packages that have been loaded and attached have a blue check box next to them, whereas this is absent from those that have not. Clicking on an empty check box will load up the package. We mention this because at some point most people realise they can use RStudio to load and attach packages. We don't recommend using this route. It's much better to put library statements into your R script. Read the relevant appendix if you're not sure what a script is yet.

5.3.5 An analogy

The package system frequently confuses new users. This stems from the fact that they aren't clear about what the <code>install.packages</code> and <code>library</code> functions are doing. One way to think about these is by analogy with smartphone "Apps". Think of an R package as analogous to a smartphone App— a package effectively extends what R can do, just as an App extends what a phone can do.

When we want to use a new App, we download it from the App store and install it on our phone. Once downloaded, the App lives permanently on the phone and can be used whenever it's needed. Downloading and installing the App is something we only have to do once. Packages are no different. When we want to use an R package we first have to make sure it is installed on the computer (e.g. using install.packages). Installing a package is a 'do once' operation. Once installed, we don't need to install a package again each time we restart R.

To actually use an App on our phone we open it up by tapping on its icon. This has to happen every time we want to use the App. The package equivalent of opening a smartphone App is the "load and attach" operation. This is what library does. It makes a package available for use in a particular session. We have to use library to load the package every time we start a new R session if we plan to access the functions in that package: loading and attaching a package via library is a "do every time" operation.

5.4 Package data

Remember what Hadley Wickam said about packages? "... include reusable R functions, the documentation that describes how to use them, **and sample data**." Many packages include sample data sets for use in examples and package

vignettes. Use the data function to list the data sets hiding away in packages:

```
data(package = .packages(all.available = TRUE))
```

The mysterious .packages(all.available = TRUE) part of this generates a character vector with the names of all the installed packages in it. If we only use data() then R only lists the data sets found in a package called datasets and any additional packages we have loaded in the current R session.

The datasets package is part of the base R distribution. It exists for one reason—to store example data sets. The datasets package is automatically loaded when we start R, i.e. there's no need to use library to access it, meaning any data stored in this package can be accessed every time we start R.

From the perspective of learning to use R, working with package data is really useful because it allows us to work with 'well-behaved' data sets without having to worry about getting them into R. Importing data is certainly an important skill but it's not necessarily something a new user wants to worry about. For this reason, we tend to use package data sets in this book.

5.5 The tidyverse ecosystem of packages

We're going to be using several packages that belong to a widely used, well-known ecosystem of packages known as the **tidyverse**. Here is the description of the tidyverse on its website:

The tidyverse is an opinionated collection of R packages designed for data science. All packages share an underlying design philosophy, grammar, and data structures.

Using the tidyverse makes data analysis simpler, faster and a more entertaining experience (honestly).

Part II Data Wrangling

Chapter 6

Getting ready to use dplyr

6.1 Introduction

Data wrangling is the process of cleaning and manipulating data to get it ready for analysis, for example, by creating derived variables and subsets of the data. Although not the most exciting part of a study—we want to answer questions, not format data—data wrangling is critical. This step in a data analysis workflow can become very time-consuming if not tackled with the right tools.

The next few chapters will introduce the **dplyr** package. **dplyr** is an important member of the tidyverse ecosystem. Its job is to provide a set of tools to address common data manipulation tasks, such as selecting subsets of data, making new variables, and summarising data in various ways.

However, before we can start to use these tools, we need a bit of a foundation. That is the aim of this chapter. We will provide some background to the so-called tidy data principles, introduce the data set we'll be using in our examples, discuss the tidyverse version of data frames known as the tibble, and finally, introduce the **dplyr** package.

6.2 Tidy data

dplyr will work with any data frame. However, it is most powerful when data are organised according to tidy data conventions for rectangular data sets. Tidy data has a specific structure that makes it easy to manipulate, model and visualise. A tidy data set is one where each variable is only found in one column and each row contains one unique observation (an imaged cell, a treated organism, an experimental plot, and so on).

The basic principles of tidy data are not too difficult to understand. We'll use an example to illustrate what the "one variable = one column" and "one

observation = one row" idea means. Let's return to the made-up experiment investigating the response of communities to fertilizer addition. This time, imagine we had only measured biomass, but that we had measured it at two time points throughout the experiment.

We'll look at two ways to organise some artificial data from this experimental setup. The first uses a separate column for each biomass measurement:

##		Treatment	BiomassT1	${\tt BiomassT2}$
##	1	Control	284	324
##	2	Control	328	400
##	3	Control	291	355
##	4	Fertilser	956	1197
##	5	${\tt Fertilser}$	954	1012
##	6	Fertilser	685	859

This feels like a reasonable way to store such data, especially for an Excel user. However, this format is **not tidy**. Why? The biomass variable has been split across two columns (BiomassT1 and BiomassT2), which means each row corresponds to two distinct observations. We won't go into the 'whys' here but take our word for it—adopting this format makes it difficult to use **dplyr** efficiently.

A tidy version of that example data set would still have three columns but now these would be: Treatment, denoting the experimental treatment applied; Time, denoting the sampling occasion; and Biomass, denoting the biomass measured:

##		${\tt Treatment}$	${\tt Time}$	${\tt Biomass}$
##	1	Control	T1	284
##	2	Control	T1	328
##	3	Control	T1	291
##	4	${\tt Fertilser}$	T1	956
##	5	${\tt Fertilser}$	T1	954
##	6	${\tt Fertilser}$	T1	685
##	7	Control	T2	324
##	8	Control	T2	400
##	9	Control	T2	355
##	10	Fertilser	T2	1197
##	11	Fertilser	T2	1012
##	12	Fertilser	T2	859

The change we made was to create an indicator variable called Time for the sampling occasion. In version one of the data, the time information was implicit—the time associated with a biomass measurement was encoded by column membership (BiomassT1 vs BiomassT2). In the second version of the data set an indicator variable, Time, was created to label the sampling occasion explicitly. That simple change means each row corresponds to one distinct observation and each variable lives in one column. These data are now tidy and ideally set up for use with dplyr.



Always try to start with tidy data

The best way to make sure your data set is tidy is to store in that format when it's first collected and recorded. Some packages can help convert non-tidy data into the tidy data format (e.g. the **tidyr** package), but life is much simpler if we ensure our data are tidy from the very beginning.

6.3 Penguins! +=

To make progress with **dplyr**, we are going to need some data to play around with. The data we'll use contains measurements taken from penguins on the Palmer Archipelago, off the north-western coast of the Antarctic Peninsula. Each row contains information about an individual penguin, including:

- the species it belongs to,
- the island it was found on,
- morphometric data (flipper length, body mass, bill dimensions)
- its body mass and sex,
- and finally, the year of capture.



Figure 6.1: Meet the Palmer penguins (artwork by Allison Horst)

Why use this data set? Apart from the fact that everyone likes penguins, obviously, these data are sufficiently complex to demonstrate everything we want to do, while remaining easily understandable. Here is the full data set shown as a table:

N.B. — the data will only show up in the HTML version of the book.

This shows the first ten rows along with the first few columns. The *Previous / Next* links at the bottom right can be used to navigate the rows; the arrows in

the top header row are used to view the different columns. Below each column name you can also see three-letter abbreviations like <dbl>, <int> and <chr>>. These are telling us that each column is a vector of some kind:

- <dbl> = a numeric 'double' vector (real numbers, i.e. with decimals)
- <int> = a numeric integer vector (integer numbers)
- <chr> = character vector.

So... data frames are table-like objects with rows and columns. They can also be seen in even simpler terms—data frames are simply collections of vectors, where each one represents a different aspect of a multi-faceted data set.

We're going to be seeing a lot of this data set in the remaining chapters so we won't say anything more about it now.



Where can we get the penguins data?

The Palmer penguins data were collected and made available by Dr. Kristen Gorman and the Palmer Station. The data set exists mostly to support people learning and teaching R. It isn't part of base R, though. We have to import it into R somehow. There are two options:

- 1. The data are available in an R package called... palmerpenguins. Like other packages, **palmerpenguins** can be installed from CRAN using either install.packages or the usual RStudio point-and-click mechanism. Once installed, we can make it available by running library(palmerpenguins) in any R session. After that, just type the name of the data set and R will find it and use it. The version in **palmerpenguins** is called **penguins**, by the way.
- 2. We could get a copy of the data set and store it as a file on our hard drive, ideally using a standard data format. The most common and portable format for storing rectangular data is as a 'Comma Separated Value' (CSV) text file. Once you have a copy of the data as a CSV file, it's just a matter of using a function like read_csv from the readr package to import the data into R. This is explained in the Managing projects, scripts and data files appendix.

6.4 Um... tibbles?

To increase the scope of what the tidyverse can do, its makers created a special kind of data object known as a 'tibble'. This is meant to sound like the word 'table' pronounced by a New Zealander. We're not lying—the person who started the tidyverse is from New Zealand. Its name is a clue that a tibble is a table-like object, i.e. a rectangular data structure similar to a data frame.

In fact, the easiest way to conceptualise a tibble is as a special data frame with a few extra whistles and bells. More often than not, it's not necessary to pay attention to whether we're working with an ordinary data frame or a tidyverse tibble—we can often treat them as though they are interchangeable.

That said, there are exceptions to this rule of thumb, and we do occasionally need to work out which one we're using. A simple way to do this is by printing the data object. Imagine that we've imported the Palmer penguins into R and stored it in a tibble called **penguins**. This is what that would look like if we printed it at the Console:

penguins

```
## # A tibble: 344 x 8
##
      species island
                          bill_length_mm bill_depth_mm flipper_length_mm body_mass_g
      <chr>
##
               <chr>>
                                   <dbl>
                                                   <dbl>
                                                                      <int>
                                                                                   <int>
##
    1 Adelie
               Torgersen
                                     39.1
                                                    18.7
                                                                        181
                                                                                    3750
    2 Adelie
               Torgersen
                                     39.5
                                                    17.4
                                                                        186
                                                                                    3800
    3 Adelie
               Torgersen
                                     40.3
                                                    18
                                                                        195
                                                                                    3250
##
##
    4 Adelie
               Torgersen
                                    NA
                                                    NA
                                                                         NA
                                                                                      NA
                                                    19.3
                                                                        193
##
    5 Adelie
               Torgersen
                                     36.7
                                                                                    3450
    6 Adelie
               Torgersen
                                     39.3
                                                    20.6
                                                                        190
                                                                                    3650
##
    7 Adelie
               Torgersen
                                    38.9
                                                    17.8
                                                                        181
                                                                                    3625
    8 Adelie
                                     39.2
                                                    19.6
                                                                        195
                                                                                    4675
               Torgersen
                                                    18.1
                                                                        193
                                     34.1
    9 Adelie
               Torgersen
                                                                                    3475
## 10 Adelie
               Torgersen
                                     42
                                                    20.2
                                                                        190
                                                                                    4250
## # ... with 334 more rows, and 2 more variables: sex <chr>, year <int>
```

The formatting applied to the output is telling us penguins is a tibble rather than an ordinary data frame (we only called it a data frame earlier because we had not introduced the concept of a tibble yet). There are a couple of clues about this in that printout. The very obvious one is the first line: # A tibble: 344 x 8. The printed output states penguins is a tibble! The output is also truncated—only the first ten lines were printed, and any columns that won't fit on one row are summarised at the bottom (i.e. sex and year).

6.5 Missing values

Take a close look at the values of the body mass data (body_mass_g) in penguins (we're using \$ to extract the whole column):

```
penguins$body_mass_g
```

```
## [1] 3750 3800 3250 NA 3450 3650 3625 4675 3475 4250 3300 3700 3200 3800 4400
## [16] 3700 3450 4500 3325 4200 3400 3600 3800 3950 3800 3800 3550 3200 3150 3950
## [31] 3250 3900 3300 3900 3325 4150 3950 3550 3300 4650 3150 3900 3100 4400 3000
```

```
##
    [46] 4600 3425 2975 3450 4150 3500 4300 3450 4050 2900 3700 3550 3800 2850 3750
                   3600 4050 2850 3950 3350 4100 3050 4450 3600 3900
##
    [61]
        3150
             4400
                                                                      3550 4150
                                                                                3700
         4250 3700 3900 3550 4000 3200 4700 3800 4200 3350 3550 3800 3500 3950 3600
##
##
        3550
              4300
                   3400 4450 3300 4300 3700 4350 2900 4100 3725
                                                                4725
                                                                      3075 4250
##
   Г1061
        3550
             3750
                   3900 3175 4775 3825 4600 3200 4275 3900 4075 2900
                                                                      3775
                                                                           3350
                                                                                3325
   Γ121
        3150 3500
                   3450 3875 3050 4000 3275 4300 3050 4000 3325 3500
                                                                      3500 4475
                                                                                3425
        3900 3175
                  3975 3400 4250 3400 3475 3050 3725 3000 3650 4250 3475 3450
   [136]
   Γ151]
        3700 4000 4500 5700 4450 5700 5400 4550 4800 5200 4400 5150 4650 5550
   [166]
        5850 4200 5850 4150 6300 4800 5350 5700 5000 4400 5050 5000 5100 4100
        4600 5550 5250 4700 5050 6050 5150 5400 4950 5250 4350 5350 3950 5700 4300
   [181]
   Г196Т
        4750 5550 4900 4200 5400 5100 5300 4850 5300 4400 5000 4900 5050 4300 5000
   [211]
        4450
             5550
                  4200 5300 4400 5650 4700 5700 4650 5800 4700 5550 4750 5000 5100
   [226]
        5200
              4700
                   5800
                        4600 6000 4750 5950 4625
                                                 5450 4725 5350
                                                                4750
                                                                      5600
                                                                           4600
                   4950 5400 4750 5650 4850 5200 4925 4875 4625 5250
   [241]
        4875 5550
                                                                      4850 5600 4975
   [256]
              4725
                   5500 4700 5500 4575 5500 5000 5950 4650 5500 4375
                                                                      5850 4875
##
   [271]
        4925
                NA
                   4850 5750 5200 5400 3500 3900 3650 3525 3725
                                                                3950 3250 3750
                                                                                4150
        3700 3800
                        3700 4050 3575
                                       4050
                                            3300 3700 3450 4400
   [286]
                   3775
                                                                 3600
                                                                      3400 2900
   [301]
        3300 4150 3400 3800 3700 4550 3200 4300 3350 4100 3600 3900 3850 4800 2700
   [316] 4500 3950 3650 3550 3500 3675 4450
                                            3400 4300 3250 3675 3325 3950 3600 4050
  [331] 3350 3450 3250 4050 3800 3525 3950 3650 3650 4000 3400 3775 4100 3775
```

The body mass information lives in a numeric (integer) vector, but not every element in that vector is a number. Two values are NA. That stands for 'Not Available'—the NA's job is to label cases where a value is missing or unknown. If you scroll around the table view of penguins above you'll find NA's in several of the columns.

Missing data crop up all the time. They are just one of those facts of life—maybe the recording machine broke one afternoon, perhaps one of our organisms was infected by a pathogen, or maybe a cow ate one of our experimental plots. One of the nice things about R is that it knows how to represent missing data. It does this using the NA symbol.

You need to be aware of missing values when they are present. Why? Because the behaviour of many functions is affected by the presence of NAs. Things get confusing if we don't understand that behaviour. We will see some examples of this in the next few chapters.

6.6 Introducing dplyr

The **dplyr** package has been carefully designed to make it easy to manipulate 'rectangular data', such as data frames. **dplyr** is very consistent in the way its functions work. For example, the first argument of the main **dplyr** functions is always an object containing our data. This consistency makes it very easy to get to grips with each of the main **dplyr** functions—it's often possible to understand how one works by seeing one or two examples of its use.

Another reason for using **dplyr** is that it is orientated around a few core functions, each designed to do one thing well. These **dplyr** functions are sometimes referred to as its 'verbs' to reflect the fact that they 'do something' to data. For example:

- select is obtains a subset of variables,
- mutate is constructs new variables,
- filter is obtains a subset of rows,
- arrange is reorders rows, and
- summarise is calculates information about groups.

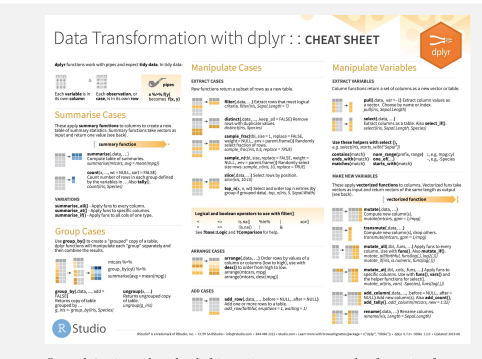
Notice that the names are chosen to reflect what each verb/function does to the input data. We'll cover each of these verbs in detail in later chapters, as well as a few additional ones, such as rename and group_by.

Apart from being easy to use, **dplyr** is also fast. This doesn't matter for small data sets but can be important when working with data sets with hundreds of thousands of rows. The **dplyr** package also allows us to work with data stored in different ways, for example, by interacting directly with several database systems. We're going to focus on using it with data frames and tibbles. But remember—once you know how to use **dplyr** with data frames its easy to use it for work with other kinds of data sources.



A dplyr cheat sheet

The developers of RStudio have produced a very usable cheat sheat that summarises the main data wrangling tools provided by **dplyr**.



Our advice is to download this, print out a copy, and refer to it often as you start working with **dplyr**.

6.6.1 A first look at dplyr

Let's take a preliminary look at the **dplyr** package. The package is not part of the base R installation, so we have to have installed it first. Remember, once installed, there is no need to install the package every time we need to use it. We do have to use library to load and attach the package every time we want to use it:

```
library("dplyr")
```

Let's look at one handy **dplyr** function now. Sometimes we just need a quick, compact summary of a data frame or tibble. This is the job of the **glimpse** function from **dplyr**:

```
glimpse(penguins)
```

The function takes one argument: the name of a data frame or tibble. It then tells us how many rows it has, how many columns there are, what these columns are called, and what type of data is associated with them. This function is useful when we need a quick overview of what's inside our data set. Some advice—use this function on any new data set before trying to do anything with it.

6.6.2 dplyr pseudocode

We'll wrap-up this chapter with a quick mention of 'pseudocode'. Pseudocode uses structural conventions of normal R code but is intended for us (humans) to read rather than the computer. We use it summarise how **dplyr** functions work. Here's an example you will encounter in the next chapter:

```
select(<data>, <variable-1>, <variable-2>, ...)
```

This is not an example we can run. It is pseudocode that serves as a template for using the dplyr select function. The words surrounded by 'angle brackets' (< >) are placeholders. If we want to turn this pseudocode into real R code we have to replace those placeholders. For example, <data> is a placeholder for the name of a data set. We would replace that with the word penguins if we want to use select on our example data set (we'd also have to do something with the other placeholders (e.g. <variable-1>)—we'll get to that later).

That's enough about pseudocode and **dplyr** templates. We'll see plenty of these in the next few chapters.

Chapter 7

Working with variables

7.1 Introduction

This chapter will explore the **dplyr** select and mutate verbs, and the closely related rename and transmute verbs. We consider these functions together because they operate on the variables (i.e. the columns) of a data frame or tibble:

- The select function selects a subset of variables to retain and (optionally) renames them in the process.
- The mutate function creates new variables from pre-existing ones and retains the original variables.
- The rename function renames one or more variables while keeping the remaining variable names unchanged.
- The transmute function creates new variables from pre-existing ones and drops the original variables.

7.1.1 Getting ready

Obviously, we need to have first installed **dplyr** package to use it. Assuming that's been done, we need to load and attach the package in the current session:

library("dplyr")

We will use the Palmer penguins data to illustrate the ideas in this chapter. Remember—the previous chapter described this data set and explained where to find it. The examples below assume it was read into R as a tibble with the name penguins.

7.2 Subset variables with select

We use select to select variables from a data frame or tibble. This is used when we have a data set with many variables but only need to work with a subset of these. Basic usage of select looks like this:

```
select(<data>, <variable-1>, <variable-2>, ...)
```

Remember—this is not an example we can run. This is a pseudocode designed to provide a generic description of how we use **select**. Let's look at the arguments of select:

- The first argument, <data>, must be the name of the object containing our data (usually a data frame or tibble). This is not optional—dplyr functions only exist to manipulate data.
- We then include a series of one or more additional arguments, where each one is the name of a variable in <data>. We've expressed this as <variable-1>, <variable-2>, ..., where <variable-1> and <variable-2> are names of the first two variables. The ... is acting as a placeholder for the remaining variables. There could be any number of these.

It's easiest to understand how a function like select works by seeing it in action. We select the species, bill_length_mm and bill_depth_mm variables from penguins like this:

```
select(penguins, species, bill_length_mm, bill_depth_mm)
```

```
## # A tibble: 344 x 3
##
      species bill_length_mm bill_depth_mm
##
      <chr>>
                         <dbl>
                                        <dbl>
    1 Adelie
                          39.1
                                         18.7
##
##
    2 Adelie
                          39.5
                                         17.4
##
    3 Adelie
                          40.3
                                         18
##
    4 Adelie
                          NA
                                         NA
    5 Adelie
                          36.7
                                         19.3
##
##
    6 Adelie
                          39.3
                                         20.6
##
    7 Adelie
                          38.9
                                         17.8
    8 Adelie
                          39.2
                                         19.6
##
   9 Adelie
                          34.1
                                         18.1
## 10 Adelie
                          42
                                         20.2
## # ... with 334 more rows
```

Hopefully, nothing about this example is too surprising. However, there are a few subtleties buried in that example:

• The select function is designed to work in a non-standard way which means variable names should **not** be surrounded by quotes. The one

exception is when a name has a space in it. Under those circumstances, it has to be quoted with backticks, e.g. `variable 1`.

- The select function does not have 'side effects'. This means is that it does not change the original penguins object. We printed the result produced by select to the Console, so we can't access the modified data set. If we need to use the result, we have to assign it a name using <-.
- The order of variables (i.e. the column order) in the resulting object is the same as the order in which they were supplied to the argument list. This means we can reorder variables at the same time as selecting them if we need to.
- The select function will return the same kind of data object we give it to work on. It returns a data frame if our data was in a data frame and a tibble if it was a tibble. In this example, R prints a tibble because penguins was a tibble.

That second point is important—we have to remember to assign the result a name using <- if we want to keep it and use it later. For example, we might call the result of that last example penguins_bill:

```
penguins_bill <- select(penguins, species, bill_length_mm, bill_depth_mm)</pre>
```

Now that we've named the new data set created by **select** we can refer to it by that name whenever we need it:

penguins_bill

```
## # A tibble: 344 x 3
##
      species bill_length_mm bill_depth_mm
##
      <chr>
                        <dbl>
                                       <dbl>
##
    1 Adelie
                         39.1
                                        18.7
    2 Adelie
                         39.5
                                        17.4
##
##
                         40.3
    3 Adelie
                                        18
    4 Adelie
                                        NA
                         NA
    5 Adelie
##
                         36.7
                                        19.3
    6 Adelie
                         39.3
                                        20.6
  7 Adelie
                         38.9
                                        17.8
## 8 Adelie
                         39.2
                                        19.6
## 9 Adelie
                         34.1
                                        18.1
## 10 Adelie
                                        20.2
## # ... with 334 more rows
```

Remember—the original penguins data is completely unchanged:

penguins

A tibble: 344 x 8

##		species	island	bill_leng	th_mm	bill_depth_m	m flippe	r_length_mm	body_mass_g
##		<chr></chr>	<chr></chr>		<dbl></dbl>	<dbl< td=""><td>></td><td><int></int></td><td><int></int></td></dbl<>	>	<int></int>	<int></int>
##	1	Adelie	Torgersen		39.1	18.	7	181	3750
##	2	Adelie	Torgersen		39.5	17.	4	186	3800
##	3	Adelie	Torgersen		40.3	18		195	3250
##	4	Adelie	Torgersen		NA	NA		NA	NA
##	5	Adelie	Torgersen		36.7	19.	3	193	3450
##	6	Adelie	Torgersen		39.3	20.	6	190	3650
##	7	Adelie	Torgersen		38.9	17.	8	181	3625
##	8	Adelie	Torgersen		39.2	19.	6	195	4675
##	9	Adelie	Torgersen		34.1	18.	1	193	3475
##	10	Adelie	Torgersen		42	20.	2	190	4250
##	#	with	334 more r	rows, and	2 more	variables:	sex <chr< td=""><td>>, year <in< td=""><td>t></td></in<></td></chr<>	>, year <in< td=""><td>t></td></in<>	t>

7.2.1 Alternative ways to identify variables with select

It's sometimes more convenient to use select to subset variables by specifying those we do **not** need, rather than specifying of the ones to keep. We can use the ! operator to indicate that certain variables should be dropped. For example, to get rid of the bill_depth_mm and bill_length_mm columns, we could use:

```
select(penguins, !bill_depth_mm, !bill_length_mm)
```

```
## # A tibble: 344 x 8
                         bill_length_mm flipper_length_mm body_mass_g sex
##
      species island
                                                                                   year
##
      <chr>
               <chr>
                                   <dbl>
                                                       <int>
                                                                    <int> <chr>
                                                                                  <int>
##
    1 Adelie
              Torgersen
                                    39.1
                                                         181
                                                                     3750 male
                                                                                   2007
##
    2 Adelie
                                    39.5
                                                         186
              Torgersen
                                                                     3800 female
                                                                                   2007
##
    3 Adelie
              Torgersen
                                    40.3
                                                         195
                                                                     3250 female
                                                                                   2007
##
    4 Adelie
              Torgersen
                                    NA
                                                          NA
                                                                       NA <NA>
                                                                                   2007
##
    5 Adelie
              Torgersen
                                    36.7
                                                         193
                                                                     3450 female
                                                                                   2007
##
    6 Adelie
               Torgersen
                                    39.3
                                                         190
                                                                     3650 male
                                                                                   2007
##
    7 Adelie
               Torgersen
                                    38.9
                                                         181
                                                                     3625 female
                                                                                   2007
##
    8 Adelie
               Torgersen
                                    39.2
                                                         195
                                                                     4675 male
                                                                                   2007
##
                                    34.1
                                                         193
                                                                                   2007
    9 Adelie
               Torgersen
                                                                     3475 <NA>
## 10 Adelie
               Torgersen
                                    42
                                                         190
                                                                     4250 <NA>
                                                                                   2007
## # ... with 334 more rows, and 1 more variable: bill_depth_mm <dbl>
```

This returns a tibble with all the other variables: species, island, flipper_length_mm, body_mass_g, sex and year.

The select function can also be used to grab (or drop) a set of variables that occur in a sequence next to one another. We specify a series of adjacent variables using the : operator. We use this with two variable names, one on the left-hand side and one on the right. When we use : like this, select will subset both those two variables along with any others that fall in between them.

For example, if we want only the morphometric variables (bill_length_mm, bill_depth_mm, flipper_length_mm and body_mass_g) we could use:

```
select(penguins, bill_length_mm:body_mass_g)
```

```
## # A tibble: 344 x 4
##
       \verb|bill_length_mm| | \verb|bill_depth_mm| | flipper_length_mm| | \verb|body_mass_g| |
##
                  <dbl>
                                   <dbl>
                                                        <int>
                                                                       <int>
##
    1
                   39.1
                                    18.7
                                                           181
                                                                        3750
##
    2
                   39.5
                                    17.4
                                                           186
                                                                        3800
                                                           195
                                                                        3250
##
    3
                   40.3
                                    18
##
    4
                   NA
                                    NA
                                                           NA
                                                                          NA
##
    5
                   36.7
                                    19.3
                                                           193
                                                                        3450
##
    6
                   39.3
                                    20.6
                                                           190
                                                                        3650
    7
##
                   38.9
                                    17.8
                                                           181
                                                                        3625
##
    8
                   39.2
                                                           195
                                                                        4675
                                    19.6
##
    9
                   34.1
                                    18.1
                                                           193
                                                                        3475
## 10
                   42
                                                           190
                                                                        4250
                                    20.2
## # ... with 334 more rows
```

The: operator can also be combined with! if we need to drop a series of variables according to their position in a data frame or tibble. For example, we can use this trick to get the complement of the previous example, i.e. throw away the morphometric variables:

```
select(penguins, !bill_length_mm:body_mass_g)
```

```
## # A tibble: 344 x 4
##
      species island
                         sex
                                 year
##
      <chr>
              <chr>>
                         <chr>>
                                <int>
    1 Adelie
              Torgersen male
                                 2007
    2 Adelie
              Torgersen female
                                 2007
##
##
    3 Adelie
              Torgersen female
                                 2007
   4 Adelie
              Torgersen <NA>
                                 2007
##
    5 Adelie
              Torgersen female
                                 2007
    6 Adelie
              Torgersen male
                                 2007
    7 Adelie
              Torgersen female
                                 2007
    8 Adelie
              Torgersen male
                                 2007
    9 Adelie
              Torgersen <NA>
                                 2007
## 10 Adelie
              Torgersen <NA>
                                 2007
## # ... with 334 more rows
```

7.2.2 Renaming variables with select and rename

The select function can also rename variables at the same time as selecting them. To do this, we name the arguments using the name = value construct,

where the name of the selected variable is placed on the right-hand side (value), and the new name goes on the left-hand side (name).

For example, to select the species, bill_length_mm and bill_depth_mm variables from penguins, and in the process, rename bill_length_mm and bill_depth_mm to BillLength and BillDepth, use:

```
select(penguins, species, BillLength = bill_length_mm, BillDepth = bill_depth_mm)
```

```
## # A tibble: 344 x 3
##
      species BillLength BillDepth
##
      <chr>
                    <dbl>
                               <dbl>
##
   1 Adelie
                     39.1
                                18.7
##
    2 Adelie
                     39.5
                                17.4
##
    3 Adelie
                     40.3
                                18
##
    4 Adelie
                     NA
                                NA
##
    5 Adelie
                     36.7
                                19.3
##
    6 Adelie
                     39.3
                                20.6
##
    7 Adelie
                     38.9
                                17.8
##
   8 Adelie
                     39.2
                                19.6
    9 Adelie
##
                     34.1
                                18.1
## 10 Adelie
                     42
                                20.2
## # ... with 334 more rows
```

Renaming the variables is a common task. What should we do if the only thing we want to achieve is to rename variables, rather than rename and select them? **dplyr** provides an additional function called **rename** for exactly this purpose. This function renames some variables while retaining all others. It works like select. For example, to rename bill_length_mm and bill_depth_mm to BillLength and BillDepth but keep all the variables, use:

```
rename(penguins, BillLength = bill_length_mm, BillDepth = bill_depth_mm)
```

```
## # A tibble: 344 x 8
##
      species island
                      BillLength BillDepth flipper_length_~ body_mass_g sex
                                                                                    year
      <chr>
                             <dbl>
##
               <chr>
                                       <dbl>
                                                          <int>
                                                                       <int> <chr>
                                                                                   <int>
                              39.1
##
   1 Adelie
              Torger~
                                        18.7
                                                            181
                                                                       3750 male
                                                                                    2007
    2 Adelie
              Torger~
                              39.5
                                        17.4
                                                            186
                                                                       3800 fema~
                                                                                    2007
                              40.3
                                                                       3250 fema~
##
    3 Adelie
              Torger~
                                        18
                                                            195
                                                                                    2007
    4 Adelie
                                                                         NA <NA>
##
              Torger~
                              NA
                                        NA
                                                             NA
                                                                                    2007
##
    5 Adelie
              Torger~
                              36.7
                                        19.3
                                                            193
                                                                       3450 fema~
                                                                                    2007
##
    6 Adelie
              Torger~
                              39.3
                                        20.6
                                                            190
                                                                       3650 male
                                                                                    2007
##
    7 Adelie
              Torger~
                              38.9
                                        17.8
                                                            181
                                                                       3625 fema~
                                                                                    2007
##
    8 Adelie
              Torger~
                              39.2
                                        19.6
                                                            195
                                                                       4675 male
                                                                                    2007
    9 Adelie
              Torger~
                              34.1
                                        18.1
                                                            193
                                                                       3475 <NA>
                                                                                    2007
                                        20.2
                                                            190
                                                                       4250 <NA>
                                                                                    2007
## 10 Adelie Torger~
                              42
```

```
## # ... with 334 more rows
```

Notice rename also preserves the order of the variables found in the original

7.3 Creating variables with mutate

We use mutate to add new variables to a data frame or tibble. This is useful if we need to construct one or more derived variables to support an analysis. Basic usage of mutate looks like this:

```
mutate(<data>, <expression-1>, <expression-2>, ...)
```

Again, this is not an example we can run—it's pseudocode highlighting how to use mutate in abstract terms.

The first argument, <data>, must be the name of the object containing our data. We then include a series of one or more additional arguments, where each of these is a valid R expression involving one or more variables in <data>. We've have expressed these as <expression-1>, <expression-2>, where <expression-1> and <expression-2> represent the first two expressions, and the . . . is acting as a placeholder for the remaining expressions. These can be any valid R code that refers to variables in <data>. This is often a simple calculation (e.g. involving arithmetic), but it can be arbitrarily complex.

To see mutate in action, let's construct a new version of penguins that contains one extra variable—body mass measured in kilograms:

```
mutate(penguins, body_mass_g / 1000)
```

```
## # A tibble: 344 x 9
##
      species island
                        bill_length_mm bill_depth_mm flipper_length_mm body_mass_g
      <chr>
                                  <dbl>
##
              <chr>>
                                                 <dbl>
                                                                    <int>
                                                                                <int>
##
                                   39.1
                                                  18.7
   1 Adelie Torgersen
                                                                      181
                                                                                 3750
    2 Adelie Torgersen
                                   39.5
                                                  17.4
                                                                      186
                                                                                 3800
##
    3 Adelie
              Torgersen
                                   40.3
                                                  18
                                                                      195
                                                                                 3250
##
    4 Adelie
              Torgersen
                                   NA
                                                  NA
                                                                      NA
                                                                                   NA
##
   5 Adelie
              Torgersen
                                   36.7
                                                  19.3
                                                                      193
                                                                                 3450
   6 Adelie
                                   39.3
                                                  20.6
                                                                      190
                                                                                 3650
##
              Torgersen
    7 Adelie
              Torgersen
                                   38.9
                                                  17.8
                                                                      181
                                                                                 3625
   8 Adelie
              Torgersen
                                   39.2
                                                  19.6
                                                                      195
                                                                                 4675
   9 Adelie
              Torgersen
                                   34.1
                                                  18.1
                                                                      193
                                                                                 3475
## 10 Adelie
              Torgersen
                                   42
                                                  20.2
                                                                                 4250
                                                                      190
## # ... with 334 more rows, and 3 more variables: sex <chr>, year <int>,
```

This creates a copy of penguins with a new column called body_mass_g/1000

body_mass_g/1000 <dbl>

(look at the bottom of the printed output). That is not a very good name but do not worry—we will improve on it in a moment. Most of the rules that apply to select also apply to mutate:

- Quotes must not be placed around an expression that performs a calculation. This makes sense because the expression is meant to be evaluated so that it "does something". It is not a value.
- The mutate function does not have side effects, meaning it does not change the original penguins in any way. In the example, we printed the result produced by mutate rather than assigning it a name using <-, which means we have no way to access the result.
- The mutate function returns the same kind of object as the one it is working on: a data frame if our data was originally in a data frame, a tibble if it was a tibble.

Creating a variable called something like body_mass_g/1000 is not ideal because that is a difficult name to work with. Fortunately, the mutate function can name new variables at the same time as it creates them. We just name the arguments using =, placing the name on the left-hand side. Look at how to use this construct to name the new area variable body_mass_kg:

```
mutate(penguins, body_mass_kg = body_mass_g / 1000)
```

```
## # A tibble: 344 x 9
##
      species island
                         bill_length_mm bill_depth_mm flipper_length_mm body_mass_g
##
      <chr>
              <chr>
                                  <dbl>
                                                 <dbl>
                                                                                <int>
                                                                    <int>
##
   1 Adelie
              Torgersen
                                   39.1
                                                  18.7
                                                                      181
                                                                                 3750
   2 Adelie Torgersen
                                   39.5
                                                  17.4
                                                                      186
                                                                                 3800
   3 Adelie Torgersen
                                   40.3
                                                  18
                                                                      195
                                                                                 3250
   4 Adelie Torgersen
##
                                   NA
                                                  NA
                                                                       NA
                                                                                   NA
   5 Adelie Torgersen
                                   36.7
                                                  19.3
##
                                                                      193
                                                                                 3450
                                   39.3
   6 Adelie Torgersen
                                                  20.6
                                                                      190
                                                                                 3650
   7 Adelie
              Torgersen
                                   38.9
                                                  17.8
                                                                      181
                                                                                 3625
##
   8 Adelie
              Torgersen
                                   39.2
                                                  19.6
                                                                      195
                                                                                 4675
   9 Adelie
             Torgersen
                                   34.1
                                                  18.1
                                                                      193
                                                                                 3475
## 10 Adelie Torgersen
                                   42
                                                  20.2
                                                                      190
                                                                                 4250
## # ... with 334 more rows, and 3 more variables: sex <chr>, year <int>,
       body mass kg <dbl>
```

We can create more than one variable by supplying mutate multiple (named) arguments:

A tibble: 344 x 10

##		species	island	bill_length_mm	bill_depth_mm	flipper_length_mm	body_mass_g
##		<chr></chr>	<chr></chr>	<dbl></dbl>	<dbl></dbl>	<int></int>	<int></int>
##	1	Adelie	Torgersen	39.1	18.7	181	3750
##	2	Adelie	Torgersen	39.5	17.4	186	3800
##	3	Adelie	Torgersen	40.3	18	195	3250
##	4	Adelie	Torgersen	NA	NA	NA	NA
##	5	Adelie	Torgersen	36.7	19.3	193	3450
##	6	Adelie	Torgersen	39.3	20.6	190	3650
##	7	Adelie	Torgersen	38.9	17.8	181	3625
##	8	Adelie	Torgersen	39.2	19.6	195	4675
##	9	Adelie	Torgersen	34.1	18.1	193	3475
##	10	Adelie	Torgersen	42	20.2	190	4250
## # with 334 more rows, and 4 more variables: sex <chr>, year <int>,</int></chr>							
<pre>## # bill_size <dbl>, scaled_bill_size <dbl></dbl></dbl></pre>							

Notice that we placed each calculation on a new line, remembering to use a comma to separate arguments. We can do this because R ignores white space. Splitting long a function call across multiple lines in this way is helpful because it makes it easier to read and understand the sequence of calculations.

This last example reveals a nice feature of mutate—we can use newly created variables in further calculations. Here we constructed a synthetic bill size variable, and used that to calculate a second variable representing the ratio of bill size to body mass.

7.3.1 Transforming and dropping variables

Occasionally we need to construct one or more new variables and then drop all the other ones in the original dataset. The **transmute** function is designed to do this. It works exactly like **mutate**, but it has a slightly different behaviour:

```
## # A tibble: 344 x 2
##
      bill_size scaled_bill_size
##
          <dbl>
                            <dbl>
##
   1
           731.
                            0.195
##
    2
           687.
                            0.181
##
    3
           725.
                            0.223
##
    4
            NA
                           NA
                            0.205
##
    5
           708.
##
    6
           810.
                            0.222
##
    7
           692.
                            0.191
##
   8
           768.
                            0.164
##
   9
           617.
                            0.178
```

```
## 10 848. 0.200
## # ... with 334 more rows
```

Here we repeated the previous example, but now only the new variables were retained in the resulting tibble. If we also want to retain additional variables without altering them, we can pass them as unnamed arguments. For example, to retain **species** identity in the output, use:

```
## # A tibble: 344 x 3
##
      species bill_size scaled_bill_size
##
      <chr>
                  <dbl>
                                    <dbl>
##
   1 Adelie
                   731.
                                    0.195
##
   2 Adelie
                   687.
                                   0.181
##
   3 Adelie
                   725.
                                   0.223
##
   4 Adelie
                                   NA
                    NA
##
   5 Adelie
                   708.
                                   0.205
   6 Adelie
##
                   810.
                                   0.222
##
   7 Adelie
                   692.
                                   0.191
## 8 Adelie
                   768.
                                    0.164
##
   9 Adelie
                   617.
                                   0.178
## 10 Adelie
                   848.
                                    0.200
## # ... with 334 more rows
```

Chapter 8

Working with observations

8.1 Introduction

This chapter will explore the filter and arrange verbs. We discuss these functions together because they manipulate observations (i.e. rows) of a data frame or tibble:

- The filter function extracts a subset of observations based on supplied criteria
- The arrange function reorders the rows according to the values in one or more variables.

8.1.1 Getting ready

We'll be using the **dplyr** package, so we need to remember to load and attach the package in the current session:

library("dplyr")

We'll use the Palmer penguins data again to illustrate the ideas in this chapter. The examples below assume those data been read into R as a tibble with the name penguins.

8.2 Relational and logical operators

Most filter operations rely on some combination of **relational and logical operators**. Relational operators allow us to ask questions like, "are the values of 'x' greater than those of 'y': x > y". These sorts of comparisons are used by R to express whether or not a particular condition is met (because they generate

a logical vector of TRUE/FALSE values). Logical operators allow us to combine such conditions, thereby building up complex conditions from simpler ones.

This is best understood by example. We'll do that in a moment. For now, simply make a mental note of the different relational and logical operators:

- 1. Use **relational operators** to make comparisons between a pair of variables on the basis of conditions like 'less than' or 'equal to':
 - x < y: is x less than y?
 - x > y: is x greater than y?
 - $x \le y$: is x less than or equal to y?
 - $x \ge y$: is x greater than or equal to y?
 - x == y: is x equal to y?
 - x != y: is x not equal to y?
- 2. Use **logical operators** to connect two or more comparisons to arrive at a single overall criterion:
 - x & y: are both x AND y true?
 - x | y: is x OR y true?

Λ

Double == or single =?

Remember to use 'double equals' == when testing for equivalence between x and y. We all forget this from time to time and use 'single equals' = instead. This will lead to an error. \mathbf{dplyr} is pretty good at spotting this mistake these days and will warn you in its error message that you used = when you meant to use ==. Of course, if you don't read the error messages, you won't benefit from this helpful behaviour.

8.3 Subset observations with filter

We use filter to subset observations in a data frame or tibble containing our data. This is useful when we want to limit an analysis to a particular group of observations. Basic usage of filter looks something like this:

```
filter(<data>, <expression-1>, <expression-2>, ...)
```

Yes, this is pseudocode again. Let's review the arguments:

- The first argument, <data>, must be the name of the object (usually a data frame or tibble) containing our data. As with all dplyr verbs, this is not optional.
- We then include one or more additional arguments. Each of these is a valid R expression involving one or more variables in <data> that returns a logical vector. We've expressed these as <expression-1>, <expression-2>,

 \dots , where <expression-1> and <expression-2> represent the first two expressions, and the \dots is acting as placeholder for the remaining expressions.

To see filter in action, we'll use it to subset observations in the penguins dataset, based on two relational criteria:

```
filter(penguins, bill_length_mm > 45, bill_depth_mm > 18)
```

```
## # A tibble: 44 x 8
##
      species
                island
                           bill_length_mm bill_depth_mm flipper_length_~ body_mass_g
##
      <chr>
                <chr>>
                                     <dbl>
                                                   <dbl>
                                                                     <int>
                                                                                  <int>
##
   1 Adelie
                Torgersen
                                      46
                                                     21.5
                                                                        194
                                                                                   4200
## 2 Adelie
                Torgersen
                                      45.8
                                                     18.9
                                                                        197
                                                                                   4150
   3 Adelie
                Biscoe
                                      45.6
                                                     20.3
                                                                                   4600
                                                                        191
  4 Chinstrap Dream
                                      50
                                                    19.5
                                                                       196
                                                                                   3900
                                      51.3
## 5 Chinstrap Dream
                                                    19.2
                                                                       193
                                                                                   3650
    6 Chinstrap Dream
                                      45.4
                                                     18.7
                                                                       188
                                                                                   3525
##
   7 Chinstrap Dream
                                      52.7
                                                     19.8
                                                                       197
                                                                                   3725
  8 Chinstrap Dream
                                      46.1
                                                     18.2
                                                                        178
                                                                                   3250
## 9 Chinstrap Dream
                                      51.3
                                                     18.2
                                                                        197
                                                                                   3750
## 10 Chinstrap Dream
                                                     18.9
                                                                        195
                                                                                   4150
                                      46
## # ... with 34 more rows, and 2 more variables: sex <chr>, year <int>
```

In this example, we've created a subset of penguins that only includes observations where the bill_length_mm variable is greater than 45 and the bill_depth_mm variable is greater than 45, i.e. both conditions must be met for an observation to be retained. This is probably starting to feel repetitious,

but there are a few features of filter that we should be aware of:

- We do not surround each expression with quotes. The expression is meant to be evaluated—it is not 'a value.
- The result produced by filter was printed to the Console in the example. The filter function did not change the original penguins in any way (no side effects!).
- The filter function will return the same kind of data object it is working
 on: it returns a data frame if our data was originally in a data frame, and
 a tibble if it was a tibble.

Notice that including are two conditions separated by a comma means both conditions have to be met. There is another way to achieve the exact same result:

```
filter(penguins, bill_length_mm > 45 & bill_depth_mm > 18)
```

```
## # A tibble: 44 x 8
## species island bill_length_mm bill_depth_mm flipper_length_~ body_mass_g
```

##		<chr></chr>	<chr></chr>	<dbl></dbl>	<dbl></dbl>	<int></int>	<int></int>
##	1	Adelie	Torgersen	46	21.5	194	4200
##	2	Adelie	Torgersen	45.8	18.9	197	4150
##	3	Adelie	Biscoe	45.6	20.3	191	4600
##	4	${\tt Chinstrap}$	Dream	50	19.5	196	3900
##	5	${\tt Chinstrap}$	Dream	51.3	19.2	193	3650
##	6	${\tt Chinstrap}$	Dream	45.4	18.7	188	3525
##	7	${\tt Chinstrap}$	Dream	52.7	19.8	197	3725
##	8	Chinstrap	Dream	46.1	18.2	178	3250
##	9	Chinstrap	Dream	51.3	18.2	197	3750
##	10	${\tt Chinstrap}$	Dream	46	18.9	195	4150

This version links the two parts with the logical & operator. That is, rather than

... with 34 more rows, and 2 more variables: sex <chr>, year <int>

supplying bill_length_mm > 45 and bill_depth_mm > 18 as two arguments, we used a single R expression, combining them with the &.

We're pointing this out because we sometimes need to create filtering criteria that cannot be expressed as 'condition 1' and 'condition 2' and 'condition 3'... etc. Under those conditions we have to use logical operators to connect conditions. A simple instance of this situation is where we need to subset on an either/or basis. For example:

```
filter(penguins, bill_length_mm < 36 | bill_length_mm > 54)
```

```
## # A tibble: 29 x 8
##
      species island
                         bill_length_mm bill_depth_mm flipper_length_mm body_mass_g
##
      <chr>
               <chr>
                                   <dbl>
                                                  <dbl>
                                                                     <int>
                                                                                  <int>
   1 Adelie
              Torgersen
                                    34.1
                                                   18.1
##
                                                                       193
                                                                                   3475
##
   2 Adelie
              Torgersen
                                    34.6
                                                   21.1
                                                                       198
                                                                                   4400
##
    3 Adelie
              Torgersen
                                    34.4
                                                   18.4
                                                                       184
                                                                                   3325
##
    4 Adelie Biscoe
                                    35.9
                                                   19.2
                                                                       189
                                                                                   3800
##
   5 Adelie Biscoe
                                    35.3
                                                   18.9
                                                                       187
                                                                                   3800
##
   6 Adelie Biscoe
                                    35
                                                   17.9
                                                                       190
                                                                                   3450
##
   7 Adelie Biscoe
                                    34.5
                                                   18.1
                                                                       187
                                                                                   2900
##
   8 Adelie Biscoe
                                    35.7
                                                   16.9
                                                                       185
                                                                                   3150
   9 Adelie
              Biscoe
                                    35.5
                                                   16.2
                                                                       195
                                                                                   3350
## 10 Adelie
              Torgersen
                                    35.9
                                                   16.6
                                                                       190
                                                                                   3050
```

... with 19 more rows, and 2 more variables: sex <chr>, year <int>

This creates a subset of penguins that only includes observation where bill_length_mm is less than 36 or (|) greater than 54. This creates a subset of the data associated with the more 'extreme' values of bill length (unusually small or large).

We're not limited to using relational and logical operators when working with filter. The conditions specified in the filter function can be any expression

that returns a logical vector. The only constraint is that the output vector's length has to equal its input's length, or be a single logical values (TRUE or FALSE).

Here's an example. The **dplyr between** function is used to determine whether the values of a numeric vector fall in a specified range. It has three arguments: the numeric vector to filter on and the lower and upper and boundary values. For example:

```
filter(penguins, between(bill_length_mm, 36, 54))
```

```
## # A tibble: 313 x 8
##
      species island
                        bill_length_mm bill_depth_mm flipper_length_mm body_mass_g
      <chr>
                                 <dbl>
##
              <chr>
                                                <dbl>
                                                                  <int>
                                                                               <int>
## 1 Adelie Torgersen
                                  39.1
                                                 18.7
                                                                    181
                                                                                3750
## 2 Adelie Torgersen
                                  39.5
                                                 17.4
                                                                    186
                                                                                3800
  3 Adelie Torgersen
                                  40.3
                                                 18
                                                                    195
                                                                                3250
## 4 Adelie Torgersen
                                  36.7
                                                 19.3
                                                                    193
                                                                                3450
## 5 Adelie
                                                 20.6
             Torgersen
                                  39.3
                                                                    190
                                                                                3650
              Torgersen
   6 Adelie
                                  38.9
                                                 17.8
                                                                    181
                                                                                3625
## 7 Adelie
                                  39.2
                                                 19.6
                                                                    195
                                                                                4675
             Torgersen
## 8 Adelie
             Torgersen
                                   42
                                                 20.2
                                                                    190
                                                                                4250
## 9 Adelie
              Torgersen
                                  37.8
                                                 17.1
                                                                    186
                                                                                3300
## 10 Adelie
             Torgersen
                                  37.8
                                                 17.3
                                                                    180
                                                                                3700
## # ... with 303 more rows, and 2 more variables: sex <chr>, year <int>
```

8.4 Reordering observations with arrange

We use arrange to reorder the rows of a data frame or tibble. Basic usage of arrange looks like this:

```
arrange(<data>, <variable-1>, <variable-2>, ...)
```

Yes, this is pseudocode. As always, the first argument, <data>, is the name of the object containing our data. We then include a series of one or more additional arguments, where each of these is the name of a variable in <data>: <variable-1> and <variable-2> are names of the first two ordering variables, and the ... is acting as a placeholder for the remaining variables.

To see arrange in action, let's construct a new version of penguins where the rows have been reordered first by flipper_length_mm, and then by body_mass_g:

```
arrange(penguins, flipper_length_mm, body_mass_g)
```

A tibble: 344 x 8

##		species	island	bill_length_mm	${\tt bill_depth_mm}$	flipper_length_~	body_mass_g
##		<chr></chr>	<chr></chr>	<dbl></dbl>	<dbl></dbl>	<int></int>	<int></int>
##	1	Adelie	Biscoe	37.9	18.6	172	3150
##	2	Adelie	Biscoe	37.8	18.3	174	3400
##	3	Adelie	Torgersen	40.2	17	176	3450
##	4	Adelie	Dream	33.1	16.1	178	2900
##	5	Adelie	Dream	39.5	16.7	178	3250
##	6	Chinstrap	Dream	46.1	18.2	178	3250
##	7	Adelie	Dream	37.2	18.1	178	3900
##	8	Adelie	Dream	37.5	18.9	179	2975
##	9	Adelie	Dream	42.2	18.5	180	3550
##	10	Adelie	Biscoe	37.7	18.7	180	3600
##	# .	with 33	34 more rov	vs, and 2 more v	variables: sex	<chr>, year <int< td=""><td>></td></int<></chr>	>

This creates a new version of penguins where the rows are sorted according to the values of by flipper_length_mm and body_mass_g in ascending order – i.e. from smallest to largest. Look at the cases where flipper length is 178 mm. What do these show? Since flipper_length_mm was placed before body_mass_g in the arguments, the values of body_mass_g are only used to break ties within any particular value of flipper_length_mm.

For the sake of avoiding doubt about how arrange works, we will quickly review its behaviour. It works the same as every other **dplyr** verb we have looked at:

- The variable names used as arguments of arrange are not surrounded by quotes.
- The arrange function did not change the original penguins in any way.
- The arrange function will return the same kind of data object it is working on.

arrange sorts variables in ascending order by default. If we need it to sort a variable in descending order, we wrap the variable name in the **dplyr desc** function:

arrange(penguins, flipper_length_mm, desc(body_mass_g))

```
## # A tibble: 344 x 8
##
      species
                 island
                           bill_length_mm bill_depth_mm flipper_length_~ body_mass_g
##
      <chr>
                 <chr>
                                     <dbl>
                                                    <dbl>
                                                                      <int>
                                                                                   <int>
   1 Adelie
                                      37.9
                                                     18.6
##
                 Biscoe
                                                                        172
                                                                                    3150
    2 Adelie
                                                     18.3
                                                                        174
                                                                                    3400
##
                 Biscoe
                                      37.8
##
   3 Adelie
                 Torgersen
                                      40.2
                                                     17
                                                                        176
                                                                                    3450
##
   4 Adelie
                 Dream
                                      37.2
                                                     18.1
                                                                        178
                                                                                    3900
##
   5 Adelie
                 Dream
                                      39.5
                                                     16.7
                                                                        178
                                                                                    3250
##
    6 Chinstrap Dream
                                      46.1
                                                     18.2
                                                                        178
                                                                                    3250
   7 Adelie
                 Dream
                                      33.1
                                                     16.1
                                                                        178
                                                                                    2900
##
   8 Adelie
                                      37.5
                                                     18.9
                                                                        179
                                                                                    2975
                 Dream
```

```
## 9 Adelie Biscoe 40.5 18.9 180 3950
## 10 Adelie Biscoe 38.8 17.2 180 3800
## # ... with 334 more rows, and 2 more variables: sex <chr>, year <int>
```

This creates a new version of penguins where the rows are sorted according to the values of flipper_length_mm and body_mass_g, in ascending and descending order, respectively. Look carefully at the values in the flipper_length_mm and body_mass_g columns to see the difference between this example and the previous one.

Chapter 9

Summarising and grouping

This chapter will explore the summarise and group_by verbs. We consider together because they are often used in combination. Their usage is also a bit different from the other **dplyr** verbs we've encountered. Here's a quick summary of what they do:

- The group_by function adds information to its input (a data frame or tibble), which makes subsequent calculations happen on a group-specific basis.
- The summarise function is a data reduction function that calculates singlenumber summaries of one or more variables, respecting the group structure if present.

We illustrate these ideas using the Paler penguins data set, which we assume has been read into a tibble called **peguins**.

9.1 Summarising variables with summarise

We use summarise to calculate summaries of variables in an object containing our data. We do this kind of calculation all the time when analysing data. In terms of pseudo-code, usage of summarise looks like this:

```
summarise(<data>, <expression-1>, <expression-2>, ...)
```

The first argument, <data>, must be the name of the data frame or tibble containing our data. We then include a series of one or more additional arguments; each of these is a valid R expression involving at least one variable in <data>. These are given by the pseudo-code placeholder <expression-1>, <expression-2>, ..., where <expression-1> and <expression-2> represent the first two expressions, and the ... is acting as placeholder for the remaining

expressions. These expressions can be any calculation involving R functions that returns a vector of some kind.

The summarise function seems to work a lot like mutate. The main difference is that the expressions mutate uses have to all return a vector of the same length as their inputs. In contrast, summarise expressions used all have to produce the same length output, but those outputs can be any length. They often return a single value because they are summarising the data in some way, but they don't have to.

The summarise verb is best understood by example. The **dplyr** function n_distinct takes a calculates the number of distinct (i.e. unique) cases in a vector. We can use n_distinct with summarise to calculate the number of unique vales of the bill_length_mm and bill_depth_mm variables like this:

summarise(penguins, n_distinct(bill_length_mm), n_distinct(bill_depth_mm))

Notice what kind of object summarise returns—it's a tibble with one row and two columns: two columns because we calculated two counts, and one row containing because we only one set of counts. There are a few other things to note about how summarise works:

- The expression that performs each calculation is not surrounded by quotes because it's an expression that it 'does a calculation'.
- The order of the columns in the output is the same as the order in which they were created in the <expression-1>, <expression-2>, ... list.
- summarise returns the same kind of data object as its input—it returns a data frame if our data was originally in a data frame, or a tibble if it was in a tibble.
- If we don't specify a name summarise uses the actual R expression to name the columns of its output (e.g. n distinct(bill length mm))

Variable names based on the calculation (e.g. n_distinct(bill_length_mm)) are not ideal because they are long and contain special reserved characters like (. This makes it difficult refer to columns in the output because we have to remember to place back ticks (`) around their name whenever we want to refer to them.

Fortunately, the summarise function can name the new variables at the same time as they are created (just like mutate). We do this by naming the arguments using =, placing the name we require on the left hand side. For example:

This time we end up with summary data set that has reasonable column names. Notice how we organised that example—we placed each calculation on a new line. We don't have to do this, but since R doesn't care about white space, we can use newlines and spaces to keep everything more human-readable. It pays to organise summarise calculations like this when they become longer.

9.1.1 More complicated calculations with summarise

Many useful base R functions can be used with summarise. Of particular value are those that calculate various summaries of numeric variables are, such as:

- min and max calculate the minimum and maximum values.
- mean and median calculate the mean and median, and
- sd and var calculate the standard deviation and variance.

We do need to pay attention when using base R functions with **dplyr**. Take a look at this attempt to use summarise to calculate the mean of bill_length_mm and bill_length_mm:

```
## n_bill_length n_bill_depth
## <dbl> <dbl>
## 1 NA NA
```

No numbers—just a pair of NAs. We forgot about the presence of missing values in the penguins data. Both bill_length_mm and bill_depth_mm contain missing values. When the mean function encounters even one missing value in its input its default behaviour is to spit out NA. It is possible to change that behaviour by setting the na.rm argument of mean:

```
## # A tibble: 1 x 2
## n_bill_length n_bill_depth
## <dbl> <dbl>
## 1 43.9 17.2
```

This example demonstrates something important—the functions we use within **summarise** often have their own arguments, and we sometimes need to set those arguments to perform the calculation we want.

Almost any R code can be used as **summarise** expressions. This means we can combine more than one function to build up arbitrarily complicated calculations. For example, if we need to know the ratio of the mean bill length and mean bill width in **penguins**, we would use:

```
summarise(penguins,
   ratio = mean(bill_length_mm, na.rm = TRUE) / mean(bill_depth_mm,
   na.rm = TRUE))

## # A tibble: 1 x 1
## ratio
## <dbl>
## 1 2.56
```

The ability to work with arbitrary expressions makes summarise (and mutate) very powerful.

9.2 Grouped operations using group_by

Performing a calculation with one or more variables using the whole data set can be useful. However, we often need to carry out calculations on different subsets of our data. For example, it's more useful to know how the mean bill length and depth vary among the different species in the penguins data set, rather than knowing the overall mean of these traits. We could calculate separate means by using filter to create different subsets of penguins, and then use summary on each of these to calculate the means. This would get the job done, but it's inefficient and quickly becomes tiresome if we have to work with many groups.

The group_by function provides an elegant solution to this kind of problem. All the group_by function does is add a bit of information to a tibble or data frame. In effect, it defines subsets of data based on one or more grouping variables. That's all it does.

The magic happens when the grouped object is used with a **dplyr** verb like summarise or mutate. Once a the data has been tagged with grouping information, operations that involve **dplyr** verbs are carried out on separate subsets of the data—defined by the values of the grouping variable(s)—and then combined.

Basic usage of group_by looks like this:

```
group_by(<data>, <variable-1>, <variable-2>, ...)
```

The first argument, <data>, must be the name of the object containing our data. We then have to include one or more additional arguments, where each one is the name of a variable in <data>. We have expressed this as <variable-1>, <variable-2>, ..., where <variable-1> and <variable-2> are names of the first two variables, and the ... is acting as a placeholder for the remaining variables.

We'll illustrate group_by by using it alongside summarise. We're aiming to calculate the mean bill length for each species in penguins. This is a two-step process. The first step uses group_by to add grouping information to penguins. Take a look at what we end up with when we do that:

```
group_by(penguins, species)
```

```
## # A tibble: 344 x 8
## # Groups:
               species [3]
##
      species island
                        bill_length_mm bill_depth_mm flipper_length_mm body_mass_g
##
      <chr>
              <chr>>
                                 dbl>
                                                <dbl>
                                                                  <int>
   1 Adelie Torgersen
##
                                  39.1
                                                 18.7
                                                                    181
                                                                                3750
   2 Adelie
             Torgersen
                                  39.5
                                                 17.4
                                                                    186
                                                                                3800
##
                                  40.3
                                                 18
                                                                    195
                                                                                3250
## 3 Adelie
             Torgersen
## 4 Adelie
             Torgersen
                                  NA
                                                 NA
                                                                     NA
                                                                                  NA
## 5 Adelie
                                  36.7
                                                 19.3
                                                                    193
                                                                                3450
             Torgersen
## 6 Adelie
             Torgersen
                                   39.3
                                                 20.6
                                                                    190
                                                                                3650
## 7 Adelie
                                                                                3625
             Torgersen
                                   38.9
                                                 17.8
                                                                    181
  8 Adelie
              Torgersen
                                   39.2
                                                 19.6
                                                                    195
                                                                                4675
## 9 Adelie
              Torgersen
                                   34.1
                                                 18.1
                                                                    193
                                                                                3475
## 10 Adelie
              Torgersen
                                   42
                                                 20.2
                                                                    190
                                                                                4250
## # ... with 334 more rows, and 2 more variables: sex <chr>, year <int>
```

Compare this to the output produced when we print the original penguins data set:

```
penguins
```

```
## # A tibble: 344 x 8
##
      species island
                        bill_length_mm bill_depth_mm flipper_length_mm body_mass_g
##
      <chr>
              <chr>>
                                  <dbl>
                                                <dbl>
                                                                   <int>
                                                                               <int>
##
   1 Adelie Torgersen
                                   39.1
                                                 18.7
                                                                     181
                                                                                3750
##
    2 Adelie
              Torgersen
                                   39.5
                                                 17.4
                                                                     186
                                                                                3800
   3 Adelie
              Torgersen
                                   40.3
                                                 18
                                                                     195
                                                                                3250
   4 Adelie
              Torgersen
                                   NA
                                                 NA
                                                                      NA
                                                                                  NA
                                                                     193
                                                                                3450
## 5 Adelie Torgersen
                                   36.7
                                                 19.3
```

```
##
    6 Adelie
              Torgersen
                                    39.3
                                                   20.6
                                                                        190
                                                                                   3650
    7 Adelie
                                    38.9
##
              Torgersen
                                                   17.8
                                                                        181
                                                                                    3625
                                    39.2
   8 Adelie
              Torgersen
                                                   19.6
                                                                        195
                                                                                   4675
   9 Adelie
              Torgersen
                                    34.1
                                                   18.1
                                                                        193
                                                                                   3475
                                                                        190
## 10 Adelie
              Torgersen
                                    42
                                                   20.2
                                                                                   4250
## # ... with 334 more rows, and 2 more variables: sex <chr>, year <int>
```

There is very little difference—group_by really doesn't do much on its own. The main change is that printing the tibble resulting from the group_by operation shows a bit of additional information at the top: Groups: species [3]. This tells us that the tibble is now grouped by the species variable. The [3] part tells us that there are three different groups (i.e. species of penguin). The only thing group_by did was add this grouping information to a copy of penguins.

The original penguins object was not altered in any way. If we want to do anything useful with the grouped tibble we need to assign it a name so that we can work with it:

```
penguins_by_species <- group_by(penguins, species)</pre>
```

Now we have a grouped tibble called penguins_by_species in which the value of species define the different groups—any row where species is equal to 'Adelie' is assigned to the first group, any row where species is equal to 'Chinstrap' is assigned to a second group, and any row where species is equal to 'Gentoo' is assigned to a third group.

dplyr operations on this tibble will now be performed on a 'by group' basis. To see this in action, we use **summarise** to calculate the mean bill length:

This is part two of the two-step process mentioned above. When we used summarise on an ungrouped object, the result was a tibble with one row—the overall global mean. Now the resulting tibble has three rows, one for each species in the data set. The mean_bill_length column shows the mean bill lengths for each species. The species column tells us what species each mean belongs to. Notice that summarise also printed an (un)helpful message:

[`]summarise()` ungrouping output (override with `.groups` argument)

There's no need to worry about this. It is simply saying that summarise has removed the grouping information from the resulting tibble.

We can also carry out multiple calculations with grouped data if we need to. For example, if we need to calculate the mean bill length and mean bill depth for each species, we would use the grouped version of penguins like this:

```
## # A tibble: 3 x 3
##
               mean_bill_length mean_bill_depth
     species
##
     <chr>
                           <dbl>
## 1 Adelie
                           38.8
                                            18.3
## 2 Chinstrap
                           48.8
                                            18.4
## 3 Gentoo
                            47.5
                                            15.0
```

9.2.1 More than one grouping variable

What if we need to calculate summaries using more than one grouping variable? The workflow is unchanged. Assume we need to know the mean body mass of males and females of each penguin species. First, we make a grouped copy of penguins using the appropriate grouping variables:

```
penguins_by_species_sex <- group_by(penguins, species, sex)</pre>
```

We called the grouped tibble penguins_by_species_sex. Look at what happens when we print this:

```
penguins_by_species_sex
```

```
## # A tibble: 344 x 8
## # Groups:
               species, sex [8]
##
      species island
                        bill_length_mm bill_depth_mm flipper_length_mm body_mass_g
      <chr>
              <chr>>
                                 <dbl>
                                               <dbl>
                                                                  <int>
                                                                              <int>
                                                18.7
## 1 Adelie Torgersen
                                  39.1
                                                                    181
                                                                               3750
## 2 Adelie Torgersen
                                  39.5
                                                17.4
                                                                    186
                                                                               3800
                                                18
                                                                               3250
   3 Adelie
             Torgersen
                                  40.3
                                                                    195
##
   4 Adelie
##
             Torgersen
                                  NA
                                                NA
                                                                    NA
                                                                                 NA
## 5 Adelie Torgersen
                                  36.7
                                                19.3
                                                                    193
                                                                               3450
## 6 Adelie
             Torgersen
                                  39.3
                                                20.6
                                                                    190
                                                                               3650
## 7 Adelie
             Torgersen
                                  38.9
                                                17.8
                                                                    181
                                                                               3625
## 8 Adelie
             Torgersen
                                  39.2
                                                19.6
                                                                    195
                                                                               4675
## 9 Adelie
             Torgersen
                                  34.1
                                                18.1
                                                                    193
                                                                               3475
## 10 Adelie Torgersen
                                                20.2
                                                                    190
                                                                               4250
                                  42
```

```
## # ... with 334 more rows, and 2 more variables: sex <chr>, year <int>
```

We see Groups: species, sex [8] near the top, which tells us that the tibble is grouped by two variables (species and sex) with eight unique combinations of values. That seems odd at first—there are three species and two sexes represented in this dataset, which gives six possible combinations at most.

The reason for the discrepancy becomes clear when we move on to calculate the mean body mass for each sex-species combination:

```
## # A tibble: 8 x 3
## # Groups:
                species [3]
##
     species
                sex
                       body_mass_g
##
     <chr>
                <chr>>
                              <dbl>
## 1 Adelie
                female
                              3369.
## 2 Adelie
               male
                              4043.
## 3 Adelie
                <NA>
                              3540
## 4 Chinstrap female
                              3527.
## 5 Chinstrap male
                              3939.
## 6 Gentoo
                female
                              4680.
## 7 Gentoo
                male
                              5485.
## 8 Gentoo
                <NA>
                              4588.
```

This shows mean body mass for each unique combination of species and sex. The first line shows that the mean body mass associated with female Adelie penguins is 3369; the second line shows us the mean body mass associated with male Adelie penguins is 4043. The third line shows us the mean body mass of Adelie penguins where sex is missing (NA). That explains why we ended up with more groups than unique combinations of species and sex — missing values create extra groups.

9.2.2 Using group_by with other verbs

The summarise function is the **dplyr** verb that is most often used with grouped data. However, all the main **dplyr** verbs will alter their behaviour to respect group information when it is present. For example, when mutate or transmute are used with a grouped object the calculation of new variables occur "by group". Here's an example:

```
## # A tibble: 344 x 3
## # Groups:
                species, sex [8]
##
                      body_mass_cen
      species sex
##
      <chr>
               <chr>
                              <dbl>
##
    1 Adelie
              male
                              -293.
    2 Adelie
              female
                               431.
    3 Adelie
              female
                             -119.
##
    4 Adelie
              <NA>
                               NA
                               81.2
##
    5 Adelie
              female
##
    6 Adelie
              male
                             -393.
##
    7 Adelie
              female
                               256.
##
    8 Adelie
              male
                              632.
    9 Adelie
              <NA>
                               -65
## 10 Adelie
              <NA>
                               710
## # ... with 334 more rows
```

This calculated a standardised measure of body mass. The new body_mass_cen variable contains the difference between the original body mass and its mean in the appropriate species-sex group (rather than the overall mean).

9.3 Removing grouping information

On occasion, it's necessary to remove grouping information and revert to operating on the whole data set. The ungroup function removes grouping information:

ungroup(penguins_by_species)

```
## # A tibble: 344 x 8
                         bill_length_mm bill_depth_mm flipper_length_mm body_mass_g
##
      species island
##
      <chr>>
               <chr>>
                                   <dbl>
                                                  <dbl>
                                                                      <int>
                                                                                  <int>
                                                                                   3750
##
    1 Adelie
              Torgersen
                                    39.1
                                                   18.7
                                                                        181
    2 Adelie
                                    39.5
                                                   17.4
                                                                                   3800
              Torgersen
                                                                        186
##
    3 Adelie
              Torgersen
                                    40.3
                                                   18
                                                                        195
                                                                                   3250
    4 Adelie
               Torgersen
                                    NA
                                                   NA
                                                                        NA
                                                                                     NA
                                    36.7
    5 Adelie
              Torgersen
                                                   19.3
                                                                        193
                                                                                   3450
    6 Adelie
              Torgersen
                                    39.3
                                                   20.6
                                                                        190
                                                                                   3650
                                                   17.8
    7 Adelie
              Torgersen
                                    38.9
                                                                        181
                                                                                   3625
##
    8 Adelie
               Torgersen
                                    39.2
                                                   19.6
                                                                        195
                                                                                   4675
##
    9 Adelie
              Torgersen
                                    34.1
                                                   18.1
                                                                        193
                                                                                   3475
## 10 Adelie
               Torgersen
                                    42
                                                   20.2
                                                                        190
                                                                                   4250
## # ... with 334 more rows, and 2 more variables: sex <chr>, year <int>
```

Looking at the top right of the printed summary, we can see that the Group: part is now gone—the ungroup function effectively recreated the original penguins tibble.

Chapter 10

Building pipelines

We don't often use the various **dplyr** verbs in isolation. Instead, they are combined in a sequence to prepare the data for further analysis. For example, we might create a new variable or two with **mutate** and then use **group_by** and **summarise** to calculate some numerical summaries. This chapter will introduce something called the pipe operator: %>%. The pipe operator's job is to allow us to represent a sequence of such steps in a transparent, readable manner.

10.1 Why do we need 'pipes'?

We've seen that carrying out calculations on a per-group basis can be achieved by grouping a tibble, assigning this a name, and then applying the summarise function to the new tibble. For example, in the previous chapter we saw that to calculate the mean bill length for each species in the Palmer penguins data set, penguins, we first create a grouped version of it:

```
penguins_by_species <- group_by(penguins, species)</pre>
```

Then we use **summarise** with the grouped version to calculate the mean bill length in each group:

```
## 3 Gentoo 47.5
```

There's nothing wrong with this way of doing things. However, building up an analysis this was is quite lengthy because we have to keep storing intermediate steps. This is especially true if an analysis involves more than a couple of steps. It also tends to clutter the global environment with many intermediate objects we don't need to keep.

One way to make things more concise is to use function nesting, like this:

```
summarise(group_by(penguins, species),
    mean_bill_length = mean(bill_length_mm, na.rm = TRUE))
```

In this version, we placed the <code>group_by</code> function call inside the list of arguments to <code>summarise</code>. Remember—we have to read nested function calls from the inside out to understand what they are doing. This is exactly equivalent to the previous example, but now we get the result without having to store intermediate data.

However, there are a couple of very good reasons why this approach is not advised:

- Experienced R users might not mind this approach because they're used to it. Nonetheless, no reasonable person would argue that nesting functions inside one another is intuitive. Reading outward from the inside of a large number of nested functions is hard work.
- Using function nesting is an error-prone approach. For example, it's very easy to accidentally put an argument on the wrong side of a closing). If we're lucky, this will produce an error and we'll catch the problem. If we're not, we may end up with complete nonsense in the output.

10.2 Using pipes (%>%)

There is a better way to combing **dplyr** functions, which has the dual benefit of keeping our code concise and readable while avoiding the need to clutter the global environment with intermediate objects. This third approach involves the 'pipe operator': %>%. Notice there are no spaces between the three characters that make up the pipe—spaces are not allowed (e.g. % > % will not be recognised as the pipe).

The pipe operator isn't part of base R. Instead, **dplyr** imports it from another package and makes it available for us to use.

The pipe has become very popular in recent years. The main reason for this is because it allows us to specify a chain of function calls in a (reasonably) human readable format. Here's how we write the previous example using the pipe operator %>%:

```
penguins %>% group_by(., species) %>% summarise(., mean_bill_length = mean(bill_length_mm, na.rm
```

How do we make sense of this? Every time we see the %% operator it means the following: take whatever is produced by the left-hand expression and use it as an argument in the function on the right-hand side. The . serves as a placeholder for the location of the corresponding argument. A sequence of calculations can then be read from left to right, just as we would read the words in a book. This example says, take the penguins data, group it by species, then take that grouped tibble and apply the summarise function to it to calculate the mean of mean_bill_length.

This is exactly the same calculation we did above.

When using the pipe operator we can often leave out the . placeholder. Remember, this signifies the argument of the function on the right of %>% that is associated with the result from the left of %>%. If we choose to leave out the ., the pipe operator assumes we meant to slot it into the first argument. This means we can simplify our example even more:

```
penguins %>% group_by(species) %>% summarise(mean_bill_length = mean(bill_length_mm, na.rm = TRUE
```

This is why the first argument of a dplyr verb is always the data. Adopting this convention ensures we can use %>% without explicitly specifying the argument to pipe into. Data goes into the pipe; data comes out of the pipe.

Remember, R does not care about white space, which means we can break a

piped set of functions over several lines to make our code more readable:

```
penguins %>%
  group_by(species) %>%
  summarise(mean_bill_length = mean(bill_length_mm, na.rm = TRUE))
```

Now each step in the pipeline is on its own line. Most **dplyr** users use this formatting convention to improve the readability of their R code.

Finally, we do have to remember to assign the result of a chain of piped functions a name if we want to capture the result and use it later. We have to break the left to right rule a bit to do this, placing the assignment at the beginning¹:

```
bill_length_means <-
  penguins %>%
  group_by(species) %>%
  summarise(mean_bill_length = mean(bill_length_mm, na.rm = TRUE))
```



Why is %>% called the 'pipe' operator?

The %>% operator takes the output from one function and "pipes it" to another as the input. It's called 'the pipe' for the simple reason that it allows us to create an analysis 'pipeline' from a series of function calls. Incidentally, if you Google the phrase "magrittr pipe" you'll see why **magrittr** is a clever name for an R package.

One final piece of advice—make an effort to learn how to use the %>% method of piping together functions. Why? Because it's the simplest and cleanest method for doing this, many of the examples in the **dplyr** help files and on the web use it, and the majority of people carrying out real-world data wrangling with **dplyr** rely on piping.

 $^{^{1}}$ Actually, there is a rightward assignment operator, ->, but let's pretend that does not exist.

Chapter 11

Helper functions

11.1 Introduction

In addition to the main **dplyr** verbs, the package provides quite a few **helper functions**. Helper functions are used in conjunction with the main verbs to make specific tasks and calculations a bit easier. Many of these are summarised in the **dplyr** cheat sheat (under *Manipulate Variables*, *Vector Functions* and *Summary Functions*). We are not going to review every single one of them in this chapter. Instead, we aim to point out where helper functions tend to be used and highlight a few of the more useful ones.

11.2 Working with select

There are a few helper functions that can be used with select. Their job is to make it easier to match variable names according to various criteria. We'll look at the three simplest of these—look at the examples in the help file for select and the cheat sheat to see what else is available.

We can select variables according to the sequence of characters used at the start of their name with the starts_with function. For example, to select all the variables in penguins that begin with the word "bill", we use:

```
select(penguins, starts_with("Bill"))
```

```
## # A tibble: 344 x 2
##
      bill_length_mm bill_depth_mm
##
                <dbl>
                               <dbl>
##
                 39.1
                                18.7
   1
    2
                 39.5
                                17.4
                 40.3
                                18
##
    3
```

```
##
                  NA
                                  NA
                  36.7
                                  19.3
##
    5
    6
                  39.3
                                  20.6
##
##
    7
                  38.9
                                  17.8
##
    8
                  39.2
                                  19.6
##
    9
                  34.1
                                  18.1
## 10
                  42
                                  20.2
## # ... with 334 more rows
```

This returns a tibble containing just bill_length_mm and bill_depth_mm. There is also a helper function to select variables according to characters used at the end of their name—the ends_with function (no surprises there).

Notice that we quote the name we want to match against because <code>starts_with</code> expects a literal character value. This is not optional. Unusually, <code>starts_with</code> and <code>ends_with</code> are not case sensitive by default. For example, we passed <code>starts_with</code> the argument "Bill" instead of "bill", yet it still selected variables beginning with the character string "bill". If we want to select variables on a case-sensitive basis, we need to set an argument <code>ignore.case</code> to <code>FALSE</code> in <code>starts_with</code> and <code>ends_with</code>.

The last select helper function we'll look at is called contains. This one allows us to select variables based on a partial match anywhere in their name. Look at what happens if we pass contains the argument "length":

```
select(penguins, contains("length"))
```

```
## # A tibble: 344 x 2
##
      bill_length_mm flipper_length_mm
##
                 <dbl>
                                     <int>
##
    1
                 39.1
                                       181
    2
                  39.5
##
                                       186
##
    3
                  40.3
                                       195
##
    4
                 NA
                                        NA
##
    5
                  36.7
                                       193
##
    6
                  39.3
                                       190
    7
                  38.9
                                       181
##
                  39.2
                                       195
##
    8
                  34.1
##
    9
                                       193
##
   10
                  42
                                       190
## # ... with 334 more rows
```

This selects all the variables with the word 'length' in their name.

There is nothing to stop us combining the different variable selection methods. For example, we can use this approach to select all the variables that have some units at the end of their names (millimetres or grams):

```
select(penguins, ends_with("_mm"), ends_with("_g"))
```

```
## # A tibble: 344 x 4
      bill_length_mm bill_depth_mm flipper_length_mm body_mass_g
##
##
                <dbl>
                                <dbl>
                                                    <int>
                                                                 <int>
##
                 39.1
                                 18.7
                                                      181
                                                                  3750
##
    2
                 39.5
                                 17.4
                                                      186
                                                                  3800
##
    3
                 40.3
                                 18
                                                      195
                                                                  3250
##
    4
                 NA
                                 NA
                                                                    NA
                                                       NA
    5
                 36.7
                                 19.3
                                                      193
                                                                  3450
##
##
    6
                 39.3
                                 20.6
                                                      190
                                                                  3650
    7
                 38.9
                                 17.8
                                                                  3625
                                                      181
##
                 39.2
                                 19.6
                                                                  4675
    8
                                                      195
##
    9
                 34.1
                                 18.1
                                                      193
                                                                  3475
## 10
                 42
                                 20.2
                                                      190
                                                                  4250
## # ... with 334 more rows
```

When we apply more than one selection criteria like this, the select function returns the variables that match any criteria, rather than the set that meets all of them.

11.3 Working with mutate and transmute

There are quite a few **helper functions** that can be used with mutate. These make it easier to carry out certain calculations that aren't easy to do with base R. We won't explore these here as they tend to be needed only in quite specific circumstances. However, in situations where we need to construct an unusual variable, it's worth looking at that handy cheat sheat to see what options might be available.

We will look at one particularly useful helper function that's used a lot when we need to recode a particular variable using mutate. The function is called case_when. It works by setting up a series of paired matching criteria and replacement values. For example, imagine that we want to replace the names in species with three-letter shortcodes for each species. This is how to achieve that using case_when with mutate:

##	## # A tibble: 344 x 8								
##		species	island	bill_length_mm	${\tt bill_depth_mm}$	${\tt flipper_length_mm}$	body_mass_g		
##		<chr></chr>	<chr></chr>	<dbl></dbl>	<dbl></dbl>	<int></int>	<int></int>		
##	1	ADL	Torgersen	39.1	18.7	181	3750		
##	2	ADL	Torgersen	39.5	17.4	186	3800		
##	3	ADL	Torgersen	40.3	18	195	3250		
##	4	ADL	Torgersen	NA	NA	NA	NA		
##	5	ADL	Torgersen	36.7	19.3	193	3450		
##	6	ADL	Torgersen	39.3	20.6	190	3650		
##	7	ADL	Torgersen	38.9	17.8	181	3625		
##	8	ADL	Torgersen	39.2	19.6	195	4675		
##	9	ADL	Torgersen	34.1	18.1	193	3475		
##	10	ADL	Torgersen	42	20.2	190	4250		

... with 334 more rows, and 2 more variables: sex <chr>, year <int>

The mutate bit of this is not new. Look at the case_when component—there are four criteria. The first of these is species == "Adelie" ~ "ADL". The way to read this is, "look for cases where the value of species is equal to "Adelie", and where that is true spit out the value "ADL". case_when steps through each criterion like this in turn, trying to find a match. The last one TRUE ~ "UNKNOWN" acts as a catch-all for the non-matches.

This looks confusing at first but it does make sense with a bit of practise, and recoding variables using case_when is a lot easier than going through a spread-sheet by hand.

11.4 Working with filter

There aren't that many **dplyr** helper function that works with **filter**. In fact, we've already looked at the most useful one: the **between** function. This is used to identify cases where the values of a numeric variable lie inside a defined range. For example, if we want all the individuals that had a body mass in the 4-5kg range, we could use:

filter(penguins, between(body_mass_g, 4000, 5000))

```
## # A tibble: 116 x 8
##
      species island
                         bill_length_mm bill_depth_mm flipper_length_mm body_mass_g
##
      <chr>
               <chr>
                                   <dbl>
                                                  <dbl>
                                                                      <int>
                                                                                  <int>
    1 Adelie
                                    39.2
                                                   19.6
##
              Torgersen
                                                                        195
                                                                                    4675
    2 Adelie
                                    42
                                                   20.2
                                                                        190
                                                                                   4250
              Torgersen
##
    3 Adelie
              Torgersen
                                    34.6
                                                   21.1
                                                                        198
                                                                                    4400
##
    4 Adelie
              Torgersen
                                    42.5
                                                   20.7
                                                                        197
                                                                                    4500
##
    5 Adelie
              Torgersen
                                    46
                                                   21.5
                                                                        194
                                                                                    4200
    6 Adelie Dream
                                    39.2
                                                   21.1
                                                                        196
                                                                                    4150
   7 Adelie Dream
                                    39.8
                                                   19.1
                                                                                    4650
                                                                        184
```

```
## 8 Adelie Dream
                                  44.1
                                                19.7
                                                                   196
                                                                              4400
## 9 Adelie Dream
                                  39.6
                                                18.8
                                                                   190
                                                                              4600
## 10 Adelie Dream
                                  42.3
                                                21.2
                                                                              4150
                                                                   191
## # ... with 106 more rows, and 2 more variables: sex <chr>, year <int>
```

11.5 Working with summarise

There are a small number **dplyr** helper functions that can be used with **summarise**. These generally provide summaries that aren't available directly using base R functions. For example, we've already seen the **n_distinct** function in action. This can be used to calculate the number of distinct values of a variable:

This confirms what we already knew—that there are three unique species and three unique islands in the penguins data set.

Part III Exporing Data

Chapter 12

Exploratory data analysis

12.1 Introduction

Exploratory data analysis (EDA) was promoted by the statistician John Tukey in his 1977 book, "Exploratory Data Analysis". The broad goal of EDA is to help us formulate and refine hypotheses that lead to informative analyses or further data collection. The core objectives of EDA are:

- to suggest hypotheses about the causes of observed phenomena.
- to guide the selection of appropriate statistical tools and techniques,
- to assess the assumptions on which statistical analysis will be based,
- to provide a foundation for further data collection.

EDA involves a mix of numerical and visual methods of analysis. Statistical methods are sometimes used to supplement EDA. However, the main purpose of EDA is to facilitate understanding before carrying out formal statistical modelling. Even if we already know what kind of analysis we plan to pursue, it's always a good idea to **explore a data set before diving into that analysis**. At the very least, this will help determine whether or not our plans are sensible. Very often, it uncovers new patterns and insights.

In this chapter, we're going to examine some basic concepts that underpin EDA. We will:

- 1. see how to classify different types of variables,
- 2. distinguish between **populations and samples**, and
- 3. review some key descriptive statistics.

This will provide a conceptual foundation and vocabulary for learning how to explore data in later chapters.

12.2 Statistical variables and data

In the Data frames chapter, we pointed out that the word 'variable' can mean one of two things. In programming, a variable is a name-value association that we create when we run some code. Statisticians use the word differently. To them, a variable is any characteristic or quantity that can be measured, classified or experimentally controlled. Much of statistics is about quantifying and explaining the variation in such quantities as best we can.

Species richness, relative abundance, infection status, enzyme activity, gene frequency, and blood glucose concentration are examples of statistical variables we encounter in the biological sciences. These as statistical variables because their values vary between different observations. For instance, 'annual fitness'—measured as the number of offspring produced—is a variable that differs both among the organisms in a population and over the life cycle of a given individual.

There are different ways to describe statistical variables according to how they can be analysed, measured, or presented. It's important to be clear about what kind of variables we're dealing with because this determines how we should visualise the data, and later, how we might analyse it statistically. There are many different ways to go about classifying variables. However, we only need to be aware of two fairly simple classification schemes in this book: numeric vs categorical variables and ratio vs interval scales.

12.2.1 Numeric vs categorical variables

Numeric variables have values that describe a measurable quantity like 'how many' or 'how much' as a number. Numeric variables are also called quantitative variables; the data collected containing numeric variables are called quantitative data. Numeric variables may be classified as either continuous or discrete:

- Continuous numeric variable: Observations can take any value in a set of real numbers, i.e. numbers represented with decimals. Examples of continuous variables include concentration, mass, age, time, and temperature. The set of numbers a continuous variable takes is typically either 'every possible number' or 'just positive numbers'. For example, a concentration may be very large or very small, but it is strictly positive, whereas a change in concentration can be positive or negative.
- Discrete numeric variable: Observations can take a value based on a count from a set of whole values, e.g. 1, 2, 3, 4, 5, and so on. A discrete variable cannot take the value of a fraction between one value and the next closest value. Examples of discrete variables include the number of individuals in a population, the number of offspring produced ('reproductive fitness'), and the number of infected individuals in an experiment. All of these are measured as whole units.

Categorical variables take values that describe a characteristic of a data unit,

like 'what type' or 'which category'. Categorical variables fall into mutually exclusive (in one category or in another) and exhaustive (include all possible options) categories. Categorical variables are qualitative variables and tend to be represented by a non-numeric value; the data collected for a categorical variable are called qualitative data. Categorical variables may be further described as ordinal or nominal:

- Ordinal variable: Categories can be logically ordered or ranked. The categories associated with ordinal variables can be ranked higher or lower than another but do not necessarily establish a numeric difference between each category. Examples of ordinal categorical variables include academic grades (e.g. A, B, C) and size classes (e.g. small, medium, large).
- Nominal variable: Categories cannot be organised into a logical sequence. Examples of nominal categorical variables include sex (see *C. elegans* example), human blood group (A, B, AB and O), genotype (e.g. AA, Aa, aa), experimental conditions (e.g. control vs enhanced nutrition), and mortality status (alive vs dead).



Do not use numbers to classify categorical variables

Be careful when classifying variables. It can be dangerous to assume that just because a numerical scheme has been used to describe a variable, it must not be categorical. There is nothing to stop someone from using numbers to describe a categorical variable (e.g. $C.\ elegans$ sex: Male = 1, Hermaphrodite = 2). That said, although we can use numbers to describe categories, it does not mean we should. Using numbers gets confusing and can lead to mistakes. It is much clearer to use a non-numeric recording scheme based on words or acronyms to record categorical variables (e.g. $C.\ elegans$ sex: Male = "Male", Hermaphrodite = "Herm").

12.2.2 Ratio vs interval scales

A second way of classifying numeric variables (**not** categorical variables) relates to the scale they are measured on. The measurement scale is important because it determines how we interpret things like differences, ratios, and variability.

• Ratio scale: This scale does possess a meaningful zero value. It gets its name from the fact that a measurement on this scale represents a ratio between a measure of the magnitude of a quantity and a unit of the same kind. What this means in simple terms is that it is meaningful to say that something is "twice as ..." as something else when working with a variable measured on a ratio scales. Ratio scales most often appear when we work with physical quantities. For example, we can say that one tree is twice as big as another or that one elephant has twice the mass of another because length and mass are measured on ratio scales.

• Interval scale: This allows for the degree of difference between measurements, but not the ratio between them. This kind of scale does not have a unique, non-arbitrary zero value. A good example of an interval scale is the date, which we measure relative to an arbitrary epoch (e.g. AD). It makes no sense to say that 2000 AD is twice as long as 1000 AD. However, we can compare ratios of differences on an interval scale. For example, it does make sense to talk about the amount of time between two dates, i.e. we can to say that twice as much time has passed since the epoch in 2000 AD versus 1000 AD.

Keep in mind that the distinction between ratio and interval scales is a property of the measurement scale, not the thing being measured. For example, when we measure temperature in $^{\circ}$ C we're working on an interval scale defined relative to the freezing and boiling temperatures of water under standard conditions. It doesn't make any sense to say that 30° C is twice as hot as 15° C. However, if we measured the same two temperatures on the Kelvin scale, it is meaningful to say that 303.2K is 1.05 times hotter than 288.2K. This is because the Kelvin scale is relative to a true zero (absolute zero).

12.3 Populations and samples

Whenever we collect data, we are almost always working with a sample drawn from a wider population. We might want to know something about that population, but since it is impossible to study the whole population, we study the properties of one or more samples instead. For example, a physiologist might want to know how exercise affects lung function. Since they obviously can't study every person on the planet, they have to study a small sample of people.

We mention the distinction between populations and samples because EDA is concerned with exploring the properties of samples. EDA aims to characterise a sample without trying to infer too much about the wider population from which it is derived. Learning about populations is the basis of much of statistics. That topic is best dealt with in a dedicated statistics book—not this one.

12.3.1 Sample distributions

When we say that 'EDA is concerned with exploring the properties samples,' we actually mean that EDA is concerned with the properties of **variables** in one or more samples. We can be even more precise—the property we are alluding to is the variable's **sample distribution**. The distribution of a variable describes the relative frequency with which different values occur. This is best understood by example...

Imagine we took a sample of undergraduates and measured their height. The majority of students would be around about 1.7m tall, even though there would be plenty of variation among students. Men would tend to be slightly taller than women, and very small or very tall people would be rare. We know from

experience that no one in this sample would be over 3 meters tall. These are all statements about a hypothetical sample distribution of undergraduate heights.

12.3.2 Associations

We've been talking about statistical variables as though we study them one at a time. However, most interesting samples involve more than one variable, and the goal of the ensuing data analysis is to understand associations among those variables. These associations might involve the same (e.g. numeric vs numeric) or different (e.g. numeric vs categorical) types of variable. Whatever the details, the aim of EDA is often to understand how each variable in a sample relates to the others.

12.4 Types of EDA

Exploratory data analysis involves questions such as:

- What are the most common values of the variable?
- How much do observations differ from one another?
- Is one variable associated with another?

Rather than describe the answers to such questions in purely verbal terms, as we did above, EDA relies on descriptive statistics and graphical summaries:

- 1. **Descriptive statistics**. Descriptive statistics are used to quantify the basic features of a sample distribution. They provide numerical summaries about the sample that can be used to make comparisons and draw preliminary conclusions. For example, we often use the mean to summarise the 'most likely' values of a variable.
- 2. Graphical summaries. Descriptive statistics are not much use on their own—a few numbers can't capture every aspect of a distribution. Graphical summaries are a powerful complement to descriptive statistics because they capture a lot of information about a sample in a way that is easy for people to understand.

Descriptive statistics are important, but they are not the main focus of this book. The next few chapters will set out how to construct a range of useful graphical data summaries. Indeed, this book's overarching goal is to demonstrate a workflow that starts with raw data and ends in one or more exploratory plots. We do need to know a bit about descriptive statistics to understand some of those plots. To that end, we'll finish this chapter with a quick survey of descriptive statistics that will pop up later.

12.5 A primer of descriptive statistics

12.5.1 Numeric variables

So far, we've only mentioned the properties of sample distributions in very general terms—using phrases like 'most common values' and 'the range of the data'—without really saying what we mean. Statisticians have devised a set of terms to describe distributions and various descriptive statistics to quantify these. The two that matter most for numeric variables are the **central tendency** and the **dispersion**:

- A measure of **central tendency** describes a typical value of a distribution. Most people know at least one measure of central tendency. The "average" that they calculated at school is the arithmetic mean of a sample. There are many different measures of central tendency, each with its own pros and cons. Take a look at the Wikipedia page to see the most common ones. Among these, the median is the one that is used most often in exploratory analyses.
- A measure of **dispersion** describes how spread out a distribution is. Dispersion measures quantify the variability or scatter of a variable. If one distribution is more dispersed than another, it means that it encompasses a wider range of values. What this means in practice depends on the kind of measure we're working with. We tend to focus on the variance, the standard deviation, and the interquartile range. There are many others, though.



Beyond central tendency and dispersion

Another important aspect of a distribution is its **skewness** (a.k.a. 'skew'). There are many different ways to quantify skewness. Unfortunately, these are quite difficult to make sense of. For now, we only need to understand what skewness means in qualitative terms. Skewness refers to the (a)symmetry of a distribution. When we talk about distributions with high skew, we mean they are very asymmetric.

12.5.1.1 Central tendency

The central tendency of a numeric variable's sample distribution is typically described using either the arithmetic mean or the median. The **arithmetic** mean of a sample is 'the mean' that everyone learns at school¹. Most people have calculated the mean by hand at some point. As R users, we can use the mean function to calculate the arithmetic mean if we need it. For example, this will calculate the mean body mass in the penguins data set:

¹People often just say 'the mean' when referring to the arithmetic sample mean. This is fine, but keep in mind that there are other kinds of mean , such as the harmonic mean and the geometric mean.

mean(penguins\$body_mass_g, na.rm = TRUE)

[1] 4201.754

Remember—we used na.rm = TRUE here because body_mass_g contains a small number of missing (NA) values. This calculation tells us the arithmetic mean of body mass is 4202 grams, i.e. in some sense, the most common body mass is about 4202 grams.

One limitation of the arithmetic mean is that it is affected by the shape of a distribution. This is why, for example, it does not make much sense to look at the mean income of workers to get a sense of what a 'typical' person earns. Income distribution are highly skewed, such that a few people receive very large salaries compared to the vast majority of the population. Those few who earn very good salaries tend to shift the mean well past anything that is really 'typical'.

Because the sample mean is sensitive to the shape of a distribution in this way, we often prefer to use a more robust measure of central tendency—the **sample median**. The median of a sample is the value that separates the upper half from the lower half. We can find the sample median with the median function in R:

median(penguins\$body_mass_g, na.rm = TRUE)

[1] 4050

This tells us that the arithmetic mean of body mass in the penguins data is 4050 grams. This is less than the mean, reflecting the fact that the body mass distribution is somewhat asymmetric.



What about 'the mode'?

The **mode** of a variable's distribution is simply the value that is most likely to occur. This is a simple idea. Unfortunately, it is often difficult to estimate the mode from a sample. Nonetheless, it is important to know what the mode represents, because the concept is useful even when the actual value is hard to estimate.

12.5.1.2 Dispersion

There are many ways to quantify the dispersion of a sample distribution. The most important quantities from the standpoint of statistics are the sample variance and standard deviation. The sample variance (s^2) is the average squared deviations of each point from the mean. Variances are non-negative. The larger the variance, the more observations are spread out around the mean. A variance of zero only occurs if all values are identical.

We won't waste time showing the formula because we'll never actually need to use it directly. As usual, R can calculate the sample variance if we need it:

```
var(penguins$body_mass_g, na.rm = TRUE)
```

[1] 643131.1

That's a big number. What does it mean? Is it 'big' or is it 'small'? No idea. That's the problem with variances—they are difficult to interpret because their calculation involves squared deviations. The variance is an important quantity in statistics because many common tools use changes in the variance as a basis for statistical tests. However, variances seldom feature in EDA because they are so hard to interpret.

A somewhat better descriptive statistic is to describe sample dispersion is a closely related quantity called the **standard deviation** of the sample, usually denoted s. The standard deviation is equal to the square root of the variance. We calculate it using the sd function:

```
sd(penguins$body_mass_g, na.rm = TRUE)
```

[1] 801.9545

Why do we prefer the standard deviation over the variance? Because it is the square root of the variance, it operates on the same scale as the variable it summarises. This means it reflects the dispersion we perceive in the data. The sample standard deviation is not without problems, though. Like the sample mean, it is sensitive to the shape of a distribution and the presence of outliers.

A measure of dispersion that is robust to these kinds of problems is the **interquartile range**. The interquartile range (IQR) is defined as the difference between the third and first quartile (see box). The IQR contains the middle 50% of the values of a variable. The more spread out the data, the larger the IQR. People prefer IQR to measure dispersion for exploratory work because it only depends on the 'middle' of a distribution. This makes it robust to the presence of outliers.

We can use the IQR function to find the interquartile range of the body mass variable:

```
IQR(penguins$body_mass_g, na.rm = TRUE)
```

[1] 1200

The IQR is used as the basis for a useful data summary plot called a 'box and whiskers' plot. We'll see how to construct this later.



What are quartiles?

We need to know what a quartile is to understand the interquartile range. Three quartiles are defined for any sample. These divide the data into four equal-sized groups, from the set of smallest numbers up to the set of largest numbers. The second quartile (Q_2) is the median, i.e. it divides the data into an upper and lower half. The first quartile (Q_1) is the number that divides the lower 50% of values into two equal-sized groups. The third quartile (Q_3) is the number that divides the upper 50% of values into two equal-sized groups.

12.5.2 Categorical variables

Descriptive statistics of categorical variables aim to quantify specific features of their sample distribution, just as with numeric variables. The general question we need to address is, what are the relative frequencies of different categories? Because categorical variables take a finite number of values, the simplest thing we can do is tabulate the number of occurrences of each type. We can use the **dplyr count** function to do this:

penguins %>% count(species)

```
## # A tibble: 3 x 2
## species n
## <chr> <int>
## 1 Adelie 152
## 2 Chinstrap 68
## 3 Gentoo 124
```

This prints that the number of observations associated with each species in penguins. The summary reveals that the most common species in the data set is the Adelie penguin, followed by the Gentoo and Chinstrap.

Can we quantify the central tendency of a categorical sample distribution? Various measures exist. We can certainly find the **sample mode** of a categorical variable easily enough. This is just the most common category. In the case of the above **species** variable, the mode is obviously Adelie. It is also possible to calculate a **sample median** of a categorical variable, but only when it is ordinal. Since the median value is the one that lies in the middle of an ordered set of values, it makes no sense to talk about the middle of a set of nominal values that have no inherent order.

What about dispersion? Well, measures of dispersion for categorical variables do exist, but they are not very easy to interpret. They seldom get used in exploratory data analysis so let's not worry about them here.

12.5.3 Associations

Statisticians have devised different ways to quantify an association between variables. The common measures calculate some kind of **correlation coefficient**. The terms 'association' and 'correlation' are closely related, so they are often used interchangeably. Strictly speaking, correlation has a narrower definition: a correlation is defined by a metric (the 'correlation coefficient') that quantifies the degree to which an association tends to a certain pattern.

12.5.3.1 Pairs of numeric variables

A widely used measure of correlation for pairs of numeric variables is **Pearson's correlation coefficient** (a.k.a. Pearson product-moment correlation coefficient, or Pearson's r). Remember, a correlation coefficient quantifies the degree to which an association tends to a certain pattern. Pearson's correlation coefficient is designed to summarise the strength of a **linear** (i.e. straight line) association.

Pearson's correlation coefficient takes a value of 0 if two variables are not linearly associated and a value of +1 or -1 if they are perfectly related and represent a straight line. A positive value indicates that high values in one variable are associated with high values of the second; a negative value indicates that high values of one variable is associated with low values of the second. High values are those that are greater than the mean; low values are those that are less than the mean.

We can use the **cor** function to calculate Pearson's correlation coefficient. For example, the Pearson correlation coefficient between flipper length and body mass is given by:

```
cor(penguins$flipper_length_mm, penguins$body_mass_g, use = "complete.obs")
```

[1] 0.8712018

This is positive, indicating flipper length tends to increase with body mass. It is also quite close to +1, indicating the association is strong. We should interpret Pearson's correlation coefficient with care. Because it is designed to summarise the strength of a **linear** relationship, Pearson's correlation coefficient will mislead when this relationship is curved. If that statement does not make immediate sense, take a look at the famous Anscombe's quartet.



Other measures of correlation

What should we do if we think the relationship between two variables is non-linear? Calculate something called a **rank correlation**. The idea is quite simple. Instead of working with the actual values of each variable, we rank them by value, i.e. sort each variable from lowest to highest and

the assign the labels $1^{\rm st}$, $2^{\rm nd}$, $3^{\rm rd}$, ... to observations. Rank correlations are based on the association among the ranks of two variables.

The two most popular rank correlation coefficients are Spearman's ρ ('rho') and Kendall's τ ('tau'). The differences are minimal:

- Spearman's ρ is a bit more sensitive to outliers in the data.
- Kendall's τ can be slow to calculate for large data sets.

We can either coefficient using the cor function, setting the method argument to the appropriate value: method = "kendall" or method = "spearman". A rank correlation coefficient is interpreted in the same way as Pearson's correlation coefficient. It takes a value of 0 if the ranks are uncorrelated and +/-1 if they are perfectly associated (though not necessarily as a straight line). The sign tells us about the direction of the association.

12.5.3.2 Pairs of categorical variables

Quantifying associations between pairs of categorical variables is not as simple as the numeric case. The general question is, "do different **combinations** of categories seem to be under- or over-represented?" We need to understand which combinations are common and which are rare. The simplest thing we can do is 'cross-tabulate' the number of occurrences (i.e. the 'frequencies') of each combination. The resulting table is called a **contingency table**.

The xtabs function (= 'cross-tabulation') can do this. For example, the frequencies of each penguin species and island combination is given by:

```
xtabs(~ species + island, data = penguins)
```

```
##
               island
## species
                 Biscoe Dream Torgersen
##
     Adelie
                     44
                            56
                                       52
##
     Chinstrap
                      0
                            68
                                        0
                             0
                                        0
##
     Gentoo
                    124
```

The first argument sets the variables to cross-tabulate. xtabs uses R's special formula language, which means we must include that ~ symbol at the beginning. After the ~, we provide the list of variables to cross-tabulate, separated by the + sign. The second argument tells the function which data set to use.

The table above shows us how many observations are associated with each combination of the **species** and **island** categories. This particular case represents a fairly extreme example of (dis)association; the Chinstrap and Gentoo species simply don't occur on certain islands (or perhaps they weren't sampled on those islands for some reason).

What about measures of association such as correlation coefficients? Spearman's ρ and Kendall's τ are designed for numeric variables, but these can also be used to measure the correlation between ordinal variables (Kendall's τ is best). Various measures of association have been constructed for pairs of nominal variables (e.g. Cramér's V). However, none of these are widely used in exploratory data analyses—people tend to stick with graphical tools for categorical data.

Chapter 13

Introduction to ggplot2

One of the main reasons data analysts turn to R is for its strong data visualisation capabilities. The R ecosystem includes many different packages that support data visualisation. The three most widely used are: 1) the base graphics system, which uses the **graphics** package; 2) the **lattice** package; and 3) the **ggplot2** package. Each system has its own strengths and weaknesses:

- Base graphics is part of base R, which means it's always available. It's very flexible and allows us to construct more or less any plot we like. This flexibility comes at a cost, though. While it is certainly easy to get up and running with base graphics—there are specialised functions making common plots—building complex figures quickly becomes time-consuming. We have to write a lot of code to prepare even moderately complex plots, there are many graphical parameters to learn, and many of the standard plotting functions are inconsistent in how they work.
- Deepayan Sarkar developed the lattice package to implement the ideas of Bill Cleveland in his 1993 book, Visualizing Data. The package implements something called Trellis graphics, a very useful approach for graphical exploratory data analysis. Trellis graphics are designed to help us visualise complicated, multiple variable relationships. The lattice package has many "high level" functions to make this process easy. The lattice package is very powerful, but it is hard to master.
- Hadley Wickham developed the **ggplot2** package to implement the ideas in a book called The Grammar of Graphics by Wilkinson (2005). It produces Trellis-like graphics but is quite different from **lattice** in the way it goes about this. It uses its own mini-language to define graphical objects, adopting the language of Wilkinson's book to define these. It takes a little while to learn the basics, but once these have been mastered, it's very easy to produce sophisticated plots with very little R code. The downside of working with **ggplot2** is that it isn't as flexible as base graphics.

We are not going to survey all these plotting systems. It's entirely possible to meet most data visualisation needs by becoming proficient with just one of them. This book focuses on the **ggplot2** package. In many ways, **ggplot2** hits the 'sweet spot' between base graphics and **lattice**. It enables complex visualisations without the need to write many lines of R code, but remains flexible enough to allow the kind of customisation required to produce publication-quality figures.

13.1 The anatomy of ggplot2

The easiest way to learn **ggplot2** is by using it. However, before we dive into **ggplot2** code, we need to review the essential features of its 'grammar'—the rules of how to specify a graph. This grammar is fairly abstract and won't make much sense on first reading. That is fine. Ideas like 'aesthetic mappings' and 'geoms' will start to make sense as we work through various examples.

The design of **ggplot2** reflects Wilkinson's grammar of graphics. A complete **ggplot2** object is defined by a combination of:

- 1) one or more layers,
- 2) a set of scales (one for each 'aesthetic mapping'),
- 3) a coordinate system (one per plot),
- 4) a **facet** specification (if using a multi-panel plot).

The underlying idea is that we construct a visualisation by defining one or more layers. Each layer is associated with some data and a set of rules for how to display the data. Let's review these components before moving onto the business of actually using **ggplot2**.

13.1.1 Layers

Each layer in a **ggplot2** plot has five different components, though we don't necessarily have to specify all of these because most have some kind of default setting:

- The data. At a minimum, every plot needs some data. Unlike base R graphics, ggplot2 always accepts data in one format, an R data frame (or tibble). Each layer can be associated with its own data set. We don't have to add data to each layer explicitly. If we choose not to specify the data set for a layer, ggplot2 will use the default data (if defined).
- A set of **aesthetic mappings**. These describe how variables in the data are associated with the aesthetic properties of the layer. Aesthetic properties include things we perceive, such as position, colour, and size of the points. Each layer can be associated with its own unique aesthetic mappings. When we choose not to specify these for a layer **ggplot2** will use the defaults (if defined).

- A geometric object (a.k.a. 'geom'). The geom part of layer tells ggplot2 how to represent the information—i.e. it refers to the objects we see on a plot, such as points, lines, bars or even text. Each geom only works with a particular subset of aesthetic mappings. We always have to define a geom when specifying a layer.
- A statistical transformation (a.k.a. 'stat'). A stat takes the raw data and transforms it in some way. A stat allows us to plot useful summaries of our raw data. We won't explicitly use them in this book because we prefer to produce summary figures by first processing the data with dplyr. Nonetheless, the stat facility is often doing useful work in the background for some kinds of plots.
- A position adjustment. These apply small tweaks to the position of layer elements. These are typically used when we need to define how the information for different categories are separated. For example, when making a bar plot, we may need to specify whether bars should be stacked on top of one another (the default) or plotted side-by-side.

13.1.2 Scales

The scale part of a **ggplot2** object controls how the information in a variable is mapped to the aesthetic properties. A scale takes the data and converts it into variation we can perceive, such as an x/y location or the colour and size of points in a plot. The two most important things to understand about scales are:

- 1. A scale must be defined for every aesthetic in a plot. It doesn't make sense to define an aesthetic mapping without a scale because there is no way for **ggplot2** to know how to go from the data to the aesthetics without one.
- 2. When we include two or more layers, they all have to use the same scale for any shared aesthetic mappings. This behaviour is necessary to ensure information is displayed consistently.

If we choose not to explicitly define a scale for an aesthetic **ggplot2** will use a default. This will often be a 'sensible' choice, which means we can get quite a long way with **ggplot2** without ever really understanding scales. We will take a brief look at a few of the more common options, though.

13.1.3 Coordinate system

A ggplot2 coordinate system takes the position of objects (e.g. points and lines) and maps them onto the 2d plane a plot lives on. Most people are already very familiar with the most common coordinate system (even if they didn't realise it). That's the Cartesian coordinate system. This is the one we've all been using since we first constructed a graph with paper and pencil at school. All the most common statistical plots use this coordinate system, so we won't consider any others in this book.

13.1.4 Faceting

The idea behind faceting is very simple. Faceting allows us to break a data set up into subsets according to the unique values of one or more variables and then produce a separate plot for each subset. The result is a multi-panel plot where each panel shares the same layers, scales, etc. The data is the only thing that varies from panel to panel. The result is a kind of 'Trellis plot', similar to those produced by the **lattice** package. Faceting is a very powerful tool that allows us to slice up our data in different ways and understand the relationship between different variables.

13.2 A quick introduction to ggplot2

Now that we've briefly reviewed the **ggplot2** grammar, we can start learning how to use it. The package uses this grammar as the basis of a sort of mini-language within R. It uses functions to specify components like aesthetic mappings and geoms, which are combined with data to define a **ggplot2** graphics object. Once we've constructed a suitable object, we can use it to display our graphic on the computer screen or save it using a common graphics format (e.g. PDF, PNG or JPEG).

Rather than orientating this introduction around each of the key functions, we're going to develop a simple example to help us see how **ggplot2** works. Many of the key ideas about how **ggplot2** works can be taken away from this one example. Hence, it's worth investing the time to understand it—i.e. use the example to understand how the different **ggplot2** functions are related to the grammar outlined above.

Our goal is to produce a simple scatter plot. The scatter plot is one of the most commonly used visualisation tools in the EDA toolbox. A scatter plot uses horizontal and vertical positions (the 'x' and 'y' axes) to visualise pairs of related observations as a series of points in two dimensions. It's designed to show how one numeric variable is associated with another. We'll use the penguins data to construct the scatter plot. The questions we want to explore are:

- what is the relationship between bill depth and bill length, and
- how does this vary in relation to other variables?

13.2.1 Making a start

We will begin our work with **ggplot2** by setting up a minimal **graphical object**. That's a job for the **ggplot** function. Using **ggplot** without any arguments builds an empty graphical object:

```
# construct empty ggplot2 graphical object
plot_obj <- ggplot()</pre>
```

print summary of empty graphical object

This constructs the skeleton object and assigns it a name, plot_obj. We can use the summary function to inspect the object to find out a bit more about it:

```
summary(plot obj)
## data: [x]
## faceting: <ggproto object: Class FacetNull, Facet, gg>
##
       compute_layout: function
##
       draw_back: function
       draw_front: function
##
##
       draw_labels: function
##
       draw_panels: function
##
       finish_data: function
##
       init_scales: function
##
       map_data: function
##
       params: list
##
       setup_data: function
##
       setup params: function
##
       shrink: TRUE
##
       train scales: function
```

The output of summary is quite verbose, but the important parts are near the top, just before the faceting: section. In this case, the 'important part' is basically empty. All we did was set up an empty graphical object—there are no data, aesthetic mapping, layers, etc associated with plot_obj.

super: <ggproto object: Class FacetNull, Facet, gg>

It is not necessary to inspect the insides of every **ggplot2** object with **summary**. However, it can be instructive to do this when first learning about the package.



##

ggplot2 vs. ggplot

vars: function

Notice that the package is called **ggplot2**, but the actual function that does the work of setting up the graphical object is called **ggplot**. Try not to mix the names up—this is a common source of errors.

How can we improve on this? We could add a default data set. This is easy. We do it by passing the name of a data frame or tibble to ggplot. Let's try doing this with penguins:

```
# construct ggplot2 graphical object with data
plot_obj <- ggplot(penguins)</pre>
```

Then print out the summary of the updated plot_obj:

```
# print summary of updated graphical object
summary(plot_obj)
```

```
## data: species, island, bill_length_mm, bill_depth_mm,
## flipper_length_mm, body_mass_g, sex, year [344x8]
## faceting: <ggproto object: Class FacetNull, Facet, gg>
## ...facet summary suppressed...
```

We suppressed the facet information this time because it takes up a lot of space and we're not interested in it. The important point is that the plot_obj summary now contains some information related to the data. The variables from penguins now comprise the data associated with the plot_obj object.

The next step is to add a default aesthetic mapping to the graphical object. Remember, this describes how variables in the data are mapped to the aesthetic properties of the layer(s).

One way to think about aesthetic mappings is they define what kind of relationships the plot will describe. Since we're making a scatter plot, we need to define mappings for positions on the 'x' and 'y' axes. We want to investigate how bill depth depends on bill length, so we need to associatebill_length_mm with the x position and bill_depth_mm with the y position.

We define an aesthetic mapping with the aes function. One way to use aes is like this:

```
# add aesthetic mappings to graphical object
plot_obj <- plot_obj + aes(x = bill_length_mm, y = bill_depth_mm)</pre>
```

This little snippet of R code looks odd at first glance. There are a couple of things to take away from this:

- 1. We can 'add' the aesthetic mapping to the plot_obj object using the + operator. This has nothing to do with arithmetic. The ggplot2 package uses some clever programming tricks to redefine the way + works so that it can be used to combine graphical objects. It takes a bit of getting used to but this is useful because it makes building a plot from the components of the grammar very natural.
- 2. The second thing to notice is that an aesthetic mapping is defined by one or more name-value pairs, specified as arguments of aes. The names on the left-hand side of each = refer to the properties of our graphical object (e.g. the 'x' and 'y' positions). The values on right-hand side refer to variable names in the data we want to associate with these properties.

Notice that we overwrote the original plot_obj object with the updated version using the assignment operator. We could have created a distinct object, but there's usually no need to do this.

Once again, we can inspect the result using summary:

```
# print summary of updated graphical object
summary(plot_obj)
```

```
## data: species, island, bill_length_mm, bill_depth_mm,
## flipper_length_mm, body_mass_g, sex, year [344x8]
## mapping: x = ~bill_length_mm, y = ~bill_depth_mm
## faceting: <ggproto object: Class FacetNull, Facet, gg>
## ...facet summary suppressed...
```

The data (data:) from the original plot_obj are still there, but now we can also see that two default mappings (mapping:) have been defined for the x- and y-axis positions. We have successfully used the ggplot and aes functions to set up a graphical object with both default data and aesthetic mappings.

Any layers that we now add will use these data and mappings unless we choose to override them by specifying different options.

We now need to specify a layer to tell **ggplot2** how to visualise the data. Remember, each layer has five components: data, aesthetic mappings, a geom, a stat and a position adjustment. Since we have already set up the default data and aesthetic mappings, there's no need to define these again—**ggplot2** will use the defaults if we leave them out of the layer definition. This leaves the geom, stat and position adjustment.

What kind of geom do we need? A scatter plot allows us to explore a relationship as a series of points. That means we need to add a layer that uses the 'point' geom. Simple.

What about the stat and position? Not so simple. These are difficult to explain without drilling down into how **ggplot2** works. A key insight is that both the stat and the position adjustment change the data somehow before plotting it. When we want to stop **ggplot2** from doing anything to our data, the keyword is 'identity'. We use this value whenever we want **ggplot2** to plot the data without modification.

We will examine the easy way to add a layer in a moment. However, we'll use the long-winded approach first because this reveals what happens whenever we build a **ggplot2** object. The general function for adding a layer is simply called layer. Here's how it works in its most basic usage:

```
# add layer to graphical object -- using points geom
plot_obj <- plot_obj + layer(geom = "point", stat = "identity", position = "identity")</pre>
```

This adds a layer to the existing plot_obj object with the layer function and overwrites the old version. Again, we add the new component using the + symbol. We set three arguments of the layer function:

- 1. define the **geom**: the name of this argument was **geom** and the value assigned to it was "point".
- 2. define the **stat**: the name of this argument was **stat** and the value assigned to it was "identity".
- 3. define the **position adjustment**: the name of this argument was position and the value assigned to it was "identity".

Let's review the structure of the resulting graphical object one last time to see what we've achieved:

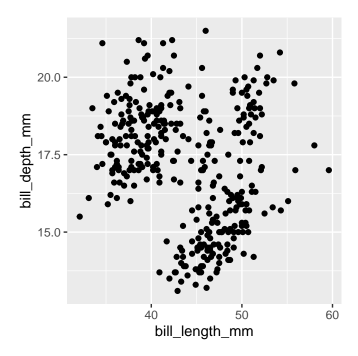
```
# print summary of updated graphical object
summary(plot_obj)
```

The text above the ---- line is the same as before. It summarises the default data and the aesthetic mapping. The text below it summarises the layer we just added. This tells us that layer uses the points geom (geom_point), the identity stat (stat_identity), and the identity position adjustment (position_identity).

Now plot_obj has everything it needs to render a figure. How do we do this? Simply 'print' the graphical object:

```
# 'print' graphical object to show it
print(plot_obj)
```

Warning: Removed 2 rows containing missing values (geom_point).



That's it! We have produced a scatter plot showing how bill depth is associated with bill length. It seems a little odd to show a plot by 'printing' it but that's just how **ggplot2** works. It soon starts to feel natural with practise.

Notice that **ggplot2** printed a warning. That is nothing to worry about. It is just letting us know a couple of rows in the data were ignored because they contained missing values. We will suppress that message from now on because it gets a bit irritating.

Here's a quick summary of what we did, all in one place:

```
# step 1. set up the skeleton graphical object with a default data set
plot_obj <- ggplot(penguins)
# step 2. add the default aesthetic mappings
plot_obj <- plot_obj + aes(x = bill_length_mm, y = bill_depth_mm)
# step 3. specify the layer we want to use
plot_obj <- plot_obj + layer(geom = "point", stat = "identity", position = "identity")
# step 4. show the plot
print(plot_obj)</pre>
```

Λ

Don't use this workflow!

It's possible to construct any **ggplot2** visualisation using the workflow outlined in this subsection, but **this is not the recommended approach**.

The workflow we adopted here was used to reveal how the grammar works, rather than its efficiency. A more concise, standard approach to using **gg-plot2** is outlined next. Use that for real-world analysis.

13.3 A standard way of using ggplot2

The **ggplot2** package is quite flexible, which means we can specify a visualisation in more than one way. To keep life simple, we're going to adopt a consistent workflow from now on. This won't reveal the full array of **ggplot2** tricks, but it is sufficient to construct a wide range of standard visualisations. Let's make the same bill depth vs. bill length scatter plot again to see the workflow in action.

We began building our **ggplot2** object by setting up a skeleton object with a default data set and then added the default aesthetic mappings. There is a more concise way to achieve the same result:

```
# create graphical object with data and aesthetic mappings
plot_obj <- ggplot(penguins, aes(x = bill_length_mm, y = bill_depth_mm))</pre>
```

In this form, the aes function appears inside ggplot as a second argument. This sets up a graphical object with default data and aesthetic mappings in a single step. We will always use this approach from now on.

The next step adds a layer. We saw that the layer function could be used to construct one from its component parts. However, **ggplot2** provides many convenience functions that construct layers according to the type of geom they need. They all look like this: **geom_TYPE**, where **TYPE** stands for the name of the geom we want to use. For example, a point geom is specified using **geom_point**. Using this function, an alternative to the last line of the example is therefore:

```
# use geom_point to add a points layer to graphical object
plot_obj <- plot_obj + geom_point()</pre>
```

We didn't have to specify the stat or the position adjustment components of the layer because the <code>geom_TYPE</code> functions all use reasonable defaults. These can be overridden if needed, but most of the time, there's no need to do this. This way of defining a layer is much simpler and less error-prone than the manual <code>layer</code> method. We will always use the <code>geom_TYPE</code> method from now on.

There's one last trick we need to learn to use **ggplot2** efficiently. We've been building a plot object in several steps, giving the intermediates the name plot_obj, and then manually printing the object to display it when it's ready. This is useful if we want to make different versions of the same plot. However, we very often just want to build the plot and display it in one go. This is done by combining everything with + and printing the resulting object directly:

```
# display bill morphology scatter plot **in one step**
ggplot(penguins,
    aes(x = bill_length_mm, y = bill_depth_mm)) +
geom_point()
```

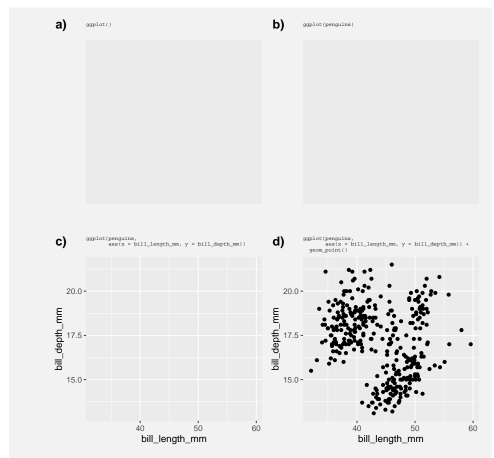
That code builds the **ggplot2** graphical object and renders it in one go. We didn't even have to use **print** to generate the output. There is a lot is going on in the background, but this small snippet of R code contains everything **ggplot2** needs to construct and display a simple scatter plot.

What does this scatter plot actually tell us about penguin bill morphology? It seems to suggest there isn't much of a relationship between bill depth and bill length. Does that seem sensible? Maybe we need a more informative plot.



Plotting incomplete ggplot2 objects

It is instructive to see what happens when we render an incomplete graphical object. The four panels below show the output produced by a) an empty **ggplot2** object, b) a partially complete **ggplot2** object with only data, c) a partially complete **ggplot2** object with data and aesthetics, and d) a complete **ggplot2** object with data, aesthetics and a geom. The actual code to make each one is shown in the title area.



We can see that **ggplot2** will try to plot something with the information it has, but we only end up with an informative plot when we specify data, aesthetic mappings, and a geom. That is the minimum we need to specify to arrive at a useful plot.

13.3.1 How should we format ggplot2 code?

Take another look at that last example. We split the **ggplot2** definition over two lines, placing each function on its own line. R doesn't care about those extra newlines. As long as we put a + at the end of a line R will assume the next line is part of the same plot definition. Splitting the different parts of a graphical object definition across lines like this makes everything more readable and helps us spot errors.

Once we split the definition across lines we can place comments between the lines of **ggplot2** code. For example, we could add comments to state what the **aes** and **geom_point** parts are doing:

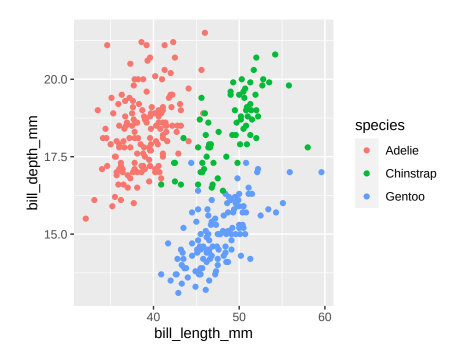
It's a very good idea to format and document **ggplot2** code in this way. That way, when we come back to it after a while, we can remember what we were trying to achieve! We will always use these conventions from now on.

13.4 Increasing the information density...

One of the great strengths of **ggplot2** is the ease with which it allows us to incorporate information from several variables into a single plot. So far, we have made a simple two-variable scatter plot to examine bill morphology—the bill depth-length relationship. There are clearly other variables in the **penguins** data set that might influence this relationship. In this section, we will highlight three approaches for including additional variables in a visualisation. That is, we will see how to increase the **information density** of a plot.

13.4.1 ...via aesthetic mappings

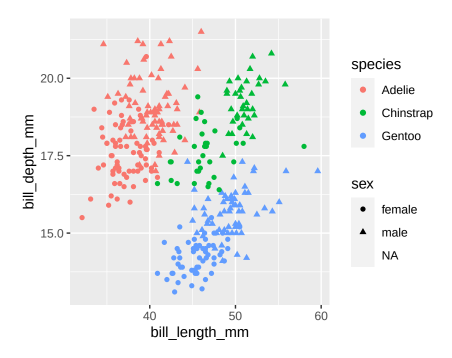
The most straightforward way to increase the information in a plot is by mapping a new variable to an unused aesthetic. For example, how might we learn whether the bill depth-length relationship varies by penguin species? We need to include information in the species variable in our scatter plot somehow. One option is to map the species to the colour aesthetic, so that the colour of the points correspond to different species. We do this by altering the aes part of the plot specification:



Individual points are now coloured according to the species they belong to. Notice **ggplot2** also adds a legend. This plot shows that bill morphology is reasonably species-specific. Separating things by species suggests a (mild) positive association between bill length and bill depth within species. This was not apparent before because it was hidden by the among species pattern.

We put the aes argument on a new line in this example because the first line was getting a bit long. This is only a readability thing—the aes part still belongs to ggplot.

We could certainly improve this visualisation. Nonetheless, it illustrates an important concept: we can add information to a plot by mapping additional variables to new aesthetics. There is nothing to stop us from using different aesthetics if we wanted to squeeze even more information into this plot. For example, we could map the sex variable (sex) to the point shape using shape = sex inside aes:



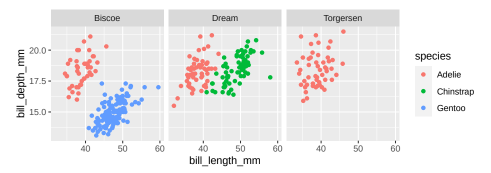
13.4.2 ...via facets

A second way to increase the amount of information on display is by making separate plots for meaningful subsets of the data. We can use the **faceting** facility of **ggplot2** to do this. Faceting allows us to define subsets of data according to the values of one or more variables and produce a separate plot for each subset, all without having to write much R code.

Faceting operates on the whole figure, which means we can't apply it by changing the properties of a layer. Instead, we have to use a new function to add the faceting information. There are two different ways to facet in **ggplot2**:

- facet_wrap forms a matrix of panels by wrapping the 1d sequence of panels into a 2d matrix with rows and columns. It is typically used with a single categorical faceting variable (though it works with two or more faceting variables).
- facet_grid forms a 2d matrix of panels defined by row and column variables. It is typically used when we have two or more categorical variables, and all combinations of the variables exist in the data.

This is best understood by example. What if we want to see how the bill morphology varies across islands? Here's how we split things up by island using the facet_wrap function:



That first vars(island) argument of $facet_wrap$ says to split up the data set according to the values of island. Simple. Notice that the panels share the same scales for the 'x' and 'y' axes. Using a common scale makes it easy to compare bill morphology across islands, but this can also be changed if necessary. For example, setting scales = "free" would ensure each panel uses its own x/y scale.

The plot indicates that bill morphology is roughly invariant across the three islands. It also inadvertently reveals something else about these penguins—the Gentoo and Chinstrap species are only found on a single island, which is different for each one, whereas Adelie penguins are found on all three islands. That's one reason exploratory analysis is so important—it throws up unexpected findings.



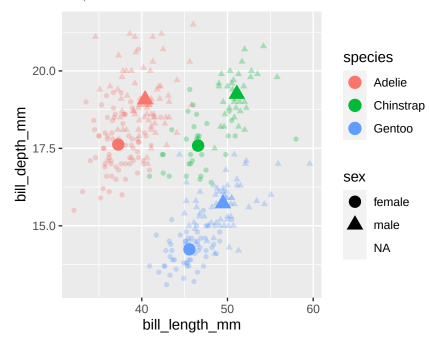
Don't forget the vars part!

We have to wrap the name of the faceting variable with the vars function in the facet_wrap specification. Trust us, facet_wrap won't work without it.

13.4.3 ...via multiple layers

A third way to add additional information to a plot is by including multiple layers. So far, we have only seen examples with a single layer—only one <code>geom_function</code> is involved in creating the plot. There is no reason we can't add multiple

layers to a plot by 'adding' two or more geom_ functions. We're not yet in a position to demonstrate how to make such multi-layer plots because they require additional skills covered in later chapters. However, as a taster, here is an example of the kind of thing we can do with this idea (we have deliberately hidden the code):



This shows how bill morphology varies by penguin species and sex, but now these relationships are summarised in two points layers. The first layer displays the raw data (small transparent points). The second layer adds the species/sex-specific means (large solid points). This places the average differences in the context of the overall variation in the raw data. Nice!

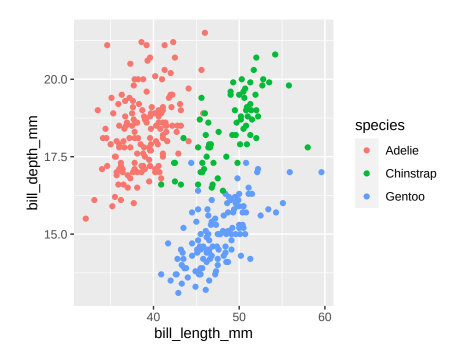
Chapter 14

Customising plots

The default formatting used by **ggplot2** has been carefully chosen to ensure the information in a plot is easy to discern. For example, by default, plots include a pale grey background and include grid lines. This is designed to emphasise the data while supporting comparisons. The grey highlights colour differences and ensures the grid lines have little visual impact beyond aiding in the assessment of position. Perfect for exploratory analysis!

Although we can justify these sorts of choices on the grounds they improve the readability of a plot, the fact remains they are somewhat unconventional and not much-loved by many users. For this reason, we often need to change the appearance of a plot before we include it in a document or presentation—e.g. most published figures use a white, rather than grey, background.

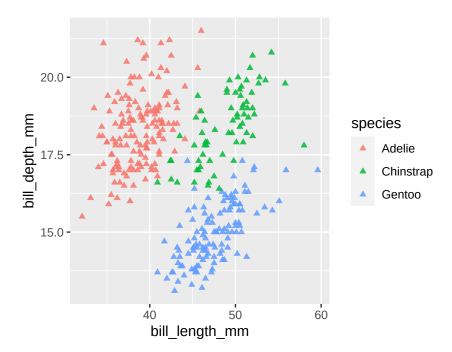
This chapter will demonstrate how to customise **ggplot2** plots. We are not going to cover every possible permutation. That would need its own book. Instead, we will explore the main routes to customisation so that we can use these as we review different visualisations later. Using the **penguins** data once again, we'll work on improving the following scatter plot from the Introduction to **ggplot2** chapter:



14.1 Geom properties

One common way to change a plot's appearance is by altering the properties of one or more geometric objects, such as points, lines, bars or polygons. How do we change the properties of the geom associated with a particular layer? We set those properties by specifying the appropriate arguments in the defining geom_NAME function.

For example, how might we change properties like the size or shape of points in our scatter plot? We used the <code>geom_point</code> function to add points to the scatter plot, which means we have to set the arguments of <code>geom_point</code> to change the point properties. We can use this idea to rebuild the example scatter plot, setting the shape, size and transparency of points:



The point shape is set with the **shape** argument. There are a few different ways to specify point shapes in R. We used the numeric coding system in this example (17 = triangle). Unfortunately, the numeric codes aren't at all intuitive. Here are the more common ones:

The point size is specified by the **size** argument. This has a baseline default value of 1. We assigned **size** a value of 1.5, thereby increasing the point size relative to the default. We could make the points smaller by using a value less than 1.

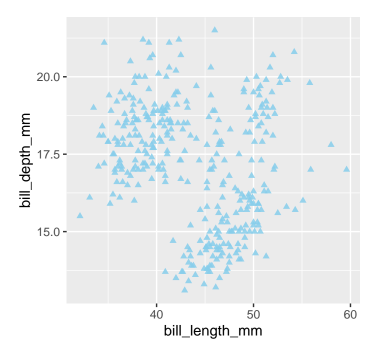
We made the points semi-transparent by setting the value of the alpha argument to be less than 1. In graphical systems, the 'alpha channel' specifies transparency—a value of 0 is taken to mean 'completely invisible', and a value of 1 means 'completely opaque'.

There are other arguments—such as fill and colour—that can also be used to adjust the way the points are rendered. We'll look at these later.

The key message to take away from this customisation example is this—if we need to alter the properties of a geometric object (points, lines, ect), we do so by specifying the appropriate arguments in the <code>geom_NAME</code> function that defines the layer it belongs to.

14.1.1 Relationship between aesthetic mappings and geom properties

In the previous chapter, we saw that we could introduce information into a plot by setting aesthetic mappings in the aes part of its specification. In the scatter plot example, we included information about species identity by mapping the species variable to the colour aesthetic. What happens if we also try to set the colour within the geom_point part?



All the points are now one colour ('sky blue') which means we have lost the species information. What does this example demonstrate? When we set the properties of a geom, this will override any aesthetic mappings that conflict with our choice of customisation. Try to remember that—failing to do so is a good way to become frustrated.



Built-in colours in R

There is nothing special about 'skyblue' other than the fact that it is colour name known to R. There are over 650 built-in colour names. To see them, use a function called **colours** to print them all to the Console. Here are 25 selected at random:

```
sample(colours(), size = 25, replace = FALSE)
    [1] "slateblue3"
                         "lavender"
                                          "gray8"
                                                            "pink"
##
    [5] "turquoise3"
                         "slategray3"
                                          "hotpink"
                                                            gray15"
    [9] "tomato4"
                         "grev50"
                                          "khaki1"
                                                            "grev65"
## [13] "darkslategray" "royalblue"
                                          "gray29"
                                                            "grey64"
        "grey6"
  [17]
                         "palegreen3"
                                          "lightyellow"
                                                            "firebrick1"
                         "darkorange1"
                                          "darkorchid1"
                                                            "goldenrod1"
## [21] "gray81"
## [25] "violetred2"
```

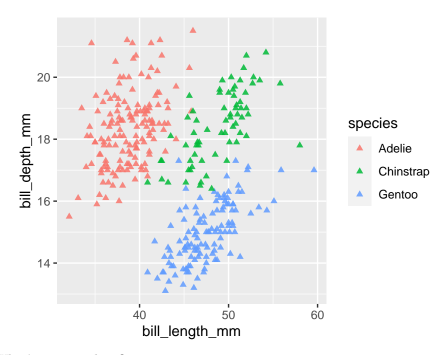
14.2 Plot scales

Setting the arguments of a <code>geom_</code> function applies changes in a layer-specific manner. Other kinds of customisation affect every layer in a plot. Remember what we said about **scales** in the previous chapter:

The scale part of a **ggplot2** object controls how the information in a variable is mapped to the aesthetic properties. A scale takes the data and converts it into variation we can perceive, such as an x/y location or the colour and size of points in a plot.

Every aesthetic mapping has a scale associated with it. We adjust 'how the information in a variable is mapped to the aesthetic properties' by changing its corresponding scale.

As always, this kind of thing is best understood by example. Let's adjust the scale associated with the y-axis aesthetic by increasing the number of the horizontal lines and their accompanying labels (the 'guides').



What's going on here?

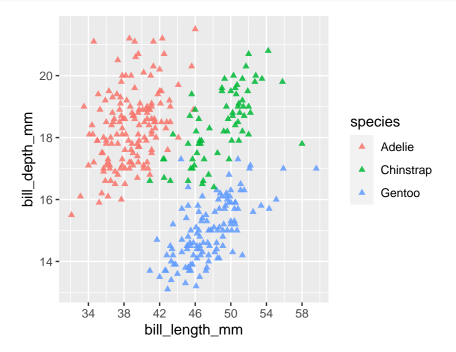
- The aesthetic we wanted to alter was the y-axis position. The y-axis
 in this particular plot is associated with a continuous scale because
 bill_depth_mm is a numeric variable. This means we have to apply the
 scale_y_continuous function to tweak the position of the y-axis guides.
- The breaks argument of scale_y_continuous takes a numeric vector to specify where the guides should be drawn. We used the base R seq function to set up a numeric vector containing the sequence: 14, 16, 18 and 20. That tells gglot2 to place the guides at 14, 16, 18 and 20 on the y-axis.

The functions that adjust a scale all have the general form scale_AES_TYP, where:

- the AES bit in the name refers to the relevant aesthetic, and
- the TYP part refers to the type of scale we want to apply.

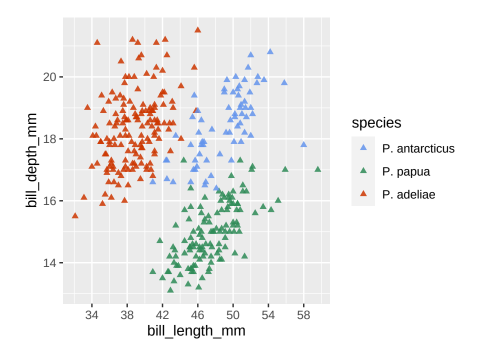
Each kind of scale_AES_TYP function has its own set of arguments to control how the scale works. For example, the breaks argument of scale_y_continuous uses a numeric vector to specify where the guides should be drawn. We can do much fancier things if we want to. For example, the trans argument can be used to apply a particular transformation to an axis, such as a logarithm.

Predictably, there is also a scale_x_continuous function. We can use this in exactly the same way to control where the guides on the x-axis appear:



The key message to take away from this is that every aesthetic mapping has a scale associated with it. If we want to change how the information associated with an aesthetic mapping is displayed we have to change the corresponding scale. That suggests we can change the way point colours are associated with the species variable in our example by using one of the scale_colour_YY functions. That is correct:

```
# customise x- and y-axis grid line and label locations
scale_x_continuous(breaks = seq(34, 62, by = 4)) +
scale_y_continuous(breaks = seq(14, 20, by = 2)) +
# customise the species colours ('value') and labels
scale_color_manual(
   limits = c("Chinstrap", "Gentoo", "Adelie"),
   values = c("cornflowerblue", "seagreen", "orangered3"),
   labels = c(Chinstrap = "P. antarcticus", Gentoo = "P. papua", Adelie = "P. adeliae")
```



This uses scale_color_manual to achieve three things:

- it defines the order to display the three species in the legend via limits,
- it sets up a custom colour palette via values, and
- it alters the category labels via labels (so that their scientific names are used).

Scales are one of the hardest aspects of **ggplot2** to master. There are many different scales, each controlled by its own unique set of arguments. Fortunately, the defaults used by **ggplot2** are often good enough to arrive at a reasonable plot without having to manipulate the scales.

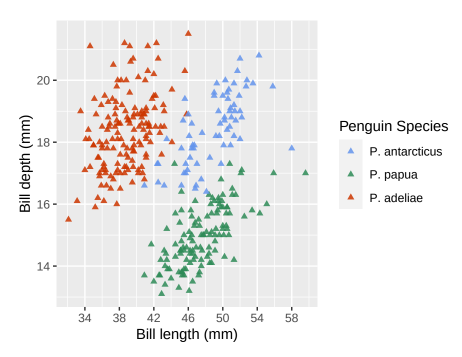
14.3. LABELS 157

14.3 Labels

What else might we like to tweak? Look at the x and y-axis labels. These are the names of the variables used in the x/y aesthetic mappings. These labels aren't too bad, but they could be nicer to look at. We know 'bill_depth_mm' stands for 'bill depth (measured in mm)', so why not use an axis label more like that?

The axes labels are a feature of the whole plot—i.e. they do not belong to a particular layer. This means we can't alter axis labels by passing arguments to the function that built a layer (geom_point). We have to use the labelling function: labs. This sets the label of any mapped aesthetics. For example, to set the labels of the x and y axes and the label associated with the colour mapping (i.e. the species legend title), use:

```
# display bill morphology scatter plot
ggplot(penguins,
       # bill depth (y) vs bill length (x) by species (colour)
       aes(x = bill_length_mm, y = bill_depth_mm, colour = species)) +
  # add points layer and customise appearance of points
  geom_point(shape = 17, size = 1.5, alpha = 0.85) +
  # customise x- and y-axis grid line and label locations
  scale_x_continuous(breaks = seq(34, 62, by = 4)) +
  scale_y_continuous(breaks = seq(14, 20, by = 2)) +
  # customise the species colours ('value') and labels
  scale_color_manual(
    limits = c("Chinstrap", "Gentoo", "Adelie"),
   values = c("cornflowerblue", "seagreen", "orangered3"),
   labels = c(Chinstrap = "P. antarcticus", Gentoo = "P. papua", Adelie = "P. adeliae")
  ) +
  # set the the x/y-axis labels and legend label using `labs`
  labs(x = "Bill length (mm)", y = "Bill depth (mm)", colour = "Penguin Species")
```



Using labs is easy—if a mapping was set up in aes we can change its label via a 'name = value' construct in labs. That's it.

Finally, another way to set the axis labels is via the xlab and ylab. We tend not to use these because labs deals with all our labelling needs on its own. Why waste effort remembering three things if you only need to remember one?

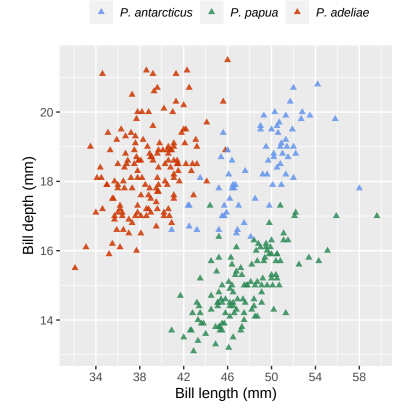
14.4 Themes

The final route to customisation happens via the 'theme' of a plot. We haven't considered the **ggplot2** theme system at all yet. A **ggplot2** theme deals with all the visual elements of a plot that aren't directly handled by adjusting geom properties or scales. These are essentially the 'non-data' parts of a plot—features such as the colour of the plotting region and the grid lines, whether or not those grid lines are even displayed, the position of labels, the font used in labels, and so on.

Here's a short example that adjusts the legend of the example plot:

14.4. THEMES 159

```
# customise x- and y-axis grid line and label locations
scale_x_continuous(breaks = seq(34, 62, by = 4)) +
scale_y_continuous(breaks = seq(14, 20, by = 2)) +
# customise the species colours ('value') and labels
scale_color_manual(
 limits = c("Chinstrap", "Gentoo", "Adelie"),
 values = c("cornflowerblue", "seagreen", "orangered3"),
 labels = c(Chinstrap = "P. antarcticus", Gentoo = "P. papua", Adelie = "P. adeliae")
) +
# use `labs` to set x/y-axis labels and remove legend label
labs(x = "Bill length (mm)", y = "Bill depth (mm)", colour = "") +
# use theme to customise position and text formatting of the legend
theme(
  legend.position = "top",
  legend.text = element_text(face="italic")
)
```



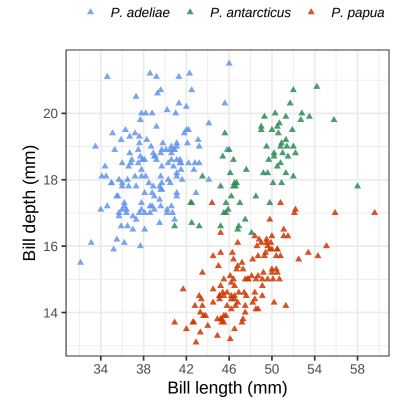
Changes to the theme are specified using... the theme function. We used theme to move the legend to the top of the plot (via legend.position) and show the

species labels in italics (via legend.text). We also got rid of the legend title by setting the colour label to empty character string in labs. We could do that via theme but using labs is simpler.

The **ggplot2** theme system is extremely powerful. Once we know how to use it, we can set up a custom theme and apply it with very little effort. However, that's not an entirely trivial thing to do because there are just so many components of every plot. Most people just google whatever adjustment they want to make when the time comes. Even then, tweaking each individual part of a theme can be very time consuming. Fortunately, there are a range of standard themes built into **ggplot2**. Here's how to apply a built-in theme:

```
# display bill morphology scatter plot
ggplot(penguins,
       # bill depth (y) vs bill length (x) by species (colour)
       aes(x = bill_length_mm, y = bill_depth_mm, colour = species)) +
  # add points layer and customise appearance of points
  geom_point(shape = 17, size = 1.5, alpha = 0.85) +
  # customise x- and y-axis grid line and label locations
  scale_x_continuous(breaks = seq(34, 62, by = 4)) +
  scale_y_continuous(breaks = seq(14, 20, by = 2)) +
  # customise the species colours ('value') and labels
  scale_color_manual(
   values = c("cornflowerblue", "seagreen", "orangered3"),
   labels = c(Adelie = "P. adeliae", Chinstrap = "P. antarcticus", Gentoo = "P. papua
  ) +
  # use `labs` to set x/y-axis labels and remove legend label
  labs(x = "Bill length (mm)", y = "Bill depth (mm)", colour = "") +
  # apply 'black and white' theme
  theme_bw(base_size = 13) +
  # use theme to customise position and text formatting of the legend
  theme(
    legend.position = "top",
    legend.text = element_text(face="italic")
```

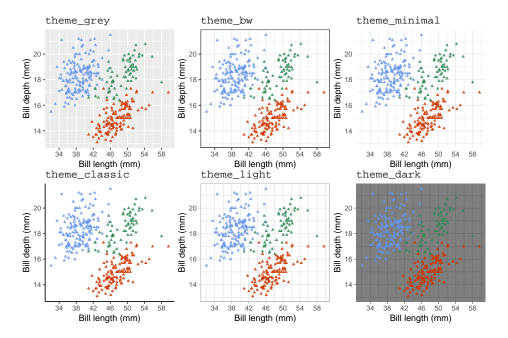
14.4. THEMES 161



This uses + with the theme_bw function to add the built-in 'black and white' theme. This theme dispenses with the grey background that so many people dislike. There are a couple of things to note from this example:

- To apply the changes to the legend position, we had to place the theme part after we set the overall theme using theme_bw. The theme_bw function will override the theme changes if we try to do it the other way round.
- We set the base_size argument of theme_bw. This rescales the text in a plot, leaving the relative size of the different elements unchanged. We did not have to do this, but it is worth showing off because it is such a useful trick.

There aren't that many themes built into **ggplot2**. Here's a quick look at them:



14.5 Advice for making plots

We'll end this section with a bit of advice about making your own plots with ggplot2. Look at the code we ended up with for the final scatter plot above. It's complicated! However, we didn't make that in one go. We built up to it slowly, tweaking one thing at a time. That's the trick—start with a basic skeleton and gradually increase the level of customisation. Begin by getting the aesthetic mappings and the faceting correct so that all the relevant variables are represented. Then introduce one bit of customisation and review the plot to ensure it worked and everything looks right. Keep doing this, one bit of customisation at a time, until the plot looks right.

Chapter 15

Exploring one variable

This chapter will consider how to go about exploring a single variable. Using the penguins data, we'll review some basic visualisations for exploring the sample distribution of numeric and categorical variables.

Exploring numerical variables 15.1

We'll work with flipper length (flipper_length_mm) and body mass (body_mass_g) to demonstrate how to explore numeric variables.

What kind of numeric variable?

The simplest way to explore data is to view it in its raw form. Take a quick look at the first 100 values of flipper length mm and body mass g. We can get these by extracting them with the \$ operator and using the head function:

flipper length —

```
head(penguins$flipper_length_mm, 50)
## [1] 181 186 195 NA 193 190 181 195 193 190 186 180 182 191 198 185 195 197 184
## [20] 194 174 180 189 185 180 187 183 187 172 180 178 178 188 184 195 196 190 180
## [39] 181 184 182 195 186 196 185 190 182 179 190 191
body mass -
head(penguins$body_mass_g, 50)
## [1] 3750 3800 3250
                        NA 3450 3650 3625 4675 3475 4250 3300 3700 3200 3800 4400
## [16] 3700 3450 4500 3325 4200 3400 3600 3800 3950 3800 3550 3200 3150 3950
```

[46] 4600 3425 2975 3450 4150

These are clearly both numeric variables, but we can say a bit more:

- They are both measured on a ratio scale. Zero really is zero, meaning that it makes sense to say that a length of 200 millimetres is twice as long as 100 millimetres or that 6000 grams is twice the mass of 3000 grams.
- They are continuous variables. Think about the possible values flipper length and body mass can take. A length of 200.52 millimetres and a mass of 3000.71 grams are both perfectly reasonable, so fundamentally, these are continuous variables.

Notice that even though body mass is a continuous variable body_mass_g looks discrete because body mass measurements have been taken to the nearest 25 grams. This reflects methodological limitations—presumably, weighing a live penguin to more than 25 grams precision in the field is challenging. This illustrates an important idea: we can't just look at the values a numeric variable takes to determine whether it is discrete or continuous.

Whether we treat a variable as continuous or discrete is sometimes an analysis decision. We have to decide based on knowledge of a variable's true nature and the measurement process. For example, if we were only able to measure body mass to the nearest 500 grams, we would only "see" a few different body mass categories. It might then be reasonable to treat the body_mass_g variable as an ordinal, categorical variable.

15.1.2 Histograms

We only looked at the first 50 values of the flipper_length_mm and body_mass_g variables because the penguins data set is a bit too big to view everything at once easily. It's hard to say much about the sample distribution of a numeric variable by looking at the raw data because there are so many unique values. Those first 50 are also not representative of the whole sample.

What else might we do? One useful tool is 'binning'. The idea behind binning a numeric variable is very simple. It involves two steps.

- 1. Divide the number line into equal-sized, non-overlapping intervals. These are the 'bins'. We pick the interval size to ensure bins (typically) include at least a few cases.
- 2. We then have to count up the number of times the variable falls inside each bin. The resulting set of counts summarises the sample distribution of the numeric variable.

Binning is very tedious to do by hand. Fortunately, R can do this for us. This shows how to use the cut function with the **dplyr** transmute and count verbs to bin the flipper_length_mm variable into 5 mm intervals:

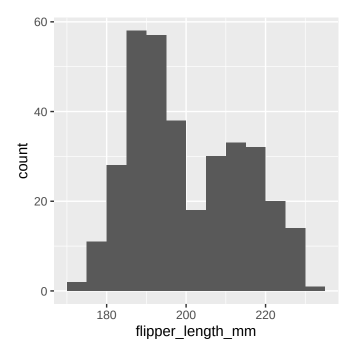
```
penguins %>%
  transmute(bins = cut(flipper_length_mm, breaks = seq(170, 235, by = 5))) %>%
  count(bins)
```

```
## # A tibble: 14 x 2
##
      bins
                     n
##
      <fct>
                 <int>
##
    1 (170,175]
                     2
    2 (175,180]
                    11
##
    3 (180,185]
                     28
    4 (185,190]
                     58
##
    5 (190,195]
##
                    57
##
    6 (195,200]
                     38
    7 (200,205]
##
                    18
    8 (205,210]
                    30
    9 (210,215]
                    33
## 10 (215,220]
                    32
## 11 (220,225]
                     20
## 12 (225,230]
                     14
## 13 (230,235]
                     1
## 14 <NA>
                      2
```

We won't explain how this works because we only need to understand the output. This is a small tibble where the bins column shows the intervals, and the n column gives the associated counts.

What does this tell us? It shows that the most common flipper length is around 190 mm. Values much below 190 mm are rare, but a range of values above this are possible, with higher values becoming less frequent. These binned data are telling us a lot about the sample distribution of flipper_length_mm.

It is still difficult to perceive distributional information when presented as a series of numbers. What we need is a visualisation based on the above numbers. This is what a **histogram** provides. Histograms summarise the sample distribution of a variable by showing the counts of binned data as a series of bars. The position and width of each bar correspond to an interval and the height shows the count. Here is a histogram based on the binned data we made:



This gives a clear summary of the sample distribution of body mass. It reveals:

- 1. the most common values, which are round about 190 mm;
- 2. the range of the data, which is about 60 mm; and
- 3. the shape of the distribution, which appears to be **bimodal**—it has two peaks.

That last observation wasn't immediately obvious when looking at the raw counts in the binned summary.

We used **ggplot2** to make that histogram. We could have used **dplyr** to make the binned data set and then used this with **ggplot2** to construct the plot manually. However, there is a much easier way to make a histogram. We'll demonstrate this now. Rather than do it using one **ggplot2** expression, we will break the process up to demonstrate how it works.

The first step uses the ggplot function with aes to set up the default data and aesthetic mapping:

```
# create graphical object with data and aesthetic mapping
plot_hist <- ggplot(penguins, aes(x = flipper_length_mm))</pre>
```

This is no different than the scatter plot example we stepped through. The only difference is that a histogram requires only one aesthetic mapping. We supplied the argument $x = flipper_length_mm$ to aes because we want to display the map intervals associated with $flipper_length_mm$ to the x-axis. We don't need

to supply an aesthetic mapping for the y-axis because **ggplot2** is going to handle this for us.

The second step adds a layer to the plot_hist object. We need to find the right geom_XX function to do this. This is called geom_histogram:

```
# use geom_histogram to add histogram layer
plot_hist <- plot_hist + geom_histogram()</pre>
```

It is instructive to look at the summary of the **ggplot2** object at this stage:

```
# print summary of graphical object
summary(plot_hist)
```

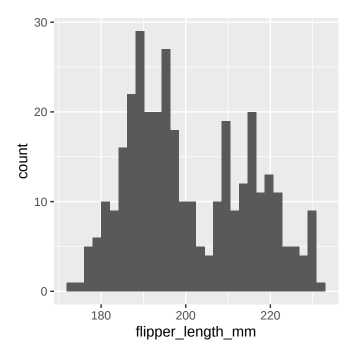
```
## data: species, island, bill_length_mm, bill_depth_mm,
     flipper_length_mm, body_mass_g, sex, year [344x8]
## mapping: x = ~flipper_length_mm
## faceting: <ggproto object: Class FacetNull, Facet, gg>
       compute_layout: function
##
       draw_back: function
       draw_front: function
##
##
       draw labels: function
##
       draw_panels: function
##
       finish_data: function
##
       init_scales: function
       map_data: function
##
##
       params: list
##
       setup_data: function
##
       setup_params: function
##
       shrink: TRUE
##
       train_scales: function
##
       vars: function
       super: <ggproto object: Class FacetNull, Facet, gg>
## geom_bar: na.rm = FALSE, orientation = NA
## stat_bin: binwidth = NULL, bins = NULL, na.rm = FALSE, orientation = NA, pad = FALSE
## position_stack
```

Look at the text below the ----. This shows that <code>geom_histogram</code> adds a 'stat' to the layer: <code>stat_bin</code>. This means <code>ggplot2</code> is going to take the raw <code>flipper_length_mm</code> data and bin it for us. Remember—the stat facility of <code>ggplot2</code> is its mechanism for creating summaries of data from the raw inputs. <code>stat_bin</code> deals with the binning process for us.

Everything we need to plot a histogram is now set up. Here is the resulting plot:

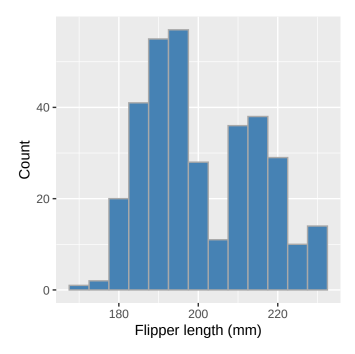
```
# show the histogram
print(plot_hist)
```

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



This is not quite the same as the example we saw above because it uses different bins. We can set the properties of the <code>geom_histogram</code> to tweak this kind of thing—the <code>binwidth</code> argument adjusts the width of the bins used. Let's construct the histogram again with 5 mm wide bins, as well as adjust the colour scheme and axis labels a bit:

```
# display flipper length histogram
ggplot(penguins, aes(x = flipper_length_mm)) +
    # use geom_histogram to add histogram layer + customisation
geom_histogram(binwidth = 5, fill = "steelblue", colour="darkgrey") +
    # set the the x- and y-axis labels
labs(x = "Flipper length (mm)", y = "Count")
```



Notice that the effect of increasing the bin width is to 'smooth' the histogram, i.e. this version looks less jagged than the last. Whether or not that colour scheme is an improvement is a matter of taste. Mostly we wanted to demonstrate how the fill and colour arguments change the output—fill sets the fill colour of the bar and colour deals with the line around each bar.



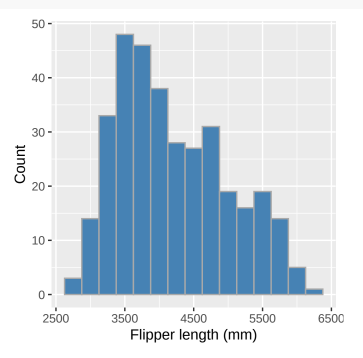
Choose your own bin widths

It is good practice to experiment with binwidth because the value selected by $\mathbf{ggplot2}$ is seldom optimal. Finding a 'good' value is as much art as science. One way of doing this is to start with a value that is probably too large and then refine it down. For example, we could work out the approximate range of the data by eye and then use $1/10^{\text{th}}$ of that range as an initial value for binwidth. Unless the data set is very small, this will result in a histogram that aggregates too much information. Once we have that rough starting value, we then reduce binwidth in small steps, re-rendering the plot each time until we end up with something that summarises the distribution well.

We can use pretty much the same R code to produce a histogram summarising the body mass sample distribution:

```
# display body mass histogram
ggplot(penguins, aes(x = body_mass_g)) +
```

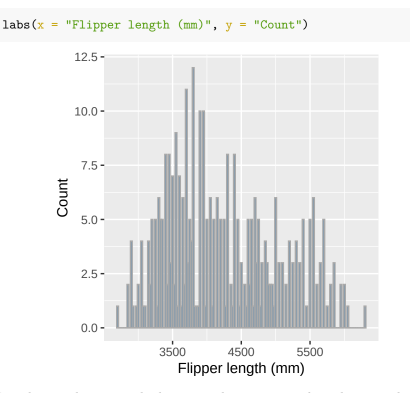
```
# use geom_histogram to add histogram layer + customisation
geom_histogram(binwidth = 250, fill = "steelblue", colour="darkgrey") +
# set the the x- and y-axis labels
labs(x = "Flipper length (mm)", y = "Count")
```



The only things that changed in this example were the aesthetic mapping and the bin width, which we set to 250 to reflect the range of observed body masses. The histogram reveals that the most common body mass (the mode) is around 3500 grams and the range of masses is also about 3500 grams. It also shows that the distribution is **skewed**, i.e. it is asymmetric. We can say even more—the distribution is right-skewed (a.k.a. positive-skewed) because the tail on the right gets further away from the mode than the left tail.

We have to choose those bin widths carefully. Body mass is measured to the nearest 25 grams. This means we should choose a bin width that is some multiple of 25 to produce a meaningful histogram. Look what happens if we set the bin width to 25:

```
# display body mass histogram
ggplot(penguins, aes(x = body_mass_g)) +
    # use geom_histogram to add histogram layer + customisation -- choose
# a sensible bin width
geom_histogram(binwidth = 25, fill = "steelblue", colour="darkgrey") +
# set the the x- and y-axis labels
```



We end up with gaps in the histogram because many bins do not include any data. The take-home message is that we have to pay attention to our data to produce meaningful summaries.

15.1.3 Dot plots

Histograms are good for visualising sample distributions when we have a reasonably large sample size, e.g. >100 observations. They are less effective when a sample is small. In this 'small data' situation, it is better to use something called a **dot plot**¹.

We have created subset of penguins that only retains female Adelie penguin observations from 2007, called penguins_small. This is considerably smaller than the original:

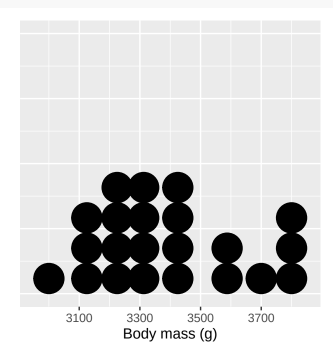
```
## # A tibble: 22 x 8
##
      species island
                         bill_length_mm bill_depth_mm flipper_length_mm body_mass_g
##
      <chr>
               <chr>
                                   <dbl>
                                                  <dbl>
                                                                     <int>
                                                                                  <int>
                                    39.5
##
    1 Adelie
               Torgersen
                                                   17.4
                                                                       186
                                                                                   3800
```

¹Not to be confused with the 'Cleveland dot plot'. A standard dot plot summarises a sample distribution. The Cleveland dot plot is something quite different, which summarises the frequencies of a categorical variable. It's meant to serve as a simple alternative to bar charts and pie charts.

```
##
    2 Adelie
              Torgersen
                                   40.3
                                                  18
                                                                     195
                                                                                 3250
   3 Adelie
              Torgersen
                                   36.7
                                                  19.3
                                                                     193
                                                                                 3450
##
   4 Adelie Torgersen
                                   38.9
                                                  17.8
                                                                                 3625
                                                                     181
   5 Adelie Torgersen
##
                                   41.1
                                                 17.6
                                                                     182
                                                                                 3200
              Torgersen
##
   6 Adelie
                                   36.6
                                                 17.8
                                                                     185
                                                                                 3700
##
   7 Adelie Torgersen
                                   38.7
                                                 19
                                                                     195
                                                                                 3450
   8 Adelie Torgersen
                                   34.4
                                                 18.4
                                                                                 3325
                                                                     184
   9 Adelie Biscoe
                                                 18.3
##
                                   37.8
                                                                     174
                                                                                 3400
## 10 Adelie Biscoe
                                   35.9
                                                  19.2
                                                                                 3800
                                                                     189
## # ... with 12 more rows, and 2 more variables: sex <chr>, year <int>
```

This is about the right size for using a dot plot. Constructing a dot plot with **ggplot2** is easy once you understand the histogram case. The code is are very similar:

```
# display body mass dot plot
ggplot(penguins_small, aes(x = body_mass_g)) +
    # use geom_dotplot to add layer + choose a sensible bin width!
geom_dotplot(binwidth = 100) +
    # set the the x-axis label and remove y-axis label
labs(x = "Body mass (g)", y = "") +
    # remove y-axis guides
theme(axis.text.y = element_blank(), axis.ticks.y = element_blank())
```



Each observation in the data adds one dot, and dots that fall into the same bin

are stacked up on top of one another. Notice the bins are not evenly spaced, however. The resulting plot displays the same information as a histogram but tends to be more informative when there are few observations.

The theme part of that example is entirely optional. It removes the grid lines and y-axis labels because these are not meaningful when the number of stacked dots represents 'height'. It is unfortunate that **ggplot2** displays them. But then, no package is perfect.

15.2 Exploring categorical variables

We will work with the penguin species (species) and observation year (year) variables to demonstrate how to explore categorical variables.

15.2.1 What kind of categorical variable?

Exploring categorical variables is generally simpler than the numeric case. The simplest thing we can do is examine the set of categories. For example, we can print these by extracting a variable with the \$ operator and using the unique function:

```
species —
```

```
unique(penguins$species)
```

```
## [1] "Adelie" "Gentoo" "Chinstrap"
```

observation year —

unique(penguins\$year)

```
## [1] 2007 2008 2009
```

These are obviously categorical—there are no numeric values associated with either variable. The obvious question to ask about each variable is, is it ordinal or nominal?

It is probably safe to assume that **species** should be treated as a nominal variable. There is no reasonable way to order the three species—i.e. a statement such as 'Adelie > Gentoo' is nonsense.

What about year? That's a bit more tricky. If we were interested in how some aspect of the data changes over time, we might consider treating year as a numeric variable, or perhaps, as an ordinal categorical variable. Alternatively, if the question is simply, 'do the data vary from one year to the next' without any concern for trends, it's reasonable to treat year as a nominal categorical variable.

This once again illustrates an important idea—the classification of a variable sometimes depends on the analysis goals. The decision matters because it influences how we choose to summarise a variable, how we interpret its relationship with other variables, and whether a specific statistical model is appropriate for our data or not.

For now, let's assume it's fine to treat year as a nominal variable.

15.2.2 Bar plots

We can't say anything about the sample distribution of a categorical variable by just looking at the possible categories. We need to explore the relative frequency of those categories. There is no need to carry out anything as complicated as binning. Instead, we simply count up the number of times each category occurs in the data. The **dplyr** count verb will does this:

```
penguins %>% count(species)
```

```
## # A tibble: 3 x 2
## species n
## <chr> <int>
## 1 Adelie 152
## 2 Chinstrap 68
## 3 Gentoo 124
```

We saw this result earlier in the book—the most common species is Adelie, followed by the Gentoo and Chinstrap. Gentoo is roughly as common as Adelie, whereas Chinstrap is about half as frequent.

These numbers lead directly to an informative visualisation. The usual graphical tool for summarising categorical variables is the **bar chart**. A bar chart presents summaries of grouped data with rectangular bars. The lengths of each bar is proportional to the value it represents. When summarising categorical variables, the bar lengths show the raw counts or proportions of each category.

Constructing a bar graph to display category counts is easy with **ggplot2**. We'll do this for the **species** variable. As always, we start by using the **ggplot** function to construct a graphical object containing the default data and aesthetic mapping.

```
# create graphical object with data and aesthetic mapping
plot_bar <- ggplot(penguins, aes(x = species))</pre>
```

We've called the object plot_bar for the obvious reason. Notice that we only need to define one aesthetic mapping—we mapped species to the x-axis. This will produce a bar plot with vertical bars.

The next step is to add a layer using one of the geom_XX functions. There are two

functions we can use to create bar charts in ggplot, geom_bar and geom_col. By default, geom_col counts the number of observations in each category, whereas geom_bar plots the actual numbers we provide.

We want to show the counts of each species category, meaning we should use geom_bar:

```
# use geom_bar to add bar plot layer
plot_bar <- plot_bar + geom_bar()</pre>
```

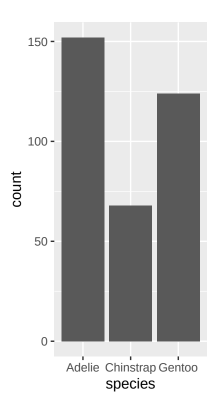
Now examine the summary of the **ggplot2** object:

```
# print summary of graphical object
summary(plot_bar)
```

```
## data: species, island, bill_length_mm, bill_depth_mm,
     flipper_length_mm, body_mass_g, sex, year [344x8]
## mapping: x = \simspecies
## faceting: <ggproto object: Class FacetNull, Facet, gg>
##
       compute_layout: function
##
       draw back: function
       draw_front: function
##
##
       draw_labels: function
##
       draw panels: function
##
       finish_data: function
##
       init_scales: function
##
       map_data: function
##
       params: list
##
       setup_data: function
##
       setup_params: function
##
       shrink: TRUE
##
       train_scales: function
##
       vars: function
##
       super: <ggproto object: Class FacetNull, Facet, gg>
## geom bar: width = NULL, na.rm = FALSE, orientation = NA
## stat_count: width = NULL, na.rm = FALSE, orientation = NA
## position stack
```

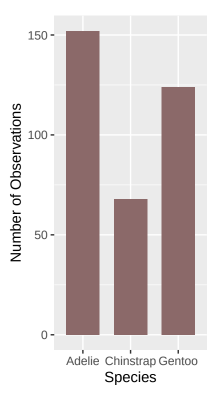
Look at the layer information below ----. The <code>geom_bar</code> function uses <code>stat_count</code>. This means <code>ggplot2</code> summed the number of observations associated with each category of <code>species</code>. Counting a categorical variable is analogous to binning a numeric variable. The only difference is that there is no need to specify bin widths because <code>species</code> is categorical. Here's the resulting figure:

```
# show the bar plot
print(plot_bar)
```



This is exactly the same information produced by the count function, only now it's presented in graphical form. We can customise this bar graph with functions like labs and by setting various properties inside <code>geom_bar</code>. For example:

```
# display bar plot of species counts
ggplot(penguins, aes(x = species)) +
    # use geom_bar to add bar plot layer of **counts**
geom_bar(fill = "rosybrown4", width = 0.7) +
    # set the the x- and y-axis labels
labs(x = "Species", y = "Number of Observations")
```



The only new thing here is that we used the width argument of geom_bar to make the bars a little narrower than the default. Notice that we use fill (not colour) to change the colour of the bars.

What might we like to change about this plot? The different species appear in alphabetical order. There's nothing particularly meaningful about that order. Perhaps it would make more sense to show them in frequency order from least to most common?

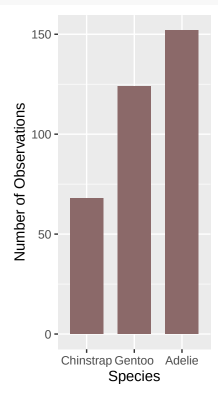
We need to customise the scale associated with the 'x' aesthetic to achieve this. We can start by making a short character vector containing the species names, listing them in the order they need to be shown:

```
# character vector of species names **in required order**
ords <- c("Chinstrap", "Gentoo", "Adelie")</pre>
```

Keep an eye on the spelling—R is not forgiving of spelling errors. We can use this with the limits argument of the scale_x_discrete function to adjust the ordering:

```
# display bar plot of species counts
ggplot(penguins, aes(x = species)) +
```

```
# use geom_bar to add bar plot layer of **counts**
geom_bar(fill = "rosybrown4", width = 0.7) +
# customise order of x-axis (limits)
scale_x_discrete(limits = ords) +
# set the the x- and y-axis labels
labs(x = "Species", y = "Number of Observations")
```

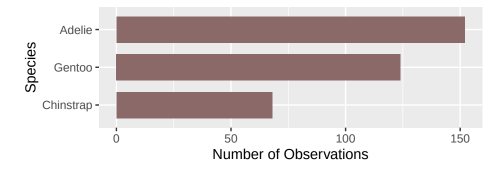


We had to use one of the scale_x_YY functions to change the way the 'x' aesthetic appears. We use scale_x_discrete because 'discrete' is ggplot2-speak for 'categorical'.

Is there anything else we might want to change? Sometimes it is clearer to plot horizontal bars, particularly if the categorical axis labels are all bunched together (not really the case here). Flipping the x and y axes makes a horizontal bar chart, which we can achieve with the coord_flip function (this is new):

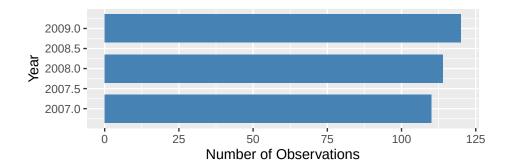
```
# display bar plot of species counts
ggplot(penguins, aes(x = species)) +
    # use geom_bar to add bar plot layer of **counts**
geom_bar(fill = "rosybrown4", width = 0.7) +
    # customise order of x-axis (limits)
```

```
scale_x_discrete(limits = ords) +
# set the the x- and y-axis labels
labs(x = "Species", y = "Number of Observations") +
# flip x- and y-axes
coord_flip()
```



We use very similar R code to produce a bar plot summarising the annual counts, i.e. those associated with the year categories:

```
# display bar plot of year counts
ggplot(penguins, aes(x = year)) +
    # use geom_bar to add bar plot layer of **counts**
geom_bar(fill = "steelblue", width = 0.7) +
    # set the the x- and y-axis labels
labs(x = "Year", y = "Number of Observations") +
    # flip x- and y-axes
coord_flip()
```

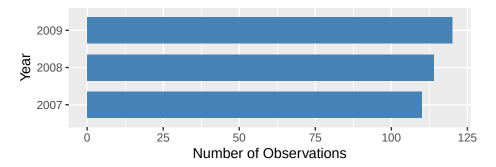


To make this, we changed the aesthetic mapping, colour and the labels, and dropped the scale_x_discrete part because the default ordering is acceptable. The resulting plot shows that there are roughly equal numbers of penguins

sampled in each year.

There is one small irritation. Look at the year axis labels and guides. These include non-existent 'half years' (e.g. 2007.5) rather than being limited to only the observed years (2007, 2008 and 2009). This has happened because **ggplot2** doesn't know that **year** is meant to be treated as a categorical variable. It sees a number, and so it constructs a continuous numeric axis for **year**.

We need to force **ggplot2** to treat year as a categorical 'thing' to avoid this behaviour. There is more than one way to do this. The simplest is to convert year into a character vector. The as.character function will do this. What's more, **ggplot2** will allow us to do this when we set up the aesthetic mappings within aes:



That's better! This may seem like a bit of an edge case but it comes up often enough to be worth highlighting. More generally, this example demonstrates that our conception of a variable needs to matches the way it is interpreted by **ggplot2** to arrive at a sensible plot.

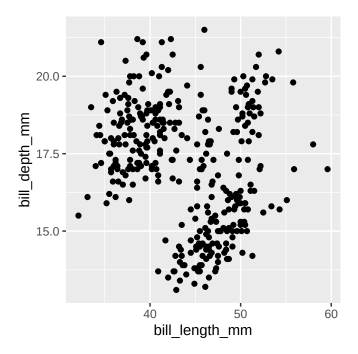
Chapter 16

Exploring associations

A pair of variables are said to be associated when knowing the value of one variable gives us information about the possible values of another. This chapter's main goal is to show how to use visualisations to explore associations among different kinds of variables.

16.1 Associations between numeric variables

The standard graph for displaying an association among a pair of numeric variables is the scatter plot. This uses horizontal and vertical axes to show two variables as a series of points. We saw how to construct a scatter plot in the Introduction to ggplot2 chapter. Here is the initial plot we made to show the association between bill depth and bill length (ignoring species information):



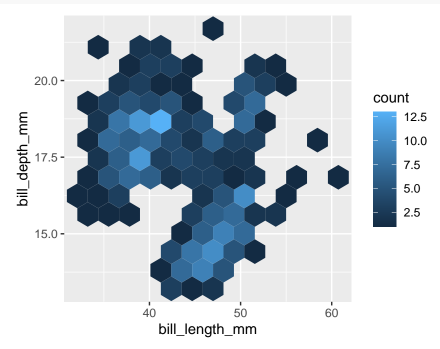
Most of the time, a scatter plot is exactly the right plot for visualising an association among numeric variables. One limitation of scatter plots is that they tend not to reveal associations when there is a lot of over-plotting happening. This occurs when:

- 1. there is a lot of data (e.g. 1000s of data points), meaning the distance between points is very small; or
- 2. there are many identical pairs of values, typically because both variables are discrete or a continuous variable is measured on a coarse scale.

In these circumstances, over-plotting of points tends to obscure the association. Sometimes we can mitigate this by making the points very small (problem #1) or semi-transparent (problem #1 or #2). But what should we do when those sorts of tricks fail to improve a plot? Luckily, **ggplot2** provides alternative geoms for visualising numeric associations in situations where over-plotting is a problem.

Two options for dealing with large data sets (problem #1) are the <code>geom_bin_2d</code> and <code>geom_hex</code> functions. These work like histograms in two dimensions. The <code>geom_bin_2d</code> divides the plane into rectangles, counts the number of cases in each rectangle, and then uses the number of cases to assign the rectangle's fill colour. The <code>geom_hex</code> function does essentially the same thing but instead divides the plane into regular hexagons.

Here's an example of geom_hex in action (n.b. geom_hex relies on the hexbin package, so this need to be installed to use it):



The bins argument controls how many hexagonal bins to use when spanning each dimension. The penguins data set is a bit small for this kind of plot (a scatter plot is actually better), so we had to use relatively coarse binning by setting the number of bins to 12.

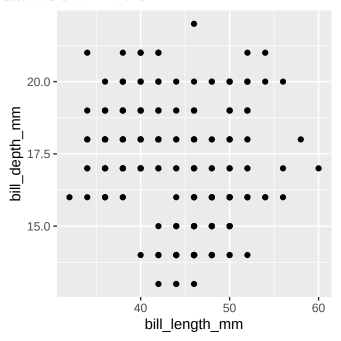
Notice how similar this looks to the **ggplot2** code for a scatter plot. We simply used **geom_hex** in place of **geom_point**. This reveals a powerful feature of **ggplot2**—once we have a working plot, we can often switch to a reasonable alternative by simply changing the type of geom. This makes it very quick to explore a data set in different ways.

What should we do if there are there are many identical pairs of values? For example, imagine that bill length had only been measured to the nearest 2 mm and bill depth had only been measured to the nearest 1 mm, like this:

```
## # A tibble: 344 x 2
## bill_length_mm bill_depth_mm
## <dbl> <dbl>
```

##	1	40	19
##	2	40	17
##	3	40	18
##	4	NA	NA
##	5	36	19
##	6	40	21
##	7	38	18
##	8	40	20
##	9	34	18
##	10	42	20
##	#	with 334 more rows	

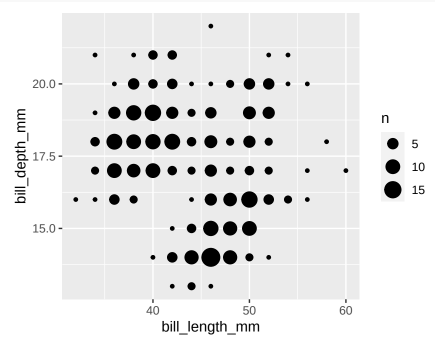
A scatter plot bill_depth_mm against bill_length_mm produced with those coarsened data would look like this:



This is not very informative because groups of points appear in exactly the same \mathbf{x}/\mathbf{y} location.

The geom_count function solves this problem by producing a scatter plot where point sizes are scaled according to how many cases are associated with each point. For example, the following code produces a modified scatter plot using a coarsened bill morphology data set (called penguins_coarse):

```
aes(x = bill_length_mm, y = bill_depth_mm)) +
# use geom_count to add sample size scaled points layer
geom_count()
```

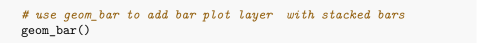


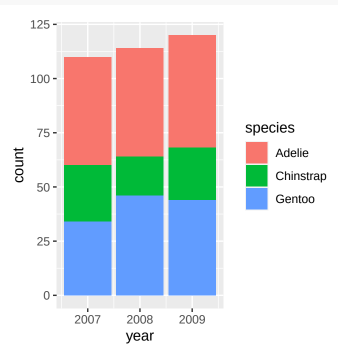
Again, notice how similar this looks to the **ggplot2** code for a standard scatter plot. We simply used **geom_count** in place of **geom_point**.

16.2 Associations between categorical variables

Bar charts are often used to summarise associations between categorical variables. The basic idea is to produce a separate bar for *each combination* of categories in the two variables. The lengths of each bar is proportional to the value it represents, which is either a raw count or the proportion in each category combination.

Using **ggplot2** to display this information is not very different from producing a bar graph to summarise a single categorical variable. Let's do this for the **species** and **year** variables in **penguins**:





We want to display information from two categorical variables, so we have to define two aesthetic mappings. We mapped the year variable (year) to the x axis and species (species) to the fill colour. Predictably, we construct the layer using geom_bar.

This produced a **stacked bar chart**. Each year has its own bar (x = year), and each bar has been divided up into different coloured segments, the length of which is determined by the number of observations associated with each species in that year (fill = species).

We have all the right information in this graph, but it could be improved in a couple of ways:

- Look at the guides on the x axis. There are some pointless extra lines. We saw this issue in a different guise in the previous chapter. It occurs because year is stored as a numeric vector—ggplot2 has no way of knowing we want to treat it as a non-numeric, categorical variable.
- The ordering of the species in the stacks does not reflect the relative frequency of each species. Again, we saw this problem in the previous chapter—ggplot2 treats does not 'know' how we want to order the species categories, and so it uses alphabetical order.

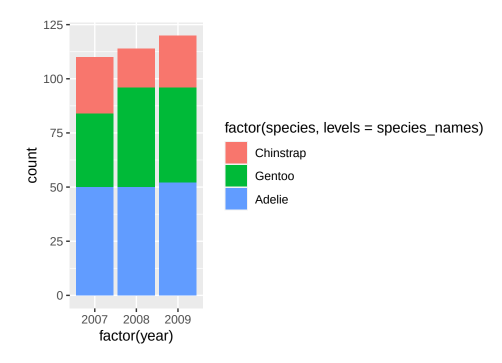
We need to ensure year is interpreted as a categorical variable and provide information about the required category order of species. R has a special kind of vector, called a **factor**, that is designed to tackle these kinds of issues. Factors are used by R to represent categorical variables.

We need to convert year and species to factors. How do we convert a numeric or character vector into a factor? Use the factor function (this is the first time we've mentioned this function). This has an argument called levels that sets the allowed categories and their order.

One way to start is by setting up a character vector of species names in the required order (species_names):

```
# levels of new species factor **in required order**
species_names <- c("Chinstrap", "Gentoo", "Adelie")</pre>
```

We supplied the names in the order we want them to appear. Be careful with the spelling—the values of species_names have to match those used in species. We can now remake the bar plot, this time converting species and year to factors 'on the fly':



This uses species_names to set the levels of species when we convert it to a factor. The stacking order of the bars then reflects their order in species_names. There was no need to set the levels of year because R will use the numeric order by default. Yes... the labels in that plot are ugly, but we could easily change these using labs function if we wanted to.



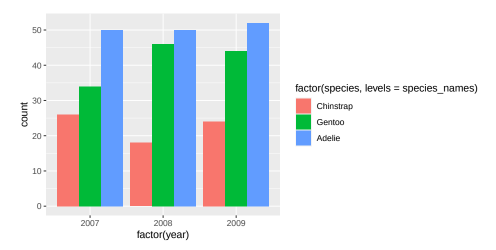
Factors

Factors are very useful and crop up all the time in R. A complete treatment of factors would require a whole new chapter. We've just shown one way to work with them via the factor function. This is enough to solve the reordering trick required to get ggplot2 to work the way we want it to, but there's a lot more to learn about factors. There is even a package (forcats) that exists solely to make working with factors a bit easier.

One thing to be aware of is that it is sometimes simpler to convert variables to factors in a data set, rather than doing it every time we use them with **ggplot2**. For example, we could use mutate to create factor versions of year and species:

Any ggplot2 code that uses the updated version of penguins would then automatically respect the intended categorical nature of year of and the species ordering we desire.

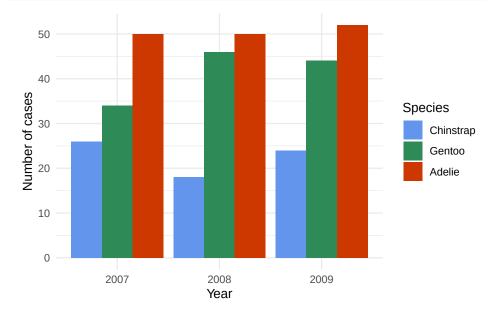
A stacked bar chart is the <code>geom_bar</code> default. If we want to know how two categorical variables are associated it is sometimes better to plot all bars side-by-side. This is not hard to do. We switch to a side-by-side bar chart by assigning a value of "dodge" to the <code>position</code> argument of <code>geom_bar</code>:



The position = "dodge" argument says that we want the bars to 'dodge' one another along the x axis. This slightly odd language is how **ggplot2** specifies the position of grouped items so they are displayed next to one another.

Remember—the customisation techniques introduced in the Customising plots chapter are completely general. For example, we could improve the plot by using labs to set the labels, setting custom fill colours with scale_fill_manual, and

applying a theme via theme_minimal:

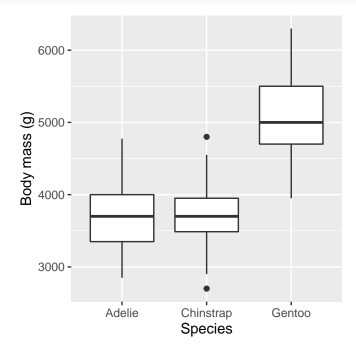


16.3 Categorical-numerical associations

The next obvious question is, "How do we display the association between a categorical and numeric variable?" As usual, there are a range of different options.

The most common is the 'box and whiskers plot' (or just 'box plot'). This is best understood by example. To construct a box and whiskers plot we need to set 'x' and 'y' axis aesthetics for the categorical and numeric variable; then use the <code>geom_boxplot</code> function to add the appropriate layer. To examine the association between penguin species (<code>species</code>) and body mass measurements

(body_mass_g), use:



We can see why this is called a box and whiskers plot. There are four components of each box and whiskers:

- The horizontal line inside the box is the sample median of the numeric variable—a measure of its central tendency. This allows us to compare the most likely value of the numeric variable across the different categories.
- The boxes display the interquartile range (IQR) of the numeric variable in each category, i.e. the middle 50% of each group. This allows us to compare the spread (=dispersion) of the numeric values across categories.
- The vertical lines above and below each box are the "whiskers". The interpretation of these depends on which kind of box plot we are making. **ggplot2** produces a traditional Tukey box plot by default. Each whisker is drawn from each end of the box (the upper and lower quartiles) to a well-

defined extreme point. To find where the upper whisker ends, we have to find the largest observation that is no more than 1.5 times the IQR above the upper quartile. The lower whisker ends at the most extreme point that is no more than 1.5 times the IQR below the lower quartile.

 Any cases that do not fall inside the whiskers are plotted as an individual point. These may be outliers, although they could also be perfectly consistent with the wider distribution.

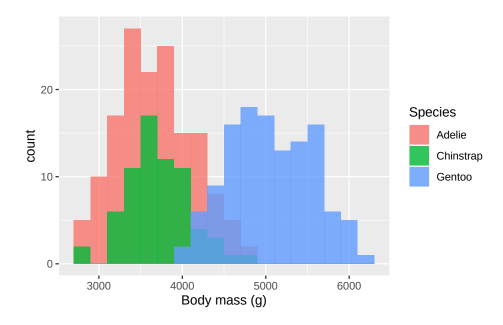
The resulting plot compactly summarises the distribution of the numeric variable within categories. It provides information about the central tendency, dispersion and skewness of each distribution. We also get a sense of whether there are potential outliers by noting the presence of individual points outside the whiskers.

What does the above plot tell us about body mass and species? It shows the body mass distributions of Chinstrap and Adelie penguins are very similar, in terms of both central tendency and dispersion. In contrast, Gentoo penguins are generally larger and exhibit great absolute variation. The Gentoo body mass distribution is also the only one that seems to display much skew.

16.3.1 Alternatives to box and whiskers plots

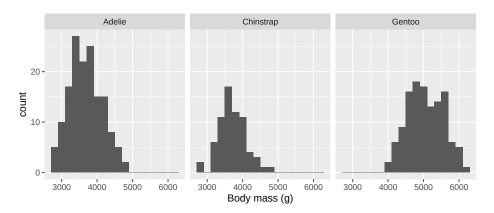
Box and whiskers plots are a good choice for exploring categorical-numerical associations. They provide a lot of information about how the distribution of the numeric variable changes across categories. Sometimes we may want to squeeze even more information about these distributions into a plot. One way to do this is to make multiple histograms.

We know how to make a histogram, and we have seen how aesthetic properties such as colour and fill are used to distinguish different categories of a variable. This suggests we could overlay more than one histogram on a single plot. Let's use this idea to see how the sample distribution of body mass differs among the species again:



We defined two mappings: the continuous variable (body_mass_g) was mapped to the x axis and the categorical variable (species) was mapped to the fill colour. Notice that we also set the position argument to "identity". This tells ggplot2 not to stack the histograms on top of one another. Instead, they are allowed to overlap. It's for this reason that we also made them semi-transparent by setting the alpha argument.

Plotting several histograms in one layer like this places a lot of information in one plot, but it can be hard to make sense of when the histograms overlap a lot. If the overlapping histograms are too difficult to interpret, we might consider producing a separate one for each category. We've already seen a quick way to do this. Faceting works well here:



The two histogram plots tell much the same story as the box and whiskers plot. The body mass distributions of Chinstrap and Adelie penguins are very similar, Gentoo penguins are generally a bit larger, but all three distributions overlap to some extent.

16.4 Multivariate associations

We have now examined several ways for summarising associations between two variables. How do we explore associations between more than two variables in a single graph? That is, how do we explore **multivariate associations**? It is difficult to give a concrete answer because it depends on the scientific question we're addressing, the kinds of variables we're working with, and to a large extent, our creativity and aptitude with a graphing framework like **ggplot2**.

We have already seen enough of **ggplot2** works to build some fairly sophisticated visualisations. The key point is that we don't really need to learn anything new to visualise multivariate associations. There are two ways to add additional information to a visualisation, both of which we have already reviewed:

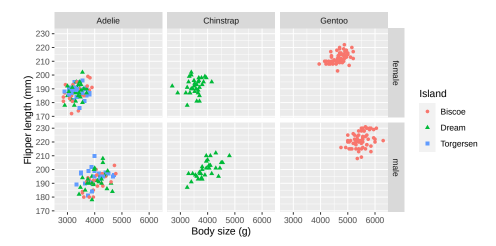
- 1. Define aesthetic mappings to allow a layer's properties to depend on the values of one or more variables.
- 2. Use faceting to construct a multi-panel plot according to the values of categorical variables.

Constructing a multivariate visualisation is a matter of combining these approaches to squeeze information from several variables into a single graph.

We saw these approaches used together in the Introduction to ggplot2 chapter. Let's look at one more example to cement the idea. We want to understand how the flipper length relates to body mass and how this numeric-numeric relationship varies by species, sex and island. That's five variables in one plot!

The obvious idea is to produce a multi-panel scatter plot, where each panel is associated with combinations of the categorical variables. We have already seen how to do this kind of thing with facet_wrap. This time, we'll use the

facet_grid function to organise separate panels for each sex-species combination into a nice 2d grid:



Simple. We set some aesthetics and used faceting to squeeze the five variables into one plot. The facet_grid function represents the values two or more categorical variables by row and column position, which means we have to set rows and cols arguments. The vars(...) bit is not optional by the way. We also took the opportunity to introduce a couple of new tricks:

- We used the base R na.omit function on penguins to strip out any rows with a missing value. This avoids introducing an 'NA' category in the faceting. This trick can be a bit dangerous because na.omit removes every case where a value is missing.
- We introduced some redundancy by mapping one variable (island) to

two different aesthetics (colour and shape). This achieves two things. It makes it easier to differentiate the points, and it ensures a plot can be understood if printed in black and white.

There is lots of information in this plot—for example, the association between flipper length and body mass is weakly positive, sexual dimorphism in body size seems to be greatest in Gentoo penguins, and the Adelie species is the only one present on all three islands.

Chapter 17

Doing more with ggplot2

We have now seen a range of different ways to visualise and explore data using **ggplot2**. This last chapter will cover a few bits of miscellanea that don't particularly fit anywhere else. That doesn't mean the ideas aren't important!

17.1 Comparing descriptive statistics

So far, we have focussed on displaying the raw data (e.g. scatter plots) or a distributional summary (e.g. box plots). What other types of quantities might we need to visualise? Descriptive statistics such as the sample mean are one possibility. These often feature in data analysis when 'comparing groups'.

We need to know how to construct plots that display such summaries. Let's start with a simple question: how does the (arithmetic) mean body mass varies by sex and across penguin species? One option is to produce a bar plot for which the lengths of bars represent the mean body mass in each category.

There are two different routes to produce this with **ggplot2**. One way to build such a plot is to break the problem into two steps. In the first step, we calculate whatever it is we want to display, i.e. the species- and sex-specific mean body mass. **dplyr** is usually the best tool to use for this first step:

```
penguins_summary <- penguins %>%
  # group data by species and penguins
group_by(species, sex) %>%
  # calculate mean body mass
summarise(mean_mass = mean(body_mass_g)) %>%
  # remove rows generated by sex = NA cases
na.omit()
```

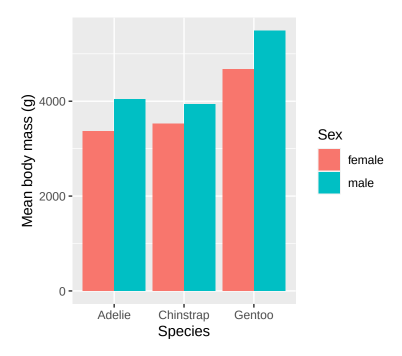
`summarise()` has grouped output by 'species'. You can override using the `.groups` argument.

We used group_by and summarise to calculate the set of means, which we decided to call mean_mass. Notice that we used na.omit to exclude the NA categories that arise when sex is unknown. The resulting penguins_summary object is a small 6 row by 3 column tibble:

```
# show the summary data
penguins_summary
```

```
## # A tibble: 6 x 3
## # Groups: species [3]
    species sex
                     mean_mass
    <chr>
##
              <chr>
                         <dbl>
## 1 Adelie
              female
                         3369.
## 2 Adelie
              male
                         4043.
## 3 Chinstrap female
                         3527.
## 4 Chinstrap male
                         3939.
## 5 Gentoo
              female
                         4680.
## 6 Gentoo
              male
                         5485.
```

The second step uses penguins_summary with ggplot2 to create the required bar plot:



Two points about this are worth noting:

- 1. We used geom_col instead of geom_bar. geom_bar counts the observations in each category by default. Using geom_col tells ggplot2 to plot the information in mean_mass 'as is'.
- We paid close attention to the names of things. The plotting data lives in penguins_summary which means the y aesthetic must be associated with mean_mass rather than body_mass_g.

It is possible to produce this plot with less code by using **ggplot2**'s built-in 'stat' facility (see box). We recommend the long-winded way when starting out with R because it makes it a bit easier to fix mistakes—we can check whether the right information is in a summary data set before plotting it.



Using ggplot2 'stats'

Another way to arrive at the above bar plot is by using the built-in **gg-plot2** stat facility:

```
# display bar plot of body mass means
ggplot(na.omit(penguins), # <- remove missing values</pre>
                                  # aesthetic mappings: mass (y) vs species (x) by sex (fill)
                                  aes(x = species, fill = sex, y = body_mass_g)) +
         # use geom_bar to add bar plot layer using a stats summary
         geom_bar(stat = "summary", fun.y = mean, position = "dodge") +
         # specify labels for all mappings
         labs(x = "Species", y = "Mean body mass (g)", fill = "Sex")
## No summary function supplied, defaulting to `mean_se()`
                      Mean body mass (g) 4000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000 - 2000
                                                                                                                                                                                                                                          Sex
                                                                                                                                                                                                                                                                female
                                                                                                                                                                                                                                                                 male
                                                  0 -
                                                                                                                         Chinstrap
                                                                                                                                                                               Gentoo
                                                                              Adelie
                                                                                                                         Species
```

This involves a couple of new tricks. When we add a layer using <code>geom_bar</code> we set two additional arguments:

- stat = "summary" tells ggplot2 not to plot the raw values of the y aesthetic mapping, but instead, to calculate and then plot a summary of the 'y' variable.
- fun.y = mean tells ggplot2 how to summarise the 'y' variable. The function on the right-hand side can be any function that takes a vector of values and returns a single number. We want the mean function.

Notice that this works on the raw data (penguins)—there is no need to

manually calculate the means because that happens inside **ggplot2**.

17.1.1 Error bars

A descriptive statistic like the mean isn't much use on its own. To properly interpret means we also need to know something about their uncertainty. There are many ways to quantify the uncertainty of an estimate. Whichever one we are using, displaying the uncertainty typically involves adding **error bars** to a plot.

We'll demonstrate how to do this by extending the previous example to show the mean and standard error of body mass for each species.



Er.... standard error?

The standard error is a measure of how precise an estimate like the sample mean is. A small standard error indicates that we can have more confidence the estimate reflects the 'true' value. The standard error of the mean can be calculated from a well-known formula:

$$\text{Standard Error} = \frac{\text{Standard Deviation}}{\sqrt{(\text{Sample Size})}} = \frac{\sigma}{\sqrt{n}}$$

It is hard to give an explanation of that formula without knowing a bit of statistical theory—it is just one of those things we have to learn. In terms of code that works with **dplyr**, that calculation looks like this:

```
standard_error = sd(x) / sqrt(n())
```

This assumes the numeric variable we are working with is called x, and we want the result to be called $\operatorname{standard_error}$. Notice this uses a special dplyr function called n to find the sample size.

Once again, constructing the plot is a two-step process. Start by calculating the means and standard error of body mass for each species and sex combination:

```
penguins_summary <- penguins %>%
  # group data by species and penguins
group_by(species, sex) %>%
  # calculate summaries
summarise(
  # mean mass
  mean_mass = mean(body_mass_g, na.rm = TRUE),
  # standard error
```

```
ster_mass = sd(body_mass_g, na.rm = TRUE) / sqrt(n())
) %>%
# remove rows generated by sex = NA cases
na.omit()
```

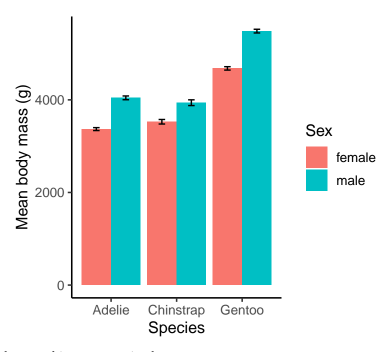
`summarise()` has grouped output by 'species'. You can override using the `.groups`

We use **dplyr** to calculate the means and standard errors of each species' body mass, which we called **mean_mass** and **ster_mass**, respectively. Again, we used **na.omit** to exclude the **NA** categories cases. The new **penguins_summary** object is a 6 row by 4 column tibble:

```
# show the new summary data penguins_summary
```

```
## # A tibble: 6 x 4
## # Groups: species [3]
##
    species sex
                    mean mass ster mass
##
    <chr> <chr>
                       <dbl>
                                 <dbl>
## 1 Adelie female
                        3369.
                                  31.5
## 2 Adelie male
                       4043.
                                  40.6
## 3 Chinstrap female
                        3527.
                                  48.9
## 4 Chinstrap male
                        3939.
                                  62.1
## 5 Gentoo
             female
                        4680.
                                  37.0
## 6 Gentoo
              male
                        5485.
                                  40.1
```

Next we use penguins_summary to make the plot. We can use geom_col to show the means as a bar plot, but what about the error bars? Use the unsurprisingly named geom_errorbar to add those! Here is the code:



The only new things appear in the geom_errorbar part:

- The ymin and ymax arguments of the geom_errorbar function give the lower and upper limits of error bars. Here, we have plotted the mean +/-1 standard error, i.e. each error bar is two standard errors long.
- Use the width argument to set the width of the 'hat' on each error bar. A value of zero gets rid of it altogether.
- The position has to be set in two places: geom_col and geom_errorbar. For a bar plot, we use the not-at-all-intuitive position = position_dodge(0.9) to locate the error bars at the centre of each bar.

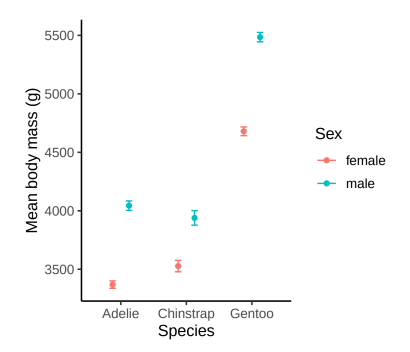


Warning!

Data visualisations can include many different kinds of 'error bar'— standard deviation, standard error, 95% confidence intervals, etc. When we include any error bars on a plot, we must remember to state which kind was used in the figure legend. Otherwise, readers will have no way of knowing how to interpret the plot.

17.1.2 Alternatives to bar plots

Do we have to use a bar plot to display means? Certainly not. In some ways bar plots are a bit old fashioned. People often prefer to show a set of means as points with error bars. This kind of visualisation is very easy to make now that we already have working bar plot code:



We made three small changes to the bar plot code to produce this. We...

- mapped sex to the colour aesthetic instead of fill,
- switched from using geom_col to geom_point, and
- altered the position adjustments to 'dodge' by a smaller amount.

That's it!

17.2 Adding text annotations

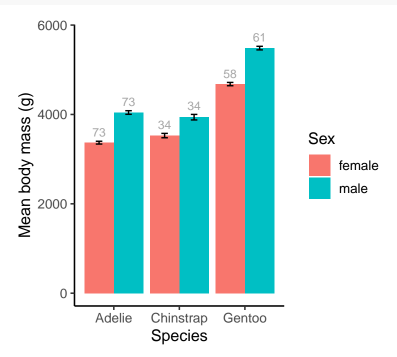
We often need to include text annotations on a plot. For example, we might want to show sample sizes or highlight 'statistically significant' differences. Adding text is no different from adding other kinds of objects to a plot—we use an appropriate geom (e.g. geom_text) to include textual information. Let's see how this works by adding sample size labels the bar chart we've been building.

We start by placing the labels we want to use in a data frame (or tibble) alongside any other variables used in aesthetic mappings. This can be distinct from the data set used to construct the plot but its often simplest to put everything together in one place. Let's do that by rebuilding the penguins_summary data set to include the mean body mass, standard error and sample size for each group.

```
penguins_summary <- penguins %>%
  # group data by species and penguins
group_by(species, sex) %>%
  # calculate summaries
summarise(
  # mean mass
  mean_mass = mean(body_mass_g, na.rm = TRUE),
  # sample size
  samp_size = n(),
  # standard error
  ster_mass = sd(body_mass_g, na.rm = TRUE) / sqrt(n())
) %>%
  # remove rows generated by sex = NA cases
na.omit()
```

 $\verb|##"summarise()" has grouped output by 'species'. You can override using the `.groups` argument.$

Then we add the text showing the sample size to our plot using the function geom_text:



The y = mean_mass + ster_mass + 200 aesthetic mapping in geom_text positions each number just above the error bar. This works because we placed the textual information in the plotting data set, meaning we can use the x/y variables to set the position of the annotations. We also have to use the correct position adjustment to ensure the annotations are located at each bar's centre. Everything else inside geom_text is adjusting the appearance of the actual text.

17.3 Saving plots

Plots can be saved using the **Export** button in the RStudio **Plots** tab. However, saving plots this way often leads to a pixelated, low resolution image. The **ggsave** function in **ggplot2** produces much better output and supports a wide range of image file types. There's not much to it. Using **ggsave** looks like this:

```
ggsave("penguins-plot.pdf", height = 5, width = 5)
```

By default, ggsave will save the last plot we made to file. If we had made the bar chart and then ran that ggsave line, R would save a copy of the plot to as a PDF file called penguins-plot.pdf.

Here is a quick summary of how the function works:

- The first argument is the path and name of the file we want to create. This example would save the plot to a file called <u>penguins-plot.pdf</u>. Because we only provide a file name, ggsave would save it to the current working directory.
- ggsave supports many different formats (e.g. "pdf", "jpeg", "png", "svg"). This can be set via the device argument (e.g. device = "jpeg"). Alternatively, if we don't set the format, ggplot2 will guess it from the file extension.
- The width and height arguments specify the plot dimensions. These usually require a bit of experimentation to get right. If we do not specify them, ggsave will use the current size of the plotting window.

17.4 Multi-panel plots

We have seen how to use facet_wrap and facet_grid to produce multi-panel plots. These functions are useful when a plot needs to show the same visualisation for different subsets of data. How do we produce a figure that shows a different visualisation in each panel?

There are many ways to do this using an external add-on package. We'll examine the framework provided by the **cowplot** package. Cowplot provides various features that help create high-quality figures, including custom themes and facilities to arrange plots into multi-panel figures.

Constructing a multi-panel plot with **cowplot** happens in two stages. First, make the individual component plots using **ggplot2** as usual, but instead of 'printing' them, assign each graphical object a name. Then use the **plot_grid** function from **cowplot** to construct the multi-panel plot from the stored objects.

For example, assume we want to display at the bill morphology scatter plot and the mean body mass bar chart side-by-side. First we make the two plots assign them names (with minimal commenting to save space):

```
# first plot
plt_a <-
    ggplot(penguins,
        aes(x = bill_length_mm, y = bill_depth_mm, colour = species)) +</pre>
```

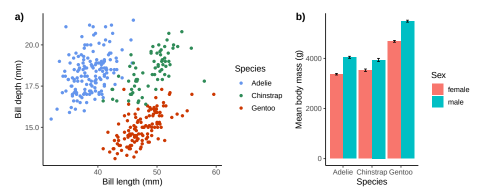
Then we use **cowplot**'s **plot_grid** to produce the two-panel plot:

```
plot_grid(
    # two-panel plot

plt_a, plt_b,
    # set location and size of panels

nrow = 1, rel_widths = c(60, 40),
    # set the panel labels

labels = c("a)", "b)"), label_size = 14
)
```



Once we've made a multi-panel plot, we can use the ggsave function to save it as before. The plot_grid function allows the figure to be customised in various ways. For example:

- nrow and ncol control the number of plots in each row/column,
- labels sets the labels used for each panel,
- label size controls the size of the label font, and
- rel_widths and rel_heights control how much space each plot gets.

The package website has some good articles that explain how this all works (along with the many other **cowplot** facilities).

Appendix A

Getting help

A.1 Introduction

R has a comprehensive built-in help system orientated around the base R functions, and every good external package also comes with its own set of **help files**. These provide information about individual package functions and summarise the included data sets. They also sometimes give descriptions of how different parts of the package should be used, and if we're lucky, one or more 'vignettes' that offer a practical demonstration of how to use the package.

We may as well get something out of the way early on. The word 'help' in the phrase 'help file' is a bit of a misnomer. It's more accurate to say R has an extensive **documentation** system. Help files are designed first and foremost to carefully document the different elements of a package, rather than explain how a particular function or the package as whole should be used to achieve a given end. They are aimed more at experienced users than novices. That said, help files often contain useful examples, and many package authors do try to make our life easier by providing functional demonstrations of their package.

It is important to get to grips with the built in help system. It contains a great deal of useful information which we need to really start using R effectively. The road to enlightenment is bumpy though.

A.2 Browsing the help system

Help files are a little like mini web pages, which means we can navigate among them using hyperlinks. One way to begin browsing the help system uses the help.start function:

help.start()

When we run this function the **Package Index** page should open up in the **Help** tab of the bottom right pane in RStudio. This lists all the packages currently installed on a computer. We can view the help files associated with a package by clicking on the appropriate link. For example, all the functions that come with the base installation of R have a help file associated with them—we can click on the link to the R base package (base) to see these.

The packages that we install separately each have their own set of associated help files. We will see how to navigate these in a moment.

The help browser has Forward, Back, and Home buttons, just like a normal web browser. If we get lost in the mire of help pages we can always navigate backward until we get back to a familiar page. Clicking on the home button takes us to a page with three sections:

- 1. The Manuals section looks like it might be useful for novice users. Unfortunately, it's not. Even the "Introduction to R" manual is only helpful for someone who understands what terms like 'data structure' and 'data type' mean. The others are more or less impenetrable unless the reader already knows quite a bit about computing in general.
- 2. The **Reference** section is a little more helpful. The "Packages" link takes us to the same page opened by help.start. From here we can browse the help pages on a package-specific basis. The "Search Engine & Keywords" link takes us to a page we can use to search for specific help pages, either by supplying a search term or by navigating through the different keywords.
- 3. The Miscellaneous Material section has a couple of potentially useful links. The "User Manuals" link lists any user manuals supplied by package authors. The "Frequently Asked Questions" link is definitely worth reviewing at some point, tough again, most of the FAQs are a little difficult for novice users to fully understand.

A.3 Searching for help files

After browsing help files via help.start for a bit it quickly becomes apparent that this way of searching for help is very inefficient. We often know the name of the function we need to use and all we want to do is open that particular help file. We can do this using the help function:

```
help(topic = Trig)
```

After we run this line RStudio opens up the help file for the trigonometry topic in the **Help** tab. This file provides information about the various trigonometric

functions such as sin or cos (we'll see how to make sense of such help pages in a bit).

The help function needs a minimum of one argument: the name of the topic or function of interest. When we use it like this the help function searches across packages, looking for a help file whose name gives an exact match to the name we supplied. In this case, we opened the help file associated with the Trig topic.

Most of the time we use the help function to find the help page for a specific function, rather than a general topic. This is fine if we can remember the name of the topic associated with different functions. Most of us cannot. Luckily, the help function will also match help pages by the name of the function(s) they cover:

```
help(topic = sin)
```

Here we searched for help on the sin function. This is part of the Trig topic so help(topic = sin) brings up the same page as the help(topic = Trig).

By default, the help function only searches for files associated with the base functions or with packages that we have loaded in the current session with the library function. If we want to search for help on the mutate function—part of the dplyr package—but we haven't run library(dplyr) in the current session this will fail:

```
help(mutate)
```

Instead, we need tell help where to look by setting the package argument:

```
help(mutate, package = "dplyr")
```

Even very experienced R users regularly forget how to use the odd function and have to dive into the help. It's for this reason that R has a built in shortcut for help accessed via? For example, instead of typing help(topic = sin) into the Console we can bring up the help page for the sin function by using? like this:

```
?sin
```

This is just a convenient shortcut that does the same thing as help. The only difference is that? does not allow us to set arguments such as package.

A.4 Navigating help files

Navigating help files is a little daunting at first. These files can be quite long and contain a lot of technical jargon. The help files associated with functions—

the most common type—do at least have a consistent structure with a number of distinct sections. Wrestling with a help file is much easier if we at least understand what each section is for. After the title, there are eight sections we need to know about:

- 1. **Description** gives us a short overview of what the function is meant to be used for. If the help page covers a family of related functions it gives a collective overview of all the functions. Always read this before diving into the rest of the help file.
- 2. Usage shows how the function(s) are meant to be used. It lists each member of the family as well as their common arguments. The argument names are listed on their own if they have no default, or in name-value pairs, where the value gives the default should we choose not to set it ourselves.
- 3. **Arguments** lists the allowed arguments along with a short description of what they do. This also tells us what what kind of data we're allowed to use with each argument, along with the allowable values (if relevant). Always read this section.
- 4. **Details** describes precisely how the function(s) behave, often in painful detail. This is often the hardest-to-comprehend section. We can sometimes get away with ignoring this section but when we really want to understand a function we need to wade through it.
- 5. Value explains what kind of object a function returns when it finishes doing whatever it does. We can often possible to guess what this will be from the type of function, but nonetheless, it is a good idea to check whether our reasoning is correct.
- 6. **References** just lists the key reference(s) for when if we really need to know the 'hows' and 'whys' of a function. We can usually skip this information. The one exception is if the function implements a particular analysis tool. It's best to know how such tools work before trying to use them.
- 7. See Also gives links to the help pages of related functions. These are usually functions that do something similar to the function of interest or are meant to be used in conjunction with it. We can often learn quite a bit about packages or related functions by following the links in this section.
- 8. Examples provides one or more examples of how to use the function. These are stand-alone examples, so there's nothing to stop us running them. This is often the most useful section of all. Seeing a function in action is a very good way to cut through the jargon and understand how it works.

A.5 Vignettes

The purpose of a package vignette is to provide a relatively brief, practical account of one or more of its features. Not all packages come with vignettes, though the best packages often do. We use the vignette function to view all the available vignettes in Rstudio. This will open up a tab that lists each vignette under their associated package name along with a brief description. A package will often have more than one vignette.

215

If we just want to see the vignettes associated with a particular package, we have to set the package argument. For example, to see the vignettes associated with **dplyr** we use:

```
vignette(package = "dplyr")
```

Each vignette has a name (the "topic") and is available either as a PDF or HTML file (or both). We can view a particular vignette by passing the vignette function the package and topic arguments. For example, to view the "grouping" vignette in the dplyr package we would use:

```
vignette(topic = "grouping", package = "dplyr")
```

The vignette function is fine, but it is more convenient to browse the list of vignettes inside a web browser. This allows us to open a particular vignette directly by clicking on its link, rather than working at the Console. We can use the browseVignettes function to do this:

```
browseVignettes()
```

This will open a page in our browser showing the vignettes we can view. As usual, we can narrow our options to a specific package by setting the package argument.

Appendix B

Managing projects, scripts and data files

Content will be added in block three.